Kinetics of Procalcitonin in the Management of Small Bowel Obstruction: A Preliminary Report

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Abstract

Background: Small Bowel Obstruction (SBO) is mainly due to adhesions acquired after abdominal surgery. Its management could be conservative or surgical but the choice is difficult because of the absence of clinico-biological markers. Serum procalcitonin (PCT) has previously been proposed as a biomarker. The aim was to present the kinetics’ profiles of serum PCT in cases of Conservative Management (CM) success and surgical management (SM) in patients with SBO.

Study design: From January to October 2013, 59 patients with adhesion-related SBO were included in a prospective, monocenter, non-randomized, clinical study. PCT was measured during 3 days maximum (or until the transit was restored), every six hours. The patients were divided into subgroups (conservative (CM) and surgical management (SM)). The predictive time points were identified with ROC curves.

Results: Patients in CM group (n=47) presented at admission a lower PCT and a higher chloride than in the SM group (n=12) (p<0.003). The time points of PCT for predicting SM are at admission (threshold>0.165 ng/mL; sensitivity (Se)=83.3% and negative predictive value (NPV)=93%), 18 hours after admission (threshold >0.275 ng/mL; Se=100% and NPV=100%) and 24 hours after admission (threshold>0.255 ng/mL; Se=83.3% and NPV=95%).

Conclusion: PCT is helpful in SBO’s management. The more predictive time points are at admission, 18 hours and 24 hours after admission to identify the patients requiring surgery.

Keywords: Procalcitonin, Kinetics; Small bowel obstruction

Materials and Methods

Population

From January to October 2013, 59 patients with adhesion-related SBO were included in a prospective, monocenter, non-randomized, clinical study (PCT kinetics’ study). The inclusion criteria were SBO diagnosed with CT scan and a clinical examination. The exclusion criteria were early postoperative obstruction, obstruction with neoplasia, obstruction with inflammatory bowel disease, colon obstruction, a history of abdominal radiotherapy, the presence of pneumoperitoneum, clinical signs of infection, age under 18, pregnancy, guardianship. Small bowel obstruction was diagnosed on the presence of standard clinical signs (abdominal pain, tenderness, nausea or vomiting, no passage of gas and/or stools) and radiological signs (presence of air fluid levels, dilated bowel loops) possibly secondary to an abdominal surgery.

The PCT kinetics’ study participants were divided into two groups: patients in whom the CM was successful (the CM group, n=47) and those in whom it had failed (the SM group, n=12).
Management

Initially, each patient was managed conservatively with nasogastric tube suction. If renewed passage of gas and stools was observed, the nasogastric tube was clamped and removed; oral nutrition was resumed and the CM was considered to be a success. When faced with clinical signs (such as fever, abdominal pain) or biological signs (hyperleucocytosis i.e. leucocyte count ≥ 16×10³/mL) and the absence of gas 48 hours after the initiation of CM, the conservative approach was considered to be a failure and the patient was scheduled for surgery.

Surgery

The open surgical procedure consisted in gut viscerolysis and (in some cases in presence of ischemia) bowel resection. An anastomosis was created depending of local conditions. In each case, the presence or absence of intra-operative bowel ischemia or necrosis was noted. Any adherence was described by the surgeon and noted in the patient’s case report form.

Study design

The PCT kinetics’ study’s primary endpoint was to present the kinetics’ profiles of serum PCT in cases of CM success and SM. The secondary endpoints were the identification of the more predictive time points to discriminate between CM success and SM and the predictive value of PCT for bowel ischemia. The study’s patient disposition is shown in Figure 1.

The following items of patient data were collected: demographic information, laboratory blood test results (white blood cell count, platelet count, C reactive protein, urea, lactate, ions, ASAT, ALAT, and PCT), comorbidities (diabetes, hypertension, chronic kidney disease or angor), and the history of abdominal surgery. Data related to surgical management included surgical recommendations, intra-operative signs and the surgical technique.

The procalcitonin assay

For each included patient, PCT test had been done on serum samples and measured on arrival to the emergency department and in the digestive surgery department during 3 days maximum (or until the transit was restored), every six hours. The Kryptor® T.R.A.C.E® assay (B.R.A.H.M.S, Clichy, France), routine laboratory test, was used on-site in the Biochemical laboratory. The reference value for PCT is <0.5 ng/mL.

Statistical analysis

Data are expressed as mean ± Standard Deviation (SD) or median (minimum – maximum) or number (percentage of patients). Univariate analyses were performed with the chi-squared test or Fisher’s exact test for qualitative variables and the Student t test for quantitative variables.

The threshold for statistical significance was set to p ≤ 0.05. To identify the more predictive time points of PCT we generated Receiver Operating Characteristic (ROC) curves and calculated the area under the ROC curve (AUROCC). The predictive time points were presented with the sensitivity (Se), specificity (Sp), Positive Predictive Value (PPV), Negative Predictive Value (NPV) and the threshold. Statistical analysis was performed with SAS 9.2 (SAS Institute Inc., Cary, NC, USA) and PASW 18 (SPSS INC., Chicago, IL, USA).

Results

Baseline characteristics of the population

The characteristics of the population are shown Table 1. There is an imbalance between the CM and SM groups in terms of PCT (p=0.002) and Chloride (p=0.003) i.e. patients in CM group presented at admission a lower PCT and a higher chloride than in the SM group.

Kinetics’ profiles of serum PCT

The kinetics’ profiles of serum PCT and the evolution of the ratio between PCT in the CM and SM are presented respectively in Figures 2 and 3. The PCT levels are always lower in the CM group than in the SM one. The ratio between PCT measured in the CM and SM groups is comprised between 2 and 4 during the first 36 hours after the admission, and is high as 20 at 48 hours. The rate of CM failure considering the PCT values is depicted in Figure 4. This rate is correlated (R²=0.989) with the serum PCT of which values correspond to the 2-folded; 3-folded and 4-folded predictive threshold at admission.

Predictive time points of PCT to discriminate between CM success and SM

At admission: Using a ROC curve, the AUROCC for PCT at admission and SM was 0.802 [95% CI, 0.65–0.954] (Figure 5). A PCT threshold >0.165 ng/mL for predicting SM yielded a sensitivity of 83.3%, a specificity of 61%, a PPV of 35.7% and an NPV of 93.3%.

18 hours after the admission: Using a ROC curve, the AUROCC for PCT 18 hours after the admission and SM was 0.89 [95% CI, 0.0 – 1] (Figure 6). A PCT threshold >0.275 ng/mL for predicting SM yielded a sensitivity of 100%, a specificity of 75%, a PPV of 30% and an NPV of 100%. 
Table 1: Baseline characteristics of the population.

<table>
<thead>
<tr>
<th>Variable</th>
<th>Conservative management (n = 47)</th>
<th>Surgical management (n = 12)</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean age, years (min – max)</td>
<td>65 (24 – 98)</td>
<td>76 (19 – 98)</td>
<td>0.15</td>
</tr>
<tr>
<td>Male sex, n (%)</td>
<td>20 (43)</td>
<td>4 (33)</td>
<td>0.56</td>
</tr>
<tr>
<td>Mean body mass index, Kg/m² (min – max)</td>
<td>29 (20 – 62)</td>
<td>25 (21 – 33)</td>
<td>0.10</td>
</tr>
<tr>
<td>Diabetes, n (%)</td>
<td>3 (6)</td>
<td>1 (8)</td>
<td>0.81</td>
</tr>
<tr>
<td>Cancer, n (%)</td>
<td>13 (28)</td>
<td>3 (25)</td>
<td>0.85</td>
</tr>
<tr>
<td>Hypertension, n (%)</td>
<td>19 (40)</td>
<td>5 (42)</td>
<td>0.94</td>
</tr>
<tr>
<td>Dyslipidemia, n (%)</td>
<td>10 (21)</td>
<td>1 (8)</td>
<td>0.30</td>
</tr>
<tr>
<td>Peripheral vascular disease, n (%)</td>
<td>6 (13)</td>
<td>0 (0)</td>
<td>0.19</td>
</tr>
<tr>
<td>Cardiopathy, n (%)</td>
<td>9 (19)</td>
<td>2 (17)</td>
<td>0.84</td>
</tr>
<tr>
<td>Acute kidney disease, n (%)</td>
<td>3 (7)</td>
<td>1 (8)</td>
<td>0.81</td>
</tr>
<tr>
<td>Smoker, n (%)</td>
<td>8 (17)</td>
<td>1 (8)</td>
<td>0.46</td>
</tr>
<tr>
<td>Chronic obstructive pulmonary disease, n (%)</td>
<td>3 (7)</td>
<td>1 (8)</td>
<td>0.81</td>
</tr>
<tr>
<td>Appendectomy, n (%)</td>
<td>12 (26)</td>
<td>4 (33)</td>
<td>0.59</td>
</tr>
<tr>
<td>Cholecystectomy, n (%)</td>
<td>8 (17)</td>
<td>2 (17)</td>
<td>0.98</td>
</tr>
<tr>
<td>Peritonitis, n (%)</td>
<td>2 (4)</td>
<td>1 (8)</td>
<td>0.57</td>
</tr>
<tr>
<td>Previous history of SBO, n (%)</td>
<td>5 (11)</td>
<td>0 (0)</td>
<td>0.24</td>
</tr>
<tr>
<td>Haemoglobin, g/dL ± SD</td>
<td>14 ± 1.8</td>
<td>14 ± 2.7</td>
<td>0.96</td>
</tr>
<tr>
<td>Leukocytes, x 10⁹/mm³ ± SD</td>
<td>12 ± 5.1</td>
<td>11 ± 6.9</td>
<td>0.81</td>
</tr>
<tr>
<td>Platelets, x10⁹/mm³ ± SD</td>
<td>310 ± 100</td>
<td>299 ± 133</td>
<td>0.81</td>
</tr>
<tr>
<td>C reactive protein, mg/L ± SD</td>
<td>31 ± 40</td>
<td>54 ± 72</td>
<td>0.31</td>
</tr>
<tr>
<td>ALAT, U/L ± SD</td>
<td>31 ± 22</td>
<td>48 ± 33</td>
<td>0.15</td>
</tr>
<tr>
<td>ASAT, U/L ± SD</td>
<td>29 ± 19</td>
<td>34 ± 25</td>
<td>0.59</td>
</tr>
<tr>
<td>Urea, mmol/L ± SD</td>
<td>8 ± 6</td>
<td>14 ± 7</td>
<td>0.49</td>
</tr>
<tr>
<td>Lactates, mmol/L ± SD</td>
<td>2.1 ± 1</td>
<td>2.9 ± 1.6</td>
<td>0.26</td>
</tr>
<tr>
<td>Amylase, U/L ± SD</td>
<td>73 ± 55</td>
<td>75 ± 33</td>
<td>0.91</td>
</tr>
<tr>
<td>Lipase, U/L ± SD</td>
<td>48 ± 50</td>
<td>60 ± 28</td>
<td>0.39</td>
</tr>
<tr>
<td>Sodium, mmol/L ± SD</td>
<td>138 ± 3</td>
<td>134 ± 7</td>
<td>0.09</td>
</tr>
<tr>
<td>Potassium, mmol/L ± SD</td>
<td>4 ± 1</td>
<td>4 ± 1</td>
<td>0.28</td>
</tr>
<tr>
<td>Chloride, mmol/L ± SD</td>
<td>101 ± 5</td>
<td>95 ± 10</td>
<td>0.003</td>
</tr>
<tr>
<td>Calcium, mmol/L ± SD</td>
<td>2.21 ± 0.2</td>
<td>2.20 ± 0.2</td>
<td>0.84</td>
</tr>
<tr>
<td>Phosphorus, mmol/L ± SD</td>
<td>1.06 ± 0.3</td>
<td>1.19 ± 0.34</td>
<td>0.25</td>
</tr>
<tr>
<td>Procalcitonin, ng/mL ± SD</td>
<td>0.16 ± 0.14</td>
<td>1.23 ± 2.3</td>
<td>0.002</td>
</tr>
</tbody>
</table>

24 hours after the admission: Using a ROC curve, the AUROC for PCT 24 hours after the admission and SM was 0.859 [95% CI, 0.692–1] (Figure 7). A PCT threshold >0.255 ng/mL for predicting SM yielded a sensitivity of 83.3%, a specificity of 73%, a PPV of 42% and an NPV of 95%.
Predictive time points of PCT to discriminate between SM without peroperative ischemia and SM with peroperative ischemia

Using a ROC curve, the AUROCC for PCT and SM with peroperative ischemia was 0.861 [95% CI, 0.641 – 1] (Figure 8). A PCT threshold >0.53 ng/mL for predicting SM yielded a sensitivity of 80%, a specificity of 84.8%, a PPV of 40% and an NPV of 90.7%.

Discussion

Our study was designed to complete our previous data related on the management of SBO and the potential use of serum PCT as a biomarker. The interest of PCT kinetics’ study has been to validate our previous data in an independent cohort. Indeed, the present population is similar in terms of demographic data as those we previously analyzed [11]. Even if the inclusion period and duration are not similar (34 months vs. 10 months) there is no imbalance between the patients’ characteristics (same age, same comorbidities). As we have already reported, at admission the threshold used to discriminate between conservative management’s success and failure is around 0.17 ng/mL meaning normal PCT levels. This concordance between the results of these two studies could be explained as follow: first the laboratory that made the assessment has already been involved in the previous study (same material, same technicians, and same techniques); secondly as Steinbach et al. have reported in their article [12] the intra-assay and inter-assay coefficients of variation for the Kryptor T.R.A.C.E® method were below 5% and below 10%, respectively. The pertinence of this method (quick results, repeatable data) now used daily in the clinical practice and emergency situations, made it sure and reliable.

The complexity for the surgeon to operate the patient is laid on the
The ratio after 36 hours could be due to many factors: the inflammatory process, the metabolic changes, and the interactions between different physiological systems. It seems that this ion enhances the production by a factor of 2. Part of chloride in the PCT's secretion is reflected by the ratio PCT SM/PCT CM. The more pronounced this phenomenon, the more chloride's osmolarity will be modified.

Symptoms longer than the others patients (nausea and vomiting; pain) are more common in patients requiring surgical management. Indeed, it is possible that peripheral cytokines such as IL6, IL8 and TNFα are released, inducing a systemic inflammatory response. Nevertheless, these signs are imperfect because in our study none of the signs was predictive of the surgery confirming the results of Evennett et al., Gearhart et al. and Delaney et al. [13-15] showing that these "old markers" are not accurate enough. At admission, with a PCT higher than 0.53 ng/mL, the surgeon could decide to operate the patient whereas a PCT between 0.17 and 0.53 ng/mL involves an aggressive monitoring. Twenty four hours after the admission a PCT higher than 0.27 ng/mL impose a surgery.

The choice of serum PCT as a biomarker for SBO's management lies on its half life time (24 hours) [16]. As reported by numerous previous works [17-21], in case of inflammatory state (for example SBO, cytokines such as IL6, IL8 and TNFα are released inducing a secretion of PCT but in case of conservative management's success the inflammatory state decreases so quickly that PCT secretion does. Complementary to these works, the data from Nagata et al. [22]; Markogiannakis et al. [23]; Karabulut et al. [24] and Ayten et al. [25] had already presented PCT as a possible biomarker. The strong correlation between PCT and the conservative management failure rate accentuates this proposition.

The difference between the two groups in the study related on chloride has previously been reported [11]. The ion seems to be involved in the SBO’s management and the disease’s severity. Indeed, it is possible that patients requiring surgical management have presented SBO’s symptoms longer than the others patients (nausea and vomiting; pain more pronounced). In that case chloride’s osmolarity will be modified enhancing PCT’s secretion to regulate the calcium’s metabolism. The part of chloride in the PCT’s secretion is reflected by the ratio PCT SM/PCT CM. It seems that this ion enhances the production by a factor 2 or 4 (if we consider that no other variable is involved). The increase in the ratio after 36 hours could be due to many factors: the inflammatory state due to the surgical procedure (because in our study the surgery was mainly performed after the 36th hours) [26], the possible infectious phenomenon engendered by the surgery etc.

To validate the use of PCT in this situation, we enrolled patients with acute kidney disease to ensure that kidney disease don’t have an impact on this biomarker’s kinetics. Indeed patients admitted with SBO are frequently aged and could present an altered renal function. Nevertheless, in our population this biological function does not have any effect on PCT’s levels. This observation confirms the data of Meissner et al. [27] who reported no difference of PCT in case of renal dysfunction.

This study validates the role of serum PCT as a biomarker of bowel ischemia. Indeed, in our previous study, we showed that PCT was a marker of ischemia with a threshold of 0.57 ng/mL. The confirmation of these data in an independent cohort let us think that PCT could be proposed as a systemic marker of bowel ischemia reflecting the inflammatory state engendering by the obstruction.

**Conclusion**

Our work confirms the previous data on the use of PCT as a biomarker in SBO’s management. The more predictive time points are at admission, 12 hours and 18 hours after admission to identify patients requiring surgery.

*This article has been presented as a poster at the UEG Week 2013 in Berlin*

**References**


