Diagnostic performance of S100B protein serum measurement in detecting intracranial injury in children with mild head trauma

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ABSTRACT

Objective To assess the accuracy of S100B serum level to detect intracranial injury in children with mild traumatic brain injury.

Methods A multicenter prospective cohort study was carried out in the pediatric emergency departments of three tertiary hospitals in Switzerland between January 2009 and December 2011. Participants included children aged <16 years with a mild traumatic brain injury (GCS ≥13) for whom a head CT was requested by the attending physician. Venous blood was obtained within 6 h of the trauma in all children for S100B measurement before a head CT was performed. As the S100B value was not available during the acute care period, the patient’s management was not altered. The main measures were protein S100B value and the CT result.

Results 20/73 (27.4%) included children had an intracranial injury detected on CT. S100B receiver operating characteristics area under the curve was 0.73 (95% CI 0.60 to 0.86). With a 0.14 μg/L cut-off point, S100B reached an excellent sensitivity of 95% (95% CI 77% to 100%) and 100% (95% CI 81% to 100%) in all children and in children aged >2 years, respectively. The specificity, however, was 34% (95% CI 27% to 36%) and 37% (95% CI 30% to 37%), respectively.

Conclusions S100B has an excellent sensitivity but poor specificity. It is therefore an accurate tool to help rule out an intracranial injury but cannot be used as the sole marker owing to its specificity. Used with clinical decision rules, S100B may help to reduce the number of unnecessary CT scans.

INTRODUCTION

Traumatic brain injury (TBI) is a very frequent cause of presentation to the emergency department. This condition accounts for more than 500 000 visits a year in children.1 The vast majority of children with mild TBI, defined as GCS 13–15,2 have no intracranial injury (ICI);3 a lesion is seen in 3–7% of these children undergoing CT3–6 and only 0.1–0.6% need a neurosurgical intervention.3–3

Consequences of a missed ICI may be devastating and its symptoms or signs, such as vomiting, headache or amnesia, are often misleading. CT is therefore performed in almost 50% of children with mild TBI7 as this is the only diagnostic tool to detect an ICI in the emergency setting.

Key messages

What is already known about this subject?

▸ Studies in adults show that serum S100B is a possible adjunct to clinical decision rules to detect intracranial injury, and the American College of Emergency Physicians suggests that S100B measurement could reduce the number of unnecessary CTs.

▸ Research in children on this biomarker is limited and it shows interesting but conflicting results.

What does this study add?

▸ In this multicentre, prospective cohort study in patients aged <16 years, S100B was found to have excellent sensitivity, making it an accurate tool to rule out intracranial injuries. Its poor specificity is a limiting factor that precludes its use as the sole marker.

▸ Used with clinical decision rules, S100B may help to reduce the number of unnecessary CTs in children.
Many adult studies have shown that serum S100B is a possible adjunct to clinical decision rules to detect ICI,17–21 with the American College of Emergency Physicians suggesting that S100B measurement could reduce the number of unnecessary CT scans by 30%.22

A few studies have evaluated S100B as a tool to reduce the number of CT scans in children, with interesting but conflicting results.23–27

The objective of our study was to assess the accuracy of S100B serum level to detect ICI in children with mild TBI.

METHODS

Study design

This was a multicentre prospective cohort study. The study received institutional review board approval of each participating hospital and was conducted in accordance with Good Clinical Practice guidelines and provisions of the Declaration of Helsinki.

Settings and selection of participants

The study was conducted in the paediatric emergency departments of three tertiary hospitals in Switzerland on a consecutive sample of patients between January 2009 and December 2011. These three centres see 33,000, 24,000 and 12,000 patients a year. We included all children aged <16 years with a mild TBI (GCS ≥13) for whom a head CT was requested by the attending physician. We excluded those children who arrived at the hospital more than 6 h after the trauma, children with Down syndrome (since in these patients S100B is overexpressed28) or patients with a history of convulsion in the past 7 days.

Study protocol

The attending paediatric emergency physicians obtained consent from the parents of children meeting the inclusion criteria for participation in the study. After consent was obtained, venous blood was drawn and centrifuged. The serum was then frozen (−20°C) and sent in batches to the division of clinical chemistry and biochemistry of the University Children’s Hospital Zurich for S100B measurement. As the S100B value was not available during the acute care period, the patient’s management was not altered.

All head CT scans were reviewed for traumatic injuries by one paediatric radiologist (CJK), who was blinded to the patient’s clinical signs and the S100B value.

S100B measurement

All S100B (αα and αβ dimers) measurements were done on an Elecsys 2010 using the commercial S100 assay from Roche (Roche Diagnostics, Rotkreuz, Switzerland), with an interassay coefficient of variation <2.8% according to the manufacturer.

Outcome measures

Primary outcome was evaluation of the diagnostic value of S100B in detecting intracranial injuries in children aged <16 years with mild head trauma. Secondary outcomes were evaluation of the same parameters excluding children aged <2 years (age group where normal S100B values vary physiologically with age) and comparison of signs and symptoms between the groups with and without ICI (all children).

Definitions

Mild TBI: acute head trauma with a GCS 13–15, with confusion or loss of consciousness (<30 min) or amnesia or transient neurological abnormality.2

ICI: any collection of blood within the cranial vault or cerebral oedema.

Data analysis and sample size

A χ² test was used to describe categorical data. A t test was used to compare continuous variables as normality was observed in our data. The 95% CIs of the results are reported. A level of p<0.05 was considered significant. Statistical analyses were performed using PASW V18 (SPSS Inc, Chicago, USA).

RESULTS

Between January 2009 and December 2011, 2595 children were admitted for a mild TBI and 342 (13.2%) had a head CT scan. A total of 80 children with mild TBI meeting the inclusion criteria were enrolled (see figure 1). Seven patients were excluded (no S100B measurement: three patients; venous puncture >6 h after trauma: one patient; no CT performed: three patients). Of the 73 included children, 20 (27.4%) had an ICI detected on CT. The lesion was an epidural haematoma in nine children, a subarachnoid haemorrhage in four, an epidural haematoma and subarachnoid haemorrhage in three. The remaining four children had respectively a subdural haematoma, an epidural and subdural haematoma, a subdural haematoma and subarachnoid haemorrhage and haemorrhagic parenchymal contusion. No surgical intervention was required. There was no statistically significant difference between the patients with or without ICI in age, gender and time between trauma and venous puncture for S100B measurement (table 1).

Clinical features that might influence a physician’s decision to obtain a CT scan are shown and compared between groups in table 1. The only statistically significant difference between groups was the S100B value.

The S100B receiver operating characteristics (ROC) curve is shown in figure 2. The area under the curve was 0.73 (95% CI 0.60 to 0.86). After excluding children aged <2 years, the area under the curve was 0.77 (95% CI 0.65 to 0.89) for the remaining 64 children.

Using the best cut-off value derived from the ROC analysis (0.14 μg/L), S100B reached a sensitivity of 95% (19/20) (95% CI 77% to 100%) and 100% (18/18) (95% CI 81% to 100%) in all children and in children aged >2 years, respectively. With this cut-off point, one (1.4%) child (aged 17 months) with a 1.5 cm temporoparietal epidural haematoma would have been missed. The corresponding specificity and the derived positive and negative likelihood ratio for all children and children aged >2 years are shown in table 2. This cut-off point has been chosen to take into account the best sensitivity/specificity ratio. If the aim was 100% sensitivity, specificity would fall to 4% (with a 0.07 μg/L cut-off value).

DISCUSSION

This study shows that S100B protein has good diagnostic properties with an area under the curve of 0.73 and, with 0.14 μg/L cut-off value, a very good sensitivity but poor specificity, in children aged <16 years with mild TBI (GCS 13–15).

We evaluated the same parameters after exclusion of children aged <2 years. As expected these are slightly better: the area under the curve rises to 0.77 and sensitivity reaches 100% with similar specificity. The reason for this is probably because the S100B value is age-dependent with physiologically higher levels in the youngest subjects.29–31 Reference value studies mainly use two age groups: under or over 2–3 years of age.29–31

Taking S100B as the only identifier of ICI, 18 (24.7%) head CT scans would have been avoided. However, one (1.4%) child...
with a 1.5 cm epidural haematoma would have been missed. Thus, the decision to perform a CT scan cannot rely on the biomarker alone. In order to reduce unnecessary CT scans and radiation-induced malignancies, several decision rules based on clinical signs have been described. The strongest validated rule to date published by Kuppermann et al suggests obtaining a CT scan in all children with GCS <15 or other signs of altered mental status or signs of basilar skull fracture, but leaves the decision to the physician if the child has any history of loss of consciousness, vomiting, a severe mechanism of injury or severe headache. According to these authors, the decision should be based, among other factors, on the physician’s experience. In our opinion, this is the situation in which S100B could help to avoid unnecessary CT.

Most evaluations of the diagnostic accuracy of S100B come from reports on adults. Unden and Romner included 12 studies with 2466 patients in a systematic review and meta-analysis. They found sensitivity and specificity results similar to ours (97% and 40%, respectively). The largest prospective cohort comparing S100B values with CT results in 1559 adults showed the same sensitivity (98.6%) but worse specificity (12%); the area under the curve was 0.76.

Our results are also comparable to those of the few paediatric studies published. Babcock et al included children aged 0–18 years with TBI of all grades; the area under the curve was 0.71. However, to achieve optimal sensitivity and specificity, the cut-off value was much lower than ours (0.006 μg/L). The reason might be a different definition of ICI, comprising skull fractures. Castellani et al selected 109 children with only mild TBI (GCS 13–15). With a cut-off value comparable to ours (0.16 μg/L) and they again found similar sensitivity (100%) and specificity (42%); the area under the curve was 0.68.

In contrast, Hallen et al published much higher S100B diagnostic properties. The area under the curve was 0.93, sensitivity 100% and specificity 88%. The difference between their study and other studies was the definition of a negative outcome. Children were considered to be without ICI if CT was normal or the course after observation was satisfactory, with only 10% of patients having a CT scan. This is an interesting point and possibly a limitation of our study. We considered positive any intracranial abnormality, even, probably, lesions without consequences, as in previous studies. It is, however, difficult to recognise those with potential for deterioration. Because a S100B value is undetectable after 6 h from injury in 36% of patients with initially detectable levels, we decided to include only children within 6 h. It is not clear whether earlier blood sampling would have altered the diagnostic properties. Bouvier et al found a 100% sensitivity and 33% specificity when including children within 3 h of the TBI. However, others had the same sensitivity with better specificity when including children within 6 h.

S100B protein is expressed in astrocytes and Schwann cells, but also in chondrocytes and adipocytes. This raised the

<table>
<thead>
<tr>
<th>Table 1</th>
<th>Comparison of demographic and clinical features between the groups with and without ICI based on CT</th>
</tr>
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<tbody>
<tr>
<td>Group</td>
<td>Without ICI (n=53)</td>
</tr>
<tr>
<td>Age (months), mean (SD)</td>
<td>94.0 (56.5)</td>
</tr>
<tr>
<td>Male, n (%)</td>
<td>35 (66.0)</td>
</tr>
<tr>
<td>Time trauma to venous puncture (min), mean (SD)</td>
<td>163.8 (84.7)</td>
</tr>
<tr>
<td>GCS &lt;15, n (%)</td>
<td>19 (35.8)</td>
</tr>
<tr>
<td>Vomiting, n (%)</td>
<td>25 (47.2)</td>
</tr>
<tr>
<td>Headache, n (%)</td>
<td>28 (52.8)</td>
</tr>
<tr>
<td>Amnesia, n (%)</td>
<td>17 (32.1)</td>
</tr>
<tr>
<td>Seizure, n (%)</td>
<td>4 (7.5)</td>
</tr>
<tr>
<td>S100B value (μg/L), mean (SD)</td>
<td>0.35 (0.45)</td>
</tr>
<tr>
<td>(range)</td>
<td>(2.79)</td>
</tr>
</tbody>
</table>

ICI, intracranial injury.
question as to whether other lesions, such as long-bone fractures, could raise the serum S100B level, with conflicting results in the paediatric literature. Unlike Berger et al.,34 who found no influence, Bechtel et al.24 showed that children with ICI and long-bone fractures had significantly higher levels of S100B than those without a fracture, possibly affecting the specificity of the biomarker. This aspect could not be evaluated in our study because only one child had long-bone fracture (with a S100B of 1.62 μg/L).

Another interesting finding is that we found no statistically significant difference between the groups in the presence of ICI-related worrying symptoms, whereas S100B levels were higher in the group with ICI. That point was also evaluated by Babcock et al.,23 but with opposite results. Their two groups significantly differed in their responses concerning nausea or vomiting, headache and amnesia. The difference might be explained by their study population that included TBI of all severities. Also, there may be a selection bias for our patients, possibly identified as needing a CT scan based on these symptoms. Indeed, ICI was found in 27.4% of our population, compared with 17.4% in the study of Babcock et al.23

Limitations
Our study has some limitations. First, is the small sample size. Because of this, we could not draw conclusions in the subgroup aged <2 years and we had no surgical intervention. It would have been interesting to compare these children with those who received conservative treatment. However, despite our small sample size, we obtained results similar to those of much larger adult20 21 and paediatric studies.23–26 32 Also, the study was conducted in three tertiary hospitals, but the inclusion period and contribution were not equal, with most of the study patients coming from the two biggest centres. As stated before, our study population is limited to children undergoing head CT, with the risk of considering positive any intracranial abnormality, even, possibly, lesions without consequences. However, the benefit is to have objective outcomes, as it is difficult to identify patients at risk of deterioration.

In conclusion, the biomarker S100B is a valuable tool to help the physician decide whether head CT is indicated for children aged <16 years with mild head trauma. Its excellent sensitivity indicates that it could be an accurate tool to ‘rule out’ an ICI. However, its poor specificity is a limiting factor that precludes...
its use as a sole marker. Used with clinical decision rules, S100B may help to reduce the number of unnecessary CT scans.

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Contributors IBH, GS and SM conceived the study, designed the trial and obtained research funding. IBH and GS supervised the conduct of the trial and the data collection. LL, DK-L, GLS and SA undertook recruitment of participating centres and patients and managed the data, including quality control. CJK interpreted all study CT scans. MH supervised the S100B laboratory analysis and its data. SM and LL provided statistical advice on study design and analysed the data. SM drafted the manuscript and all authors contributed substantially to its revision.

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Competing interests None declared.

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Correction notice Since this paper was first published online figure 2 has been updated so that it displays two different ROC curves.

REFERENCES