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Observational Study

Delirium, insulin-like growth factor I, growth hormone in older inpatients

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Abstract

BACKGROUND

Delirium is a common disorder in elderly medical inpatients with serious adverse outcomes and is characterized by sudden onset, disturbance in attention, awareness, consciousness and cognition, and often with behavioural disturbances. Central to understanding delirium, is understanding mechanisms by which body and brain wellbeing are linked and in particular how brain responses to bodily homeostatic stress is mediated. A number of studies have investigated the relationship between insulin-like growth factor I (IGF-I) and delirium in medically ill hospitalised patients with conflicting results. However, none have investigated growth hormone (GH) which is related to IGF-I via negative feedback.

AIM
Delirium is a syndrome which often presented with sudden onset, and disorders of the cognitive, consciousness, motor, affective, and perceptual domains. Those disturbances often are fluctuated\(^6\). It also manifested with different motor subtypes like hyperactive, hypoactive or mixed states\(^4\). Delirium is a common disorder in elderly medical inpatients. In Irish hospitals the point prevalence of delirium is estimated around 20\%\(^{\ref{5}}\). Higher rates are reported among patients in palliative and intensive care settings\(^{\ref{4,5}}\). Delirium has severe bad consequences like, delays in hospital discharges increased hospital budgets\(^8\), higher rates of institutionalisation\(^8\), and perhaps increased mortality\(^7\). The old-fashioned
concept of delirium as a brief, transient, and highly reversible condition is no longer supported by longitudinal studies. Accumulating evidence support that delirium is associated with persistent cognitive and functional problems and delirium may be an accelerating and possibly causal factor in the development of dementia. Notably, these adverse outcomes are independent of the severity of physical illness that cause delirium. Despite the understanding that delirium is caused by physical illness, it is not clear yet how these physical causes without direct connection with brain, can produce such a consistent complex neuropsychiatric picture as found in delirium. Thus, the pathophysiology of delirium remains unclear and many pathophysiological mechanisms have been hypothesised and suggested but none yet proved.

Delirium often occurs in the context of infectious illness and the external administration of cytokines in a number of medical illness for therapeutic reasons, can results to delirium. Thus, in an effort to clarify the pathophysiology of delirium a number of studies have investigated cytokine levels, but the results are conflicting and inconclusive. However, from our previous work we found that low circulating levels of neuroprotective factors of IGF-I and interleukin-1 receptor antagonist (IL-1RA) were connected with delirium. Although there are few studies with firm conclusions about the role of cytokines in delirium, a hypothesis in which we work is that cerebral deficits may be the reason for the occurrence of delirium. This has led our research group to work on the general hypothesis that delirium is associated with already existed deficits in the brain regarding the immunoreactivity and the readiness to respond to the external “insult” (e.g., physical illness). Low levels of neuroprotective factors may possibly explain the onset of delirium rather than the actual trigger or “insult” factor. This can explain the observations that the severity of physical illness is not a risk factor for delirium at least in older populations. One of those neuroprotective factors which has been investigated in delirium is the IGF-I.

IGF-I is regulate the body growth and metabolism but involves also in different brain functions. Circulating IGF-I is produced mainly from the liver but it can be produced by any cell type. The receptors of IGF-I are on almost all different cells including brain and have neuroprotective effects. In addition IGF-I receptors play significant role in the integrity and regulation of blood-brain barrier, they have high expression in the cells that constitute it and they facilitate the access of serum IGF-I to entry into the brain.

In the Central Nervous System the IGF-I is produced by neurons and glial cells and plays an extensive role in the development, plasticity and survival of neurons. IGF-I involves in the production of neurotransmitters, blocks apoptosis in damaged neurons, and thus has effects on cognition and cognitive decline during ageing or in other neurocognitive disorders like dementia and delirium. It has been reported that cerebrospinal levels of somatostatin (Growth hormone Inhibiting hormone) are significantly lower during delirium but and later at follow-up. This does not indicate a directly link between delirium and GH, but indicates a disturbed GH/IGF-I axis in delirium. In the same line, in patients admitted to the intensive care unit it was found that low levels of IGF-I were correlated with high levels of GH. In addition, a study reported that administration of human GH in aged women following hip operation had increased IGF-I levels. Therefore, until now we do not know the way of the interaction of GH and IGF-I during delirium. In our previous review we have identified a lack of research work in relation to GH in studies which investigated IGF-I and delirium.

Nevertheless, a number of studies have investigated the relationship of IGF-I with delirium in medically ill hospitalised patients as well as in patients undergoing surgery with conflicting results. A resent meta-analysis showed that there are indications of an association of IGF-1 and delirium but the authors also calling for further research into this area.

Given the previous studies and the new updates we carried out a new study in older medically ill hospitalised patients with the aim to find out the relationship of the circulating levels of IGF-I and GH to the delirium (in both prevalent and incident).
MATERIALS AND METHODS

The present study was designed as a pragmatic prospective, longitudinal study. The study was carried out in a University Hospital in Sligo in the North-West of Ireland. The inclusion criteria were: (1) Consecutive admitted patients in the elderly medical wards; and (2) To be 70 years old and above. Exclusion criteria were: (1) Patients who were readmitted and had already participate in the study; (2) Patients intubated or with aphasia; (3) Patients in a terminal stage of illness; and (4) Patients unable to speak English. A time frame of 72 h since admission was in place for the assessment of the eligibility for recruitment and the recruitment of participants.

Procedure
Those patients who fulfilled the inclusion criteria and consented had an assessment at first day. Then seven more assessments were followed in a regular space of 3 ± 1 d if they were still hospitalised and alive. The maximum number of assessment was eight. Non-fasting blood was withdrawn the same days of the assessments. Bloods were centrifuged within ten minutes and then stored at -70 °C until analysis. Levels of IGF-I and GH were estimated with the ELISA method. Levels of IGF-I are measured in ng/ml and levels of GH in pg/mL.

Assessments and scales
Demographic data were collected from the computer of the hospital database. In addition at each time the following measurements/scales were administered.

Cognitive scale
The Montreal Cognitive Assessment (MoCA) have been used for assessment of cognition. The maximum score in MoCA is 30 which indicates an intact cognition. In participants who were unable to complete all the sections due to a physical disability (e.g., visual impairment) MoCA results were standardized to give a maximum score of 30. To complete the MoCA it takes about 12-15 min.

Assessment/scale for delirium
The presence/absence of delirium was assessed with the CAM scale/algorithms. The CAM is based in DSM-IIIIR criteria for delirium. It asses four “cardinal” criteria for delirium.

Physical illness
To assess the severity of the underling physical illness the Acute Physiology and Chronic Health Evaluation II (APACHE-II) (Acute Physiology and Chronic Health Evaluation II) and the APS subscale (Acute Physiology Score) were used. Higher scores in both scales indicate more illness severity.

Diagnosis of previous history of dementia
Pre-existing dementia was assessed with two ways. First if it was documented clearly according to DSM-IV diagnostic criteria or if not the Short Informant Questionnaire of Cognitive Decline was used by interviewing the nearest relative. The cut-off point of ≥ 3.5 was used to define pre-existing dementia.

Ethical considerations
Informed consent was in writing using an earlier described method. A separate consent in writing was asked for phlebotomy. The Sligo University Hospital Research Ethics Committee has graded Ethical approval for the project.

Statistical analysis
SPSS v23 was used for the analysis of the data. Continuous variables were presented as mean ± SD and categorical as counts and percentages. The Generalized Estimating Equations (GEE) method was used to analyse the effects of independent variables on delirium. GEE adjusts for correlations due to repeated assessments of each participant. Because the dependent variable (delirium/no delirium) was binary the binominal distribution was used. To evaluate the fit of the model the Corrected Quasi Likelihood under Independence Model Criterion (QICC) value was used, (lower value – better fit). Because there were many missing values in the last 3 assessments (drop-outs) only the first 5 will be entered to the model.
RESULTS

Description of the sample
A total of 198 participants were analysed. The mean age of the participants was 80.63 ± 6.81; range 70-97. Of these 106 (53.5%) were males. Previous history of dementia was found in eighty six (43.4%). The characteristics of the two groups (delirium/ no delirium) at each of the five assessments, including means and standard deviations of the scales MoCA, and APACHE II scores, and IGF-I and GH levels at each assessment are shown in Table 1. Figure 1 shows the mean levels of IGF-I and GH across the assessments for those with and without delirium.

Evaluation of missing data
This was done by using the Little’s MCAR test. The results of the test was not significant (MCAR, $\chi^2 = 12.24, df = 9, P = 0.20$) which indicates that the missing values were missing completely at random.

Longitudinal analysis: GEE model
Here we examined the effects of the independent variables age, gender, previous history of dementia (binary yes/no), APACHE II, MoCA scores, and the levels of IGF-I and GH on the dependent variable delirium/no delirium (binary). The most parsimonious model (lowest QICC value) is shown in Table 2.

The results from the Table 2 shows that those with any delirium during the hospitalisation had significantly lower scores in the MoCA scale, lower levels of IGF-I and higher levels of GH compared to those without delirium. None of the other examined variables (age, gender, previous history of dementia or severity of physical illness (APACHE II) had any significant effect in the presence or absence of delirium as it was defined with CAM.

DISCUSSION

First of all, the results show that deficits in cognition as measured with the MoCA, are a significant independent predictor for the occurrence of any delirium (prevalent, incident, or fluctuating). This result is constantly found in all the studies which investigate delirium because disturbance in cognition is a central feature of delirium. Therefore this is a result which was expected. In addition the severity of the physical illness (as measured with APACHE II), previous history of dementia, and age did not have any effect on delirium. Severity of physical illness again is an expected finding since in our previous studies we did not find any effect and thus we generate the hypothesis that it is not the severity of insult that is important for causing the delirium but the reduced neuroprotection of the brain. Besides the lack of effect of age and previous history of dementia is easily explained by the more powerful predictor, scores in the MoCA.

Regarding the IGF-I, the results of the present study is in accordance with our previous study in which we use similar methodology and longitudinal design but in a different population, different hospital and in a different country and also confirmed that low levels of circulating IGF-I are significantly linked with any delirium (incident or prevalence). Similar results have been reported form other research groups, but not from all. However a resent meta-analysis showed that lower levels of circulating IGF-I are associated with higher rates of delirium among older patients. There are many reasons for those discrepancies among the studies, but not from all. First of all different populations were studied. Some studies include populations with pre-existing dementia (e.g., the present study) while others excluded them. A second reason perhaps is the setting where the study is conducted and the sample. Some of the studies were conducted in medical wards where the sample include populations with mainly medical illness while others in surgical wards in patients before and after surgery. Perhaps surgery is another stressor and perhaps pathophysiology which leads to delirium in those patients is different despite the end product being the same. In addition those studies in surgery wards have examined patients before and immediately after the surgery for delirium. However, it has been suggested that perhaps different mechanisms are underline the delirium that developed in the first 24 hours after surgery and in the delirium that developed the next one to three days after surgery. Finally one important reason which can explain those discrepancies is the different scales/measurements/criteria that have been applied to define delirium. It has been shown that applying different criteria for delirium is influence significantly
Table 1 Cases of delirium and no delirium (according to confusion assessment method) with the scores (mean ± SD) of the Montreal cognitive assessment, acute physiology and chronic health evaluation II and growth hormone, insulin-like growth factor I levels at each assessment point

<table>
<thead>
<tr>
<th>Assessment</th>
<th>CAM (n)</th>
<th>MoCA</th>
<th>APACHE II</th>
<th>IGF-I (ng/mL)</th>
<th>GH (pg/mL)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>No delirium (173)</td>
<td>mean ± SD</td>
<td>11.38 ± 7.90</td>
<td>8.59 ± 3.68</td>
<td>61.32 ± 22.95</td>
</tr>
<tr>
<td></td>
<td>Valid, n</td>
<td>168</td>
<td>173</td>
<td>68</td>
<td>68</td>
</tr>
<tr>
<td>Delirium (24)</td>
<td>mean ± SD</td>
<td>3.78 ± 2.76</td>
<td>9.00 ± 3.86</td>
<td>45.88 ± 14.66</td>
<td>693.89 ± 462.15</td>
</tr>
<tr>
<td></td>
<td>Valid, n</td>
<td>23</td>
<td>24</td>
<td>8</td>
<td>8</td>
</tr>
<tr>
<td>2</td>
<td>No delirium (141)</td>
<td>mean ± SD</td>
<td>10.42 ± 8.23</td>
<td>8.69 ± 3.69</td>
<td>53.93 ± 18.57</td>
</tr>
<tr>
<td></td>
<td>Valid, n</td>
<td>134</td>
<td>140</td>
<td>49</td>
<td>49</td>
</tr>
<tr>
<td>Delirium (11)</td>
<td>mean ± SD</td>
<td>3.67 ± 3.43</td>
<td>11.00 ± 5.20</td>
<td>41.78 ± 13.98</td>
<td>652.14 ± 346.98</td>
</tr>
<tr>
<td></td>
<td>Valid, n</td>
<td>9</td>
<td>11</td>
<td>5</td>
<td>5</td>
</tr>
<tr>
<td>3</td>
<td>No delirium (90)</td>
<td>mean ± SD</td>
<td>9.27 ± 7.58</td>
<td>8.77 ± 3.67</td>
<td>60.66 ± 18.88</td>
</tr>
<tr>
<td></td>
<td>Valid, n</td>
<td>84</td>
<td>88</td>
<td>30</td>
<td>30</td>
</tr>
<tr>
<td>Delirium (13)</td>
<td>mean ± SD</td>
<td>3.90 ± 2.51</td>
<td>10.38 ± 3.69</td>
<td>58.29 ± 14.60</td>
<td>487.25 ± 507.84</td>
</tr>
<tr>
<td></td>
<td>Valid, n</td>
<td>10</td>
<td>13</td>
<td>5</td>
<td>5</td>
</tr>
<tr>
<td>4</td>
<td>No delirium (58)</td>
<td>mean ± SD</td>
<td>10.63 ± 7.44</td>
<td>8.79 ± 3.55</td>
<td>61.97 ± 22.70</td>
</tr>
<tr>
<td></td>
<td>Valid, n</td>
<td>56</td>
<td>58</td>
<td>18</td>
<td>18</td>
</tr>
<tr>
<td>Delirium (8)</td>
<td>mean ± SD</td>
<td>6.00 ± 1.83</td>
<td>9.00 ± 2.67</td>
<td>41.43 ± 8.13</td>
<td>577.94 ± 200.90</td>
</tr>
<tr>
<td></td>
<td>Valid, n</td>
<td>7</td>
<td>8</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>5</td>
<td>No delirium (46)</td>
<td>mean ± SD</td>
<td>8.70 ± 7.20</td>
<td>8.80 ± 3.28</td>
<td>53.70 ± 14.37</td>
</tr>
<tr>
<td></td>
<td>Valid, n</td>
<td>43</td>
<td>46</td>
<td>11</td>
<td>11</td>
</tr>
<tr>
<td>Delirium (6)</td>
<td>mean ± SD</td>
<td>2.33 ± 1.03</td>
<td>9.50 ± 3.33</td>
<td>42.62 ± 2.42</td>
<td>924.34 ± 677.88</td>
</tr>
<tr>
<td></td>
<td>Valid, n</td>
<td>6</td>
<td>6</td>
<td>4</td>
<td>4</td>
</tr>
</tbody>
</table>

Table 2 Generalized estimating equations model and parameter estimates of the significant variables on delirium status

<table>
<thead>
<tr>
<th>Parameter estimates</th>
<th>Parameter</th>
<th>B</th>
<th>SE</th>
<th>95%CI</th>
<th>Hypothesis test</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Lower</td>
<td>Upper</td>
</tr>
<tr>
<td>GH (pg/ml)</td>
<td>-0.0011</td>
<td>0.0002</td>
<td>-0.001</td>
<td>0</td>
<td>6.21</td>
</tr>
<tr>
<td>IGF-I (ng/ml)</td>
<td>0.02</td>
<td>0.0074</td>
<td>0.005</td>
<td>0.034</td>
<td>7.196</td>
</tr>
<tr>
<td>MoCA</td>
<td>0.205</td>
<td>0.0586</td>
<td>0.09</td>
<td>0.32</td>
<td>12.231</td>
</tr>
</tbody>
</table>

The sign (-) shows the direction of the relationship. Reference status: No delirium. GH: Growth hormone; IGF-I: Insulin-like growth factor I; MoCA: Montreal cognitive assessment.

the rates and diagnosis of delirium and there is extensive discrepancy between the actual cases defined by each different system\textsuperscript{45-48}.

Furthermore, the present study for first time shows that GH is an important factor in the pathogenesis of delirium. However, because IGF-I and GH are correlated with a negative loop feedback we do not know which of them is more important in the pathogenesis of delirium. From the present study what we can conclude is that the somatotropic axis (IGF-I/GH) is disturbed during the delirium phase compared to those without delirium. Low IGF-I and high GH levels are related to delirium. To the best of our knowledge no previous studies have investigated the IGF-I/GH axis.
Therefore we attempt to explain our hypothesis further.

The secretion of GH is age related and a decline of GH started between the ages of 18 to 25 years. This decrease of the GH concentration in the periphery is accompanied by a gradual decline of IGF-I as well\[27,49\]. In addition, lower levels of GH are correlated with frailty\[27\] and exogenous administration of GH can increase energy, and improve mood, concentration, and memory. Those improvements in cognition may are the results of a direct effect of GH in the brain\[50,51\]. Therefore, perhaps the increased levels of GH reflect a compensatory mechanism in the brain to recover through a direct effect of GH. However, this does not explain the low levels of IGF-I which interestingly have been proven to protect neurons directly\[52\]. Experimental studies in animals have shown that after administration of IGF-I in brain injuries there was an improvement in the outcomes (regarding behaviour and cognition)\[53\]. Therefore a direct effect of IGF-I is more likely than a direct effect of GH. In the same line a second explanation has been suggested that in acute inflammation situations there is a GH resistance because the body prevents growth and energy storage in an attempt to keep the homeostasis. In those situations of acute inflammation the levels of IGF-I reduced, regardless of the increase of GH\[41,55\].

Taken together those suggestions provide a likely explanation that in delirious states GH is increased because of the disturbed IGF-I / GH axis and the lack of inhibitory mechanism of IGF-I. Whatever mechanism is involved we cannot conclude from this study and it is important to notice that we measure levels in the periphery which may or may not reflect the brain levels of IGF-I and GH. However, we can conclude that in delirium the IGF-I/GH axis is disturbed, and that low levels of IGF-I together with high levels of GH and impaired cognition are independently significant predictors of delirium.

Limitations of the study

An obvious and common limitation of those kinds of studies including the present is the small sample size. However, the drop-outs and missing data of the study were completely missing at random so no biases have introduced in the study. A second limitation of the study is the lack of generalizability. The results of this study apply only in medically ill older people and not in other populations. Perhaps similar studies are needed also in other populations like surgical patients. There are ongoing collaborative studies examined the role of serum factors in postoperative delirium. Surgery induced delirium is a good model to separate the predisposal factors (preoperative) from the precipitating factors (post-operative) in the occurrence of delirium. Furthermore, the strengths of his study are the longitudinal design and the statistical analysis accompanied the design. By having this design and analysis we have included all the deliria during hospitalisation (prevalence, incident, fluctuated, and persistent) compared with the non-delirium states (including never delirium during hospitalisation and recovered delirium) across the time in the entire examined population.
Implications of the study
As we noted above, further studies need to be done in different populations before we be asserted about the results. If the results are replicated in further studies this can lead to clinical trials for the treatment and / or prevention of delirium with small doses of IGF-I.

CONCLUSION
In conclusion, this study indicates that during delirium in older medically ill hospitalised patients the IGF-I/GH axis is disturbed but we do not know yet the mechanism behind it. However more studies are needed to confirm or disconfirm the above findings before we move further to clinical trials for treatment or prevention of delirium with small doses of IGF-I.

ARTICLE HIGHLIGHTS

Research background
Delirium is a common disorder in elderly medical inpatients, in surgical wards, and Intensive care units with serious adverse outcomes.

Research motivation
To understand delirium is important to understand the underline mechanisms by which body and brain are linked and how brain responses to bodily homeostatic stress is mediated. We have notice from our previous research work that the severity of physical illness is not a risk factor for delirium at least in older populations and perhaps delirium is associated with deficits in the immunoreactivity of the brain (low cerebral reserve). Low levels of neuroprotective factors may possibly explain the onset of delirium rather than the actual trigger or “insult” factor. A number of studies have investigated the relationship between Insulin-like growth factor I (IGF-I) and delirium with conflicting results. A relevant also factor is the Growth Hormone (GH) which is related to IGF-I via negative feedback. Therefore in the present study we included also the GH.

Research objectives
To investigate the relationship of the occurrence of delirium during hospitalisation (prevalent and incident) with the serum levels of IGF-I and GH.

Research methods
Observational, prospective, longitudinal study of older people who consecutively admitted to medical wards of a general hospital.

Research results
We found that low cognitive function, low levels of IGF-I and high levels of GH were significantly associated with any delirium (prevalence, incident, or fluctuating) during the study period.

Research conclusions
The involvement of GH in delirium is a new finding from the present study. Also the finding of the low levels of IGF-I and the association of delirium confirms some of the previous studies. Those findings together with the association of cognitive decline with delirium strength the primary hypotheses that low brain reserves are possible the predisposing factor for delirium. Those findings needs further replication in other studies and especially in surgical samples

Research perspectives
If the above findings are replicated in future studies then the next step is clinical trials with small doses of IGF-I for prevention of delirium.
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