Hypothesis on the pathophysiology of syringomyelia based on simulation of cerebrospinal fluid dynamics

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Objective: Despite many hypotheses, the pathophysiology of syringomyelia is still not well understood. In this report, the authors propose a hypothesis based on analysis of cerebrospinal fluid dynamics in the spine.

Methods: An electric circuit model of the CSF dynamics of the spine was constructed based on a technique of computational fluid mechanics. With this model, the authors calculated how a pulsatile CSF wave coming from the cranial side is propagated along the spinal cord.

Results: Reducing the temporary fluid storage capacity of the cisterna magna dramatically increased the pressure wave propagated along the central canal. The peak of this pressure wave resided in the mid-portions of the spinal cord.

Conclusions: The following hypotheses are proposed. The cisterna magna functions as a shock absorber against the pulsatile CSF waves coming from the cranial side. The loss of shock absorbing capacity of the cisterna magna and subsequent increase of central canal wall pressure leads to syrinx formation in patients with Chiari I malformation.

Syringomyelia is a disease that produces fluid containing cavities in the parenchyma of the spinal cord. Most often it is associated with Chiari type I malformation, with a herniation of cerebellar tonsils through the foramen magnum. In 1958, Gardner described his hypothesis on the pathophysiology of this disease; he considered that an obstruction of the cerebrospinal fluid (CSF) outflow from the fourth ventricle diverts the CSF pulse waves into the central canal. His hypothesis was thus called the “water-hammer theory”. Based on his theory, suboccipital craniectomy with plugging of the obex, obstructing the opening of the central canal, became common. Recently, however, most surgeons agree with this hypothesis, although several other authors agree with this hypothesis, their opinions differ about specific mechanisms. Ball and Dayan proposed that the CSF enters the syrinx through the enlarged Virchow-Robin space in the spinal cord. Although several other authors agree with this hypothesis, their opinions differ about specific mechanisms. Ball and Dayan hypothesised that the CSF is driven into the Virchow-Robin space when the spinal subarachnoid pressure is increased during coughing or straining. They postulated that this increased pressure is caused by a one way valve-like mechanism at the craniovertebral junction, which blocks the upward CSF movement. Heiss et al thought that the piston-like movement of the herniated tonsils is responsible for producing downward pulse waves in the subarachnoid space, which contribute to the downward progression of the syrinx. Stoodley et al, on the other hand, considered that increased compliance of the spinal subarachnoid space increases the arterial pulse dependent CSF flow through the Virchow-Robin space.

In this report, we took a different approach to evaluate this problem. We theoretically analysed the dynamics of the pressure wave propagating through the spinal subarachnoid space. For this purpose, we built an electric circuit model of the spinal CSF pathway, based on the standard analysis technique of the arterial pulse wave. We then applied a sudden increase of pressure on the cranial side of this model, simulating the CSF pressure wave coming from the cranial side. By examining this model, we could analyse how the pressure wave propagated along the spinal cord.

METHODS

Figure 1 shows our electric circuit model superimposed on the actual anatomical structures. The CSF pathways are broken up into multiple nodal points starting from the fourth ventricle and prepontine cistern, followed by obex and cisterna magna, and then by nine nodal points and finally the lumbar theca. (The number of nodal points is arbitrary.) We represented the compliance (or the temporary fluid storage capacity) of the cisterna magna with a capacitor named \(C_{CM} \) in the spine, two pathways of CSF, namely the central canal and the extramedullary subarachnoid space, are represented as series of multiple resistors. To represent the compliance of the spinal cord and the dural sac, we inserted two arrays of capacitors: one between these two CSF pathways, and the other between the subarachnoid CSF pathway and the ground. Thus, the capacitors between the central canal and the subarachnoid space \(C_{CC} \) to \(C_{C} \) in Fig 1) represent the elasticity (compliance) of both the central canal wall and the spinal cord. Similarly, the capacitors between the subarachnoid space and the ground \(D_{1} \) to \(D_{9} \) in Fig 1) represent the elasticity of both the dural sac and the root sleeves. Table 1 shows the values of the parameters that we used for the calculations in this study.

We could solve this electrical diagram with the standard analysis technique of a linear system. After setting the initial voltage accumulated in \(C_{1} \) to \(C_{9} \) to zero, and that in \(C_{CM} \) to \(D_{1} \) and \(D_{9} \) to 100, we applied a step increase of voltage of 500 simultaneously on the two cranial leads. Mathematically, this problem is reduced to solving a set of linear ordinary differential equations with the voltages accumulated in all the capacitors taken as unknown variables. These equations were solved on a personal computer using a software package: Mathematica version 4.0 (Wolfram Research, Champaign, IL, USA).
RESULTS

The time course of the pressure wave propagated along the central canal in response to a sudden increase of voltage on the two cranial leads is shown in figure 2. In biological terms, the voltage accumulated in these capacitors corresponds to the pressure difference between inside and outside the spinal cord at each nodal point, a positive value of the voltage indicating a higher pressure inside. In the following figures, the horizontal coordinate shows the nodal points (1 to 9 in fig 1) in our model, the left side being the cranial side and the right side being the caudal side. The time course of the accumulated voltage at each nodal point is shown as indicated in the figures starting from the time of application of the pulsatile wave at time zero. Figure 2A, B, and C show the responses of the system with three different values of capacitance at the cisterna magna (C_m in fig 1). As the capacitance of C_m decreased, there was a dramatic increase of the central canal wall pressure. The peak of this pressure increase was at the mid-portion of the spinal cord.

The result of the simulation of syringo-subarachnoid shunting is shown in figure 3. In this case, we set the value of C_m at one hundredth of the original value. In this setting, the pressure response is increased as in figure 2C. We placed an electrical shunt at the nodal point 4 (fig 1) between the central canal and the subarachnoid space. As shown in figure 3, this shunting effectively reduced the central canal wall pressure, although it was less effective on the cranial side.

DISCUSSION

Our results showed that reduced compliance at the craniovertebral junction increased the pressure wave propagated through the spinal cord. This phenomenon is analogous to the Windkessel model of the cardiovascular system. In the Windkessel model, the aorta functions as an elastic tube with temporary fluid storage capacity. This temporary fluid storage capacity is understood with an analogy to a Windkessel—a reservoir of a water pump of a fire engine (fig 4). When the aorta becomes stiff and loses this temporary fluid storage capacity, the arterial pulse wave becomes steep resulting in systolic hypertension. Likewise, our model provides a hypothesis explaining why the constriction at the craniovertebral junction causes the formation of syrinx in the spinal cord. When the temporary fluid storage capacity of the cisterna magna is reduced as in Chiari type 1 malformation, the pressure wave propagated through the central canal is increased. In other words, we can think that the cisterna magna normally functions as a shock absorber, which absorbs the pulse pressure of the CSF coming from the cranial side. If this shock absorbing capacity is lost, because of the overcrowding of the posterior fossa in Chiari I malformation, the CSF pressure wave propagated along the central canal is markedly increased; it will then lead to leakage of CSF into the parenchyma, which precedes the formation of syrinx. Thus, we may consider that our hypothesis is a modified version of Gardner’s water-hammer theory. The difference is that our theory can explain why extradural decompression at the craniovertebral junction without intradural procedure can reduce the size of the syrinx in the spinal cord.

Our hypothesis is better in some regards than the other hypotheses presupposing the role of the Virchow-Robin space. Firstly, it can explain the phenomenon of syringomyelia with no tonsillar herniation, and why foramen magnum decompression is effective in reducing the size of the syrinx. Tubbs et al showed that these patients had moderately small posterior fossae. Our hypothesis can explain this phenomenon well, because we can assume that the temporary fluid storage capacity at the cisterna magna is reduced in such patients because of the mild overcrowding of the posterior fossa. On the other hand, other hypotheses have difficulty explaining this phenomenon, because they require either a one way valve mechanism or a piston-like movement of the cerebellar tonsils, which are less likely to occur in those patients with no Chiari malformation. Secondly, our model is compatible with the fact that syringo-subarachnoid shunting is effective in reducing the size of the syrinx. Although the syringo-subarachnoid shunting is known to have poor long term

| Table 1 The values of the resistors and capacitors in our diagram |
|------------------|----------------------|
| Parameter       | Value                |
| C1 to C9        | 0.1                  |
| Cthec           | 600                  |
| Ccist           | 600000               |
| r1 to r10       | 100                  |
| R1 to R10       | 0.2                  |
| Rs              | 0.0002               |
| Rcs             | 0.02                 |
| Rcs             | 6000000              |
| Cthec           | 6000000              |
results mainly because of obstruction of the tube, it is usually effective in short-term. If we assume that the CSF enters from the subarachnoid space to the syrinx through the Virchow-Robin space, we cannot easily explain this phenomenon; a larger conduit made by a syringo-subarachnoid shunt will, simply interpreted, rather aggravate the syrinx.

The weak point of our hypothesis might be that it assumed the patency of the central canal, or an existence of some other channel connecting the fourth ventricle and the syrinx. Only in about 14% of the patients with syringomyelia associated with Chiari type I malformation, was the connection between the syrinx and the fourth ventricle demonstrated on MRI. Also, cadaver studies showed that the central canal was obliterated in a large proportion of the human population as the age increased.

We can make several arguments concerning this point. Firstly, to say the least, our theory can explain the pathophysiology in those 14% of patients who have communicating syringomyelia associated with Chiari type I malformation. Secondly, MRI may not be able to demonstrate the connection between the syrinx and the fourth ventricle. There is some evidence that the current resolution of MRI cannot detect the normal sized central canal.

Thirdly, our model showed that if there was stenosis of the central canal at some point, the pressure wave was markedly increased immediately rostral to that stenotic point (data not shown). If we postulate that this increased pressure wave ruptures the ependyma of the central canal, and drives the formation of syrinx inside the parenchyma, our theory does not necessarily contradict the necropsy finding of progressive obliteration of the central canal in human. Fourthly, even if the central canal is obliterated in a large percentage of the elderly human population, it does not necessarily mean that the
same thing happens in the patients with Chiari type I malformation. In those patients, our results suggest that the comparatively small volume of the posterior fossa causes increased pressure waves along the central canal; this increased pressure wave may then prevent the normal process of occlusion of the central canal. Fifthly, if we closely observe our model, we find that the patency of the central canal is not absolutely required. Our model only assumed some fluid channel inside the spinal cord, and this fluid channel may well be the Virchow-Robin space.

Our theory may also be criticised because we used rather arbitrary values on the resistors and capacitors of our circuit. However, we were only interested in the qualitative behaviour of our circuit, not the quantitative determination of the amplitude and the time course of the pressure waves. The qualitative behaviour of our circuit was basically unchanged with different sets of parameters that we tested. Our study is theoretical in nature; we believe that a theory that can elegantly explain the actual phenomena can be justified as far as its assumptions do not absolutely contradict the experimental findings.

Our theory is important in the clinical perspective. Even though we have a standard surgical approach to patients with syringomyelia associated with Chiari malformation, it is not satisfactory if we do not clearly understand why it works. Better understanding of the pathophysiology will certainly improve the results of our surgery. In addition to syringomyelia associated with Chiari type I malformation, we have other types of syringomyelia such as that associated with adhesive arachnoiditis. Our theory was also effective in explaining the pathophysiology of syringomyelia associated with adhesive arachnoiditis, thus providing a unifying theory on those two types of syringomyelia. We believe that our theory can serve as a working hypothesis for further clinical and experimental works.

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