

# Direction-Selective Resistance to Cerebrospinal-Fluid Flow As the Mechanism of Syrinx Generation in Syringomyelia

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## ABSTRACT

The pathophysiology of syringomyelia is not well understood. The main theoretical problem is how cerebrospinal fluid (CSF) enters from low-pressure subarachnoid space to high-pressure syrinx and remains inside the syrinx. We approached this problem with computer simulation of the CSF flow in the spine.

## Introduction

The pathophysiology of syringomyelia is still poorly understood. A number of hypotheses exist in the literature<sup>1–13</sup>, but they provide widely different explanations on the mechanisms of syrinx generation. Nevertheless, most researchers seem to agree on the following points. First, the syrinx fluid is identical to the CSF, and there should be some communication between the syrinx and the subarachnoid space. This point is supported by many studies<sup>14–17</sup>, although a few different opinions exist<sup>13,18</sup>. Second, some derangement of CSF flow in the spinal subarachnoid space causes syrinx both in Chiari-I malformation<sup>7,19–22</sup> and subarachnoid arachnopathy<sup>23–26</sup>. Notably, the cerebellar tonsil deranges the CSF flow in the former and adhesive arachnoiditis in the latter.

The problem, however, is where this communicating channel resides and what mechanism generates the syrinx. On these points, there is no solid experimental or clinical evidence, and the opinions of researchers deviate widely. Gardner et al.<sup>1</sup> thought that the central canal intercommunicates the syrinx and the fourth ventricle, and arterial pressure waves exerted on the central canal generate the syrinx. Williams et al. also postulated the communication through the central canal, but for the syrinx generation mechanism, he emphasized the craniospinal pressure gradient produced by Valsalva maneuver et al<sup>2</sup>.

On the other hand, Ball and Dayan<sup>4</sup> assumed that CSF enters the syrinx through the perivascular space of arteries penetrating the spinal cord. This idea has the following variations. Heiss et al.<sup>7</sup> proposed that the piston-like movement of the cerebellar tonsils in Chiari-I patients generates pressure waves in the spinal subarachnoid space, which subsequently drive CSF into the syrinx through the perivascular space. Stoodley et al. also considered the perivascular space as the communicating channel, but he assumed the arterial pulse pressure as the driving force of CSF<sup>27</sup>. All these assumptions are not proven and remain hypothetical. Although the perivascular-space theory seems to be favored by recent researchers, there remains the possibility that a thin communicating channel exists between the syrinx and the fourth ventricle<sup>28</sup>.

In our opinion, the main theoretical problems reside in the following points.

1. No theory can explain the pathophysiological mechanism of syringomyelia in a unified fashion.
2. No theory can explain how CSF enters from the low-pressure subarachnoid space to the high-pressure syrinx cavity and remains inside.

As to the first point, there are different syringomyelia types, such as Chiari-I-malformation and spinal-arachnopathy-related<sup>23</sup>. The Chiari-I-malformation-related syringomyelia is further divided into communicating and non-communicating<sup>29</sup>. For each of them, current theories assume a distinct mechanism of syrinx generation. However, it may be more natural to conjecture some common mechanism underlying these different types of syringomyelia<sup>27</sup>. The second point is theoretically essential but challenging to solve. Physical theories dictate that the expanded syrinx cavity has higher pressure than the subarachnoid space<sup>7,30–32</sup>. Therefore, merely assuming a communicating channel does not explain how CSF enters the syrinx and remains inside against this pressure gradient. Even if we take a specific time window where the subarachnoid pressure exceeds the syrinx pressure, it does not explain how the CSF remains inside the syrinx after it.

The current article is part of our effort to solve the above theoretical problems. In our previous paper<sup>28</sup>, we hypothesized that if there is a resistance to CSF flow in a particular direction (rostral or caudal), it causes a one-way valve-like effect

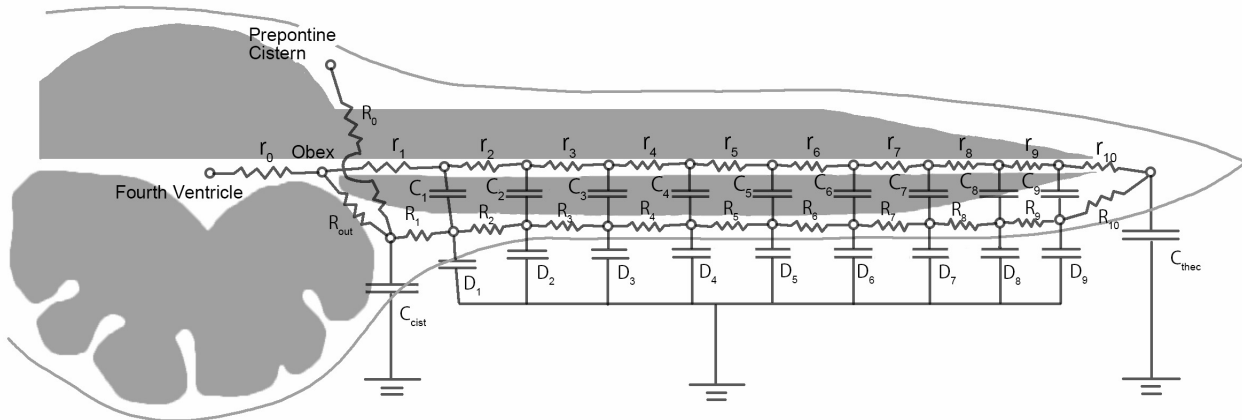
on a CSF channel inside the spinal cord, leading to the accumulation of CSF and generation of a syrinx. This hypothesis was attractive to us because it could explain the pathophysiology of both Chiari-I-malformation-related syringomyelia and arachnopathy-related syringomyelia. Namely, in Chiari-I-malformation-related syringomyelia, the herniated cerebellar tonsils may function as a direction-selective resistance. In arachnopathy-related syringomyelia, some arachnoid adhesion may play the same function<sup>26</sup>. However, in that article, we just drew a rough sketch of this process and left out a detailed explanation. The current article will describe in detail how direction-selective resistance in the subarachnoid space generates a one-way valve mechanism in the CSF channel inside the spinal cord.

For this purpose, we used a mathematical model simulating the CSF movement of the spine—a revised version of our previous model<sup>11,12</sup>. This model describes the spinal CSF movement as an electric current in a modeled circuit (a lumped parameter model with multiple compartments<sup>33</sup>), and it assumes the existence of a patent central canal. We placed a direction-selective resistance in this model at a certain point in the spinal subarachnoid space and observed how it affects the CSF flow in the central canal.

## Material and Method

Some problem exists in the computer simulation of the motion of biological fluids such as blood and CSF. Such bodily fluids move inside flexible tubes. It, therefore, requires different analysis techniques from those used in the engineering field, where the boundary of the conduit is supposed to be solid. For this purpose, researchers widely used lumped parameter models<sup>33,34</sup>. This model considers the fluid flow inside a flexible tube in analogy to the flow of electricity in an electric circuit. The accumulation of electricity in a capacitor represents the expansion of a flexible tube and the accompanying pressure elevation. An electrical resistor represents the frictional resistance to flow. This model has a wide variety. For example, it may model the whole cardiovascular system as one electric circuit, or it may model it as a synthesis of multiple compartments of electric circuits<sup>33,34</sup>. Our current model is the type of a lumped parameter model with multiple compartments. In this study, we did not intend to make a quantitatively precise model of the spinal CSF flow but to make a basic model that reveals the phenomenon underlying the generation of syringomyelia. For this purpose, we adopted a revised version of our previous lumped parameter model with multiple compartments.

Previously, we developed a mathematical model that simulated the CSF flow in the spine<sup>11,12</sup>. This model (a multiple-compartment version of a 1-dimensional lumped parameter model) could describe the CSF movement in the whole spine (Figure 1).



**Figure 1.** Schema of the electric circuit model of the CSF dynamics in the spine.

A set of differential equations can describe the behavior of this model. Using computer software, we can numerically calculate its behavior to a cranial pressure wave (defined as a boundary condition on the cranial points). This time, we improved the previous model as follows.

- We increased the number of compartments from 10 to 100, thereby making the model more precise.
- We estimated the values of the parameters (the capacitance and resistance of each component) of the model as follows so that the model will become more realistic.

- First, we set the length of the modeled spinal cord to be 1 meter.
- The resistance of the subarachnoid space ( $R$ ) was estimated using the following equations of Poisseuille<sup>35–37</sup>.

$$\Delta P = \frac{8\pi\mu LQ}{A^2} = RQ$$

- \*  $\Delta P$ : Pressure difference between the adjacent compartments
- \*  $Q$ : flow speed per unit surface
- \*  $\mu$ : viscosity coefficient. In this case, it was set to the value of water (0.0007).
- \*  $L$ : distance between the adjacent compartments. It was set to 1 cm.
- \*  $A$ : cross sectional area of the subarachnoid space. It was set to the value of a concentric annulus<sup>37</sup> with the outer diameter of 1cm and the inner diameter of 0.7 cm ( $1.6 \times 10^{-4} (m^2)$ )
- Thus,  $R$  was calculated to be  $6872 (Pa \cdot sec/m^3)$
- The resistance of the central canal ( $r$ ) was estimated using the same equation with  $A$  set to  $\pi(10^{-4})^2 (m^2)$ , i.e. the cross sectional area of a tube with a diameter of  $100 \mu m$ . Thus,  $r$  was calculated to be  $1.78 \times 10^{11} (Pa \cdot sec/m^3)$

- We determined the capacitance ( $C_{sub}$ ) corresponding to the dural elasticity so that the pressure-wave velocity determined by the time constant ( $RC$ ) will roughly correspond to the pressure-wave velocity of the downward CSF wave observed in phase-contrast MRI of normal individuals. Thus, we set  $C_{sub} = 0.1 (m^3/Pa \cdot sec)$ .

Figure 1 shows the scheme of the constructed electric circuit model. This model represents the CSF movement in the spine as electric flow through multiple compartments of capacitances connected with resistors. Table 1 shows the values of the resistors and capacitors of the system. A set of differential equations can describe the behavior of this electric circuit, and we can solve it numerically by setting the voltage at the cranial nodes as the boundary condition (Figure 2). In the previous articles<sup>11, 12</sup>, we only analyzed the transient behavior of the model to a sudden pressure increase on the cranial side of the subarachnoid space. This analysis helped simulate the situation of coughing or Valsalva maneuvers. In this article, however, we analyzed the steady-state response of the model to an oscillating cranial pressure wave simulating the normal cardiac pulsation of the CSF.

We numerically solved the differential equations using computer software (Mathematica version 12, Wolfram Research, Champaign, IL, U.S.A.). We set the boundary conditions as follows. (1) The voltage at the two cranial nodes was set to a sine wave oscillating around 100 mmHg with an amplitude of 20 mmHg at one cycle per second. (2) The initial dural pressure was set at 100 mmHg in all segments. We set the step of the numerical solution to 1/5000 second and calculated the solution from zero to 20 seconds. We displayed the obtained solution as a movie in mp4 format.

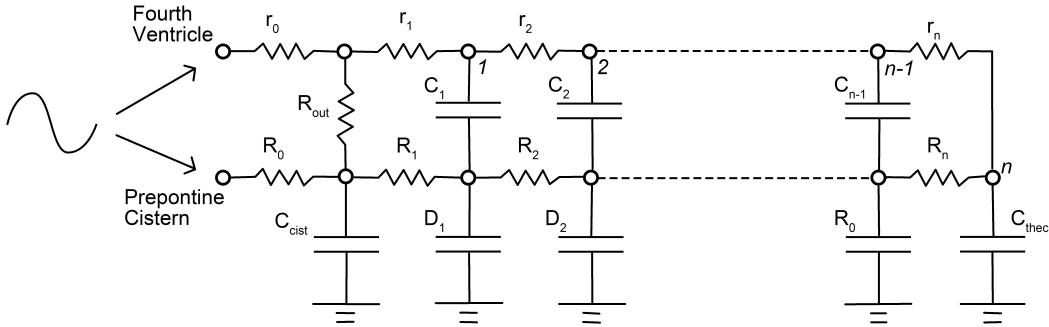
We analyzed the original normal system and two of its modifications. In the first modification, we simply increased the subarachnoid resistance at point 25 ( $R_{25}$ ) by 20 times. In the second modification, we placed a direction-selective subarachnoid resistance at point 25, so that only the resistance to the caudal flow would be increased by 20 times.

## Results

We present the systems' responses to twenty cycles of to-and-fro waves as animations. The x-axis of each animation represents the 100 nodes of the circuit laid out from the cranial to caudal direction. The y-axis displays some of the following four measures: the dural tension (voltage in D capacitors in Figure 2), the canal tension (voltages in C capacitors), the subarachnoid CSF flow (flows in R resistors), and the canal flow (flows in r resistors). We plotted the caudal flow in the positive and the rostral flow in the negative direction. The pressure values are shown in  $cmH_2O$ , the flows in  $ml/sec$ . To plot the four measures in a single animation, we multiplied the following three measures—the canal tension, the subarachnoid CSF flow, and the canal flow—with the following coefficients, respectively: namely, the canal tension with 50, the subarachnoid flow with 0.005, and the canal flow with  $1.5 \times 10^5$ .

Video 1 shows the original system's response representing the normal condition. In this state, CSF makes a smooth to-and-fro movement in the subarachnoid space with the corresponding pressure wave along the dura and the central canal. CSF also makes to-and-fro movements in the canal.

In Video 2, we increased the subarachnoid resistance  $R_{25}$  by 20 times, thereby simulating a simple block of the subarachnoid flow in both directions. In this condition, both the caudal and rostral flow along the resistance produced a drop of the dural tension downstream to the resistance, corresponding to the increased resistance both in the caudal and rostral flow phases. This pressure drop caused an increase in canal flow in the same direction as the subarachnoid flow. This increased canal flow



**Figure 2.** Electric circuit diagram representing the CSF dynamics of the spine

caused a transient increase in the canal pressure distal to the block and a decrease proximal to the block. However, these pressure changes alternated along the alternation of the flow direction and did not produce a sustained pressure increase.

In Video 3, we replaced the subarachnoid resistor  $R_{25}$  with a direction-selective resistor whose resistance to the rostral flow was unchanged but that to the caudal flow was increased by 20 times. This time, as shown in Video 3, sustained high pressure appeared in the central canal in the segment distal to the replaced resistor, and sustained low pressure in the segment proximal to it. This sustained pressure gradually accumulated as the flow cycle proceeded. The dural tension showed a pressure drop at node 25 only during the caudal-flow phase. The to-and-fro canal flow increased near the node 25 similarly to that in the simple block above, but, this time, the increase was larger in the caudal direction.

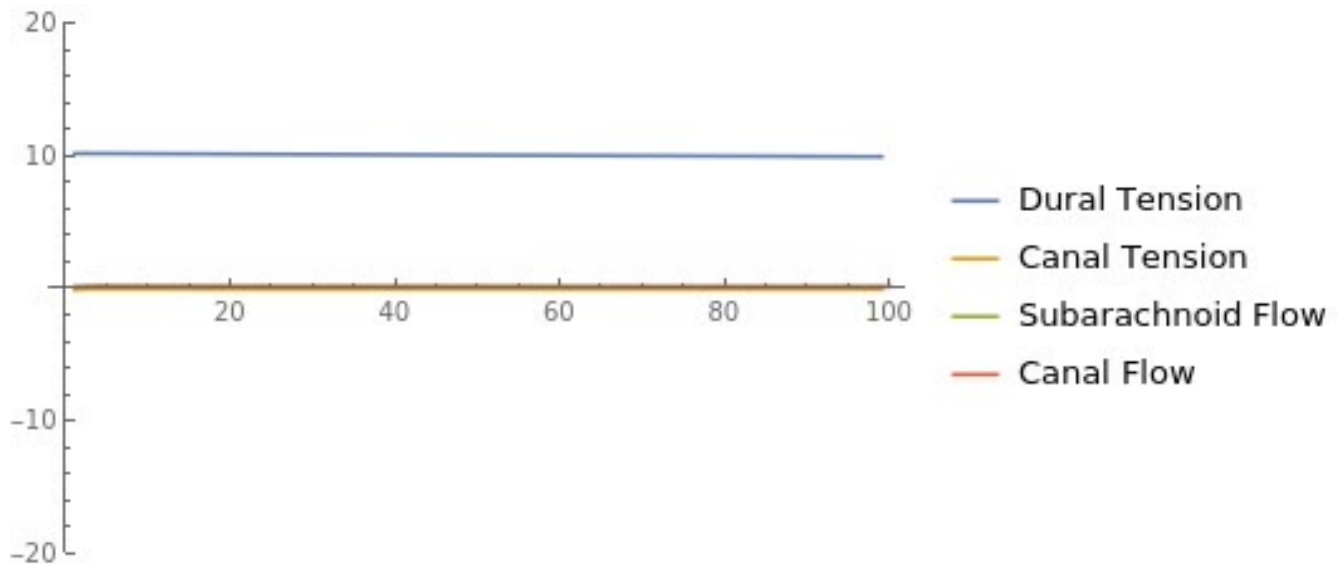
We took out the canal flow and showed it in Video 4. Observing this video, we can see that the cumulative total of the caudal flow is larger than that of the rostral flow, and it means that the CSF is virtually pumped caudally at node 25.

## Discussion

In this article, we theoretically analyzed the CSF movement in the spinal cord using a lumped parameter model with multiple components. It simulated a system with an elastic tube (dura) containing an elastic cylindrical material (spinal cord) that itself had a fluid channel inside (the central canal). When we placed a direction-selective resistor in the subarachnoid space and evoked a to-and-fro pressure wave on this system, it produced a sustained pressure elevation in the segment distal to the direction-selective resistor. This phenomenon may explain the pathogenesis of syringomyelia both in Chiari I malformation and syringomyelia associated with arachnopathy.

Subarachnoid pressure affects the pressure inside the central canal. As seen in Figure 2, the absolute pressure inside the central canal is the sum of the subarachnoid and canal pressure (voltages of  $C_k$  and  $D_k$  in electrical terms). Suppose a one-way valve selectively resists caudal flow in the subarachnoid space at point A. Caudal CSF flow creates a pressure drop across this valve, with the distal pressure smaller than the proximal one. It decreases the absolute canal pressure distal to point A because it is the sum of the subarachnoid pressure and the canal pressure. It, therefore, creates a pressure gradient in the central canal across point A, thereby increasing the distal CSF flow in the canal at that point (Figure 3).

On the contrary, reverse flow, not encountering resistance, does not create a pressure drop (Figure 4). Although some of the CSF that had been pumped caudally during the caudal-flow phase will flow back rostrally, its amount will be smaller. The net result will be that some CSF is pumped caudally in one cycle of the to-and-fro movement. Thus, CSF gradually accumulates in the distal segment of the resistance (Video 3). We hypothesize that this is the mechanism underlying the syrinx generation.



**Video 1.** Video showing the normal circuit response

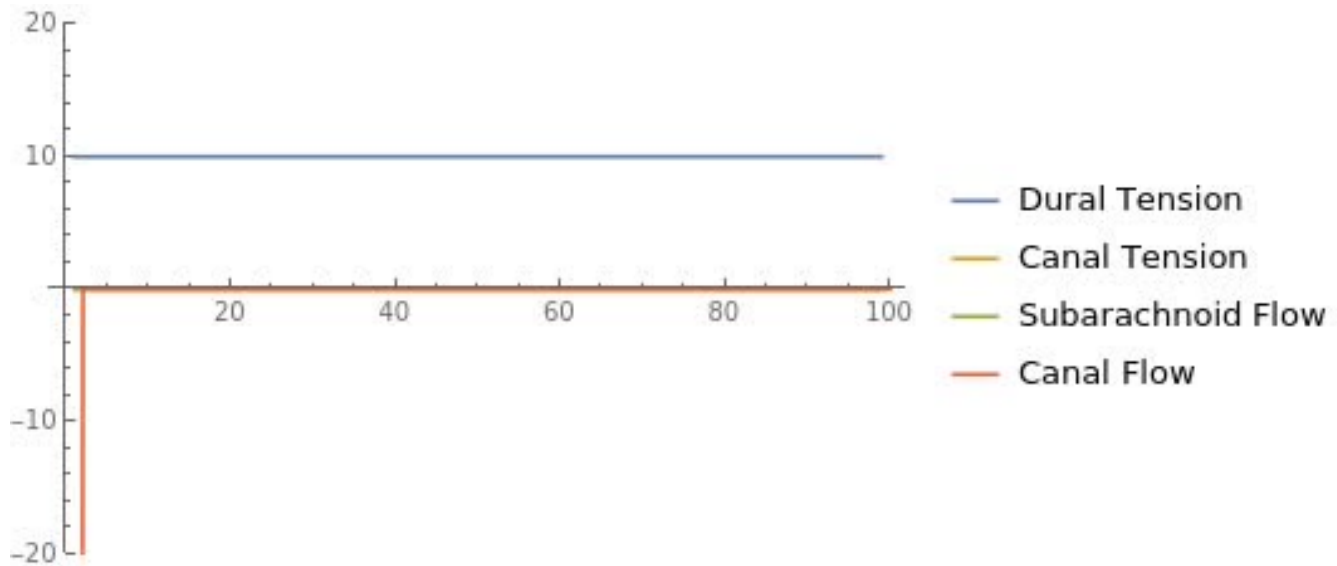
This hypothesis solves the theoretical problems pointed out in the Introduction. A one-way valve in subarachnoid space creates an asymmetry of the pressure gradient between the caudal and rostral flow phases in the central canal. This asymmetric alternation of pressure gradient effectively pumps CSF caudally, creating sustained pressure elevation in the caudal segment. In other words, the energy of the to-and-fro CSF movement is translated via the one-way valve into the creation and sustenance of syringomyelia.

Direction-selective resistance to CSF flow is not an imaginative assumption but actually exists in patients. In Chiari-I malformation, the herniated tonsils move like a ball-valve, displaced caudally during the caudal flow and rostrally during the rostral flow. Higher velocity observed in the phase-contrast MRI studies suggests that it selectively impedes the caudal CSF flow more than the cranial flow. This direction-selective resistance was demonstrated by Williams in direct measurements in Chiari-I patients and became the basis of his theory<sup>38</sup>.

Also, there is a possibility that some types of arachnoid pathology function as one-way valves. In 2014, we reported a case of thoracic arachnoid web associated with syringomyelia, in which phase-contrast MRI detected one-way-valve-like behavior of the arachnoid web<sup>26</sup>. We found an obliquely oriented arachnoid web that reminded us of a one-way valve in surgery. Thus, our hypothesis may also solve the second theoretical problem that we pointed out in the Introduction. Namely, we may consider the presence of a one-way valve in spinal subarachnoid space as a common mechanism underlying both Chiari-I-related and arachnopathy-related syringomyelia. Our theory assumes that the arterial-pulse generated to-and-fro CSF movement provides the energy for syrinx generation. It is compatible with Stoodley et al.'s experimental result that syrinx maintenance is dependent on arterial pulsation<sup>27</sup>. In this study, the authors showed that

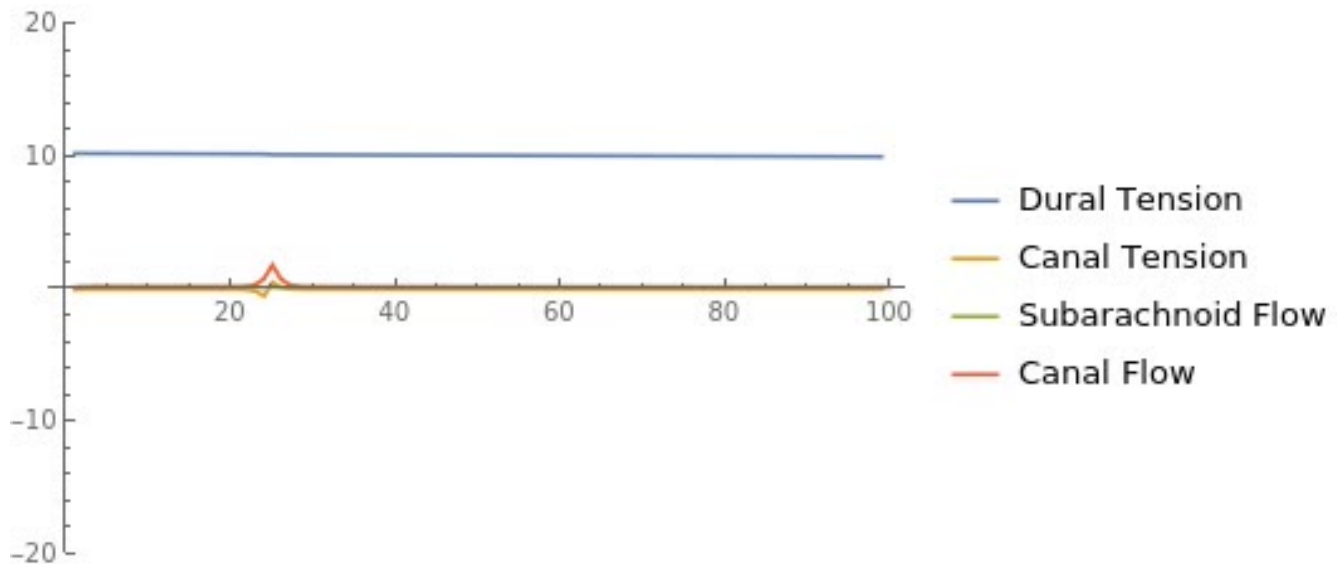
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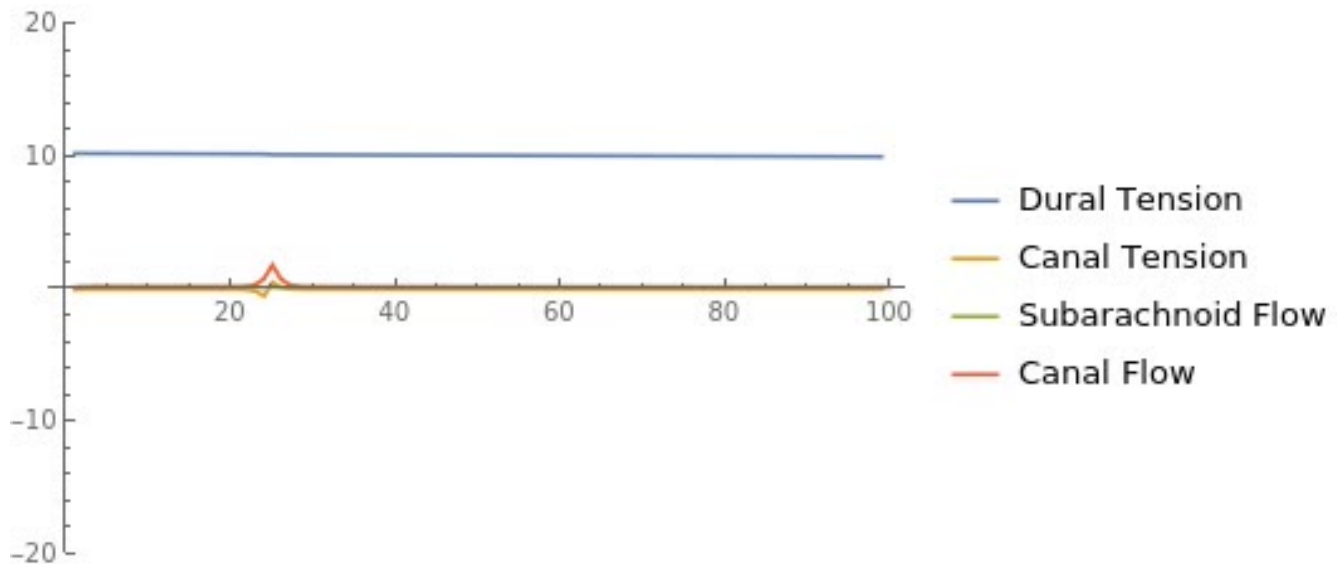
**Video 2.** Video showing the response of the circuit with a simple resistor at point 25

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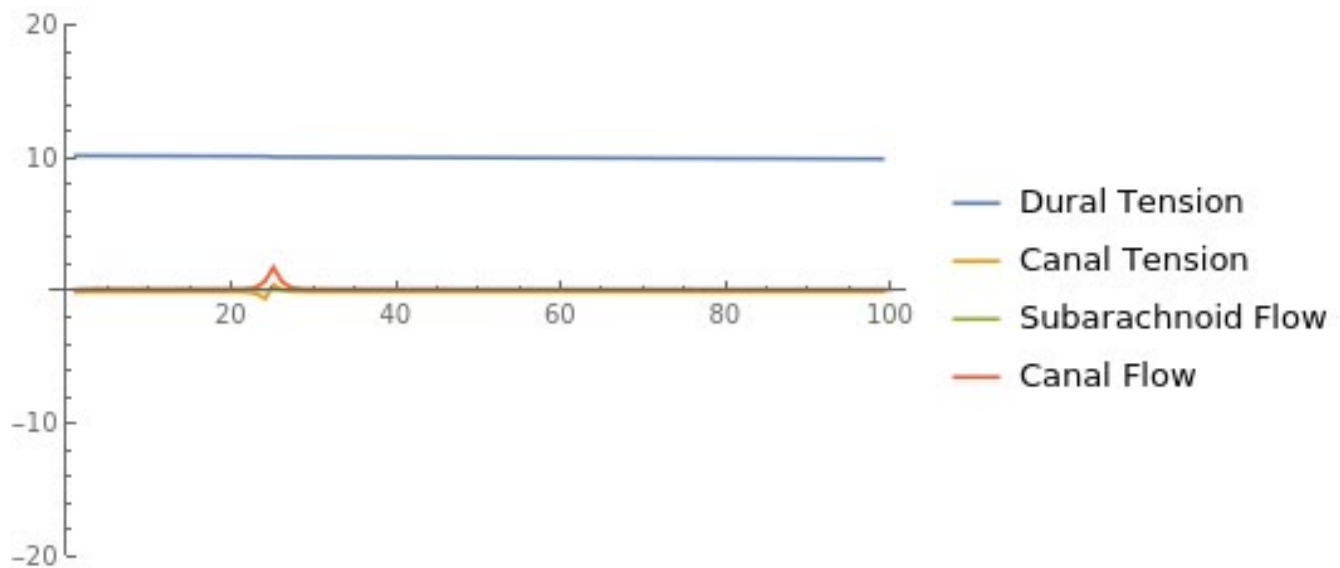
**Video 3.** Video showing the response of the circuit with a direction-selective resistor at point 25

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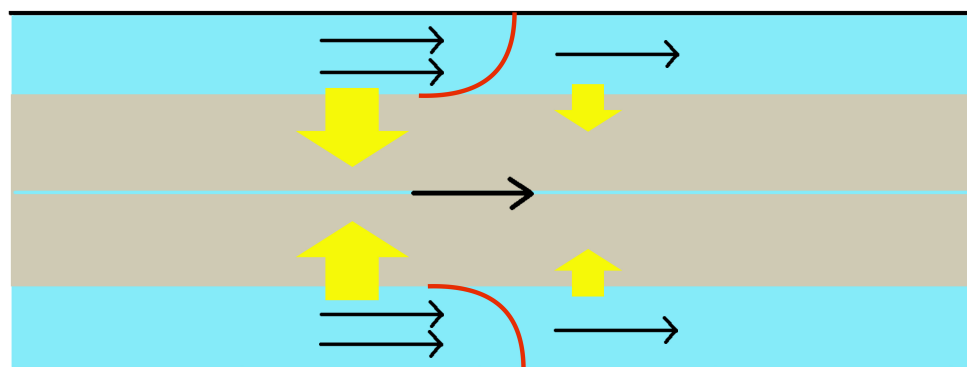


**Video 4.** Video showing the canal flow in the model with direction-selective resistance at point 25

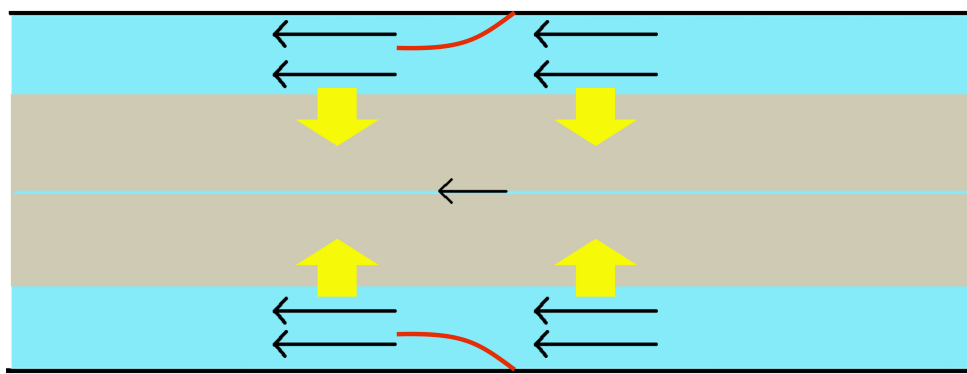
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**Video 5.** Video showing the canal flow in the two models: simple resistance at point 25 and direction-selective resistance at point 25



**Figure 3.** Increased central-canal flow across a direction-selective resistance in the subarachnoid space



**Figure 4.** Normal central-canal flow across a direction-selective resistance during the reverse flow