Protein adsorption to hydrocephalus shunt catheters: CSF protein adsorption

Howard L Brydon, Geoff Keir, Edward J Thompson, Roger Bayston, Richard Hayward, William Harkness

Abstract

Objective—To assess the quantity and nature of the proteins that adsorb to hydrocephalus shunt catheters after implantation, and to determine whether sufficient could accumulate to obstruct the catheter.

Design—Elution of proteins from 102 explanted shunt catheters, with protein assay and electrophoresis of the eluate, and scanning electron microscopy (SEM) of the catheters.

Results—The amount of protein elutable was extremely low, and significant protein, apart from a thin film, was not found on SEM. Qualitative analysis disclosed that most of the adsorbed protein was albumin.

Conclusions—Protein deposition on hydrocephalus catheters does not occur in sufficient quantities to cause catheter obstruction.

Keywords: hydrocephalus shunts; cerebrospinal fluid protein; silicone rubber; protein adsorption

It is a common belief that a high CSF protein concentration impairs hydrocephalus shunt performance. There are several mechanisms that could cause this—for example, reduced flow due to high CSF viscosity, sticking of the valve components, peritoneal malabsorption, and protein deposition obstructing the shunt lumen. We have recently investigated the first two mechanisms, and neither has been supported: protein had an insignificant effect on CSF viscosity; and the surface tension of CSF, which affects valve sticking, was lower when more protein was present. These findings have also been supported by an in vitro study on the perfusion of shunts with protein solutions.

The adsorption of CSF proteins on to shunts has not yet been studied. The aim of this project was therefore to determine the extent to which proteins accumulated on the catheter wall, and to ascertain whether this could cause catheter obstruction. In addition, a qualitative assessment of the nature of the proteins was undertaken.

The standard shunt used at Great Ormond Street Hospital for Children is the Cordis-Hakim system (Cordis UK, Brentford, Middlesex, UK), which consists of an integral valve and distal catheter. The whole shunt is often replaced during revision operations, and so a large amount of catheter material was available for study.

Materials and methods

Patients

Eighty-six permanent catheters replaced during shunt revisions were collected prospectively from 49 patients over a 15 month period (table 1): in 20 cases proximal and distal catheters were obtained from the same shunt. In addition, 16 temporary ventricular drains were collected and analysed.

Table 1 shows the patients’ original diagnoses. The mean implant time for the ventricular catheters was 25.3 months and for peritoneal catheters 34.3 months. However, this varied with the reason for removal (table 2). Most ventricular drains had been inserted for shunt infections, and they had been in situ for a mean of 10 days (range three to 15 days).

Analysis

The CSF within the catheter was aspirated, examined by light microscopy, and then cultured bacteriologically. A 1 cm length of catheter was prepared and examined by scanning electron microscopy. Adsorbed proteins were eluted from the remainder of the catheters using 0.9% saline followed by a solution of 0.5% 3-((3-cholamidopropyl)-dimethylammonio)-1-propanesulphonate (CHAPS; BDH Chemicals, Dagenham, UK) in barbitone electrophoresis buffer (pH 8.6). CHAPS was chosen as it is
stored separately at −20°C. Longer periods of elution (up to 72 hours) did not increase the yield of protein.

Attempts were made at measuring the total protein content of the eluate by protein precipitation using benzethonium chloride. Electrophoresis of the eluate (5 μl volume) was performed on agarose gel (FMC Bioproducts, Vallenbæk Strand, Denmark). The proteins were blotted on to nitrocellulose (Sartorius AG, Goettingen, Germany), and stained with gold chloride. Electrophoresis of the patients’ CSF was also performed using 2 μl volumes.

**Results**

Choroid plexus was the cause of six ventricular catheter obstructions. Another four ventricular catheters had been replaced because of malposition or disconnection. No cause for the shunt obstruction could be found in the part considered peroperatively at fault in 15 cases (eight ventricular and seven peritoneal catheters), although in seven of them the shunt valve was obstructed. The valves were subjected to a different (mechanical) analysis. In the other cases we think that the obstructive material was wiped off the shunt during its removal, although it is possible that we did not receive the part of the shunt containing the obstruction.

One peritoneal catheter was obstructed by granulation tissue growing from the outside through a hole in the wall of the catheter. This hole looked as if it had been made during insertion of the shunt, which had been implanted for 73 months. The shunt had not been infected, and the patient had normal CSF.

In no instance had sufficient protein precipitated within a catheter to obstruct it. Furthermore, scanning electron microscopy did not show a microscopic focal protein accumulation on any catheter, although a thin film was present on 15% of catheters. This was noticeable only where it had cracked during processing, and so might have been unnoticed on others (figs 1 and 2). Red and white blood cells were found on the surface of seven catheters (fig 1), degenerating cells on seven, and bacteria on four specimens. Bacteria were grown from 10 catheters removed for presumed infection, and microscopy also disclosed degenerating cell debris in these catheters.

The protein content of the shunt eluate was too low to measure (<0.01 g/l), even though 20 catheters were from patients with a CSF protein concentration >1.0 g/l (range 0.06–12.3 g/l, mean 1.06 g/l, median 0.32 g/l). Nevertheless, protein bands were identified on electrophoresis of all 102 specimens. Albumin was identified in all specimens, and in 18 (18%) this was the only significant band identified, the rest of the trace being too faint. A γ-globulin band was identified in 70 specimens (70%) and transferrin in 61 (61%). Haptoglobin was identified in 28 (28%) and tau-protein (asialotransferrin, tau-transferrin) in 13 (13%). A full electrophoretic sequence, similar to CSF, was found in only 13% (fig 3).
In 44 specimens (44%), two distinct bands were identified in the γ region on elution with both saline and CHAPS (figs 3 and 4). However, western blotting for IgG, IgM, IgA, fibrinogen, and fibronectin all proved negative. These bands were identified in ventricular and peritoneal catheters and ventricular drain specimens. They were found in catheters that had been implanted for a few days as well as for several years. Furthermore, they were not associated with any specific cause of shunt failure, being found in catheters removed for infection, overdrainage, disconnection, and obstruction. It was noted that the fraction did not store well, and on repeated electrophoresis the bands were fainter. This prevented the pooling of positive specimens for protein sequencing.

Discussion

Shunts are the standard treatment for hydrocephalus, but they are prone to complications, with up to 16% of shunts requiring revision within one month. Some consider that a high CSF protein plays a part in shunt failure, but recent work has invalidated most theories on possible mechanisms. However, the possibility of protein deposition obstructing the shunt lumen has not yet been investigated.

In 1963 Scarf stated that “...cerebrospinal fluid is laden with protein and minerals, which are gradually deposited as particulate matter on the inner surfaces of the tubes and valves, contributing to their obstruction or malfunction”. This statement, from a clinical review, was made without any supporting evidence, but is still accepted by many. Furthermore, as far as can be determined, no attempt has been made to verify or refute the statement.

The only material present in sufficient quantities to occlude the catheters in this series was choroid plexus. Several authors, analysing material found in shunt catheters by light microscopy, have reported that “fibrin”, “protein” and leucocytes have been the cause of shunt obstructions. However, they did not state whether biochemical analysis was performed to confirm the identity of the materials, nor where the presumed “fibrin” originated, as clotting proteins are normally absent from CSF. Furthermore, the composition of the patients’ CSF was not given. Bacterial slime is strand-like, and might be confused with fibrin on microscopy.

Snow and Kossovsky found a cylindrical core of tissue, up to 8 mm long, in 51% of the 57 shunts they examined. This tissue was always at the end of the catheters, and on microscopy consisted a combination of blood, fibrous tissue, neural tissue, and inflammatory tissue (sterile or infective). Similar tissues have been detected by others, but not in sufficient amounts to cause obstruction. The pathogenesis of these intralumenal tissues has not been speculated on, but there is nothing to suggest that they were related to a high CSF protein content. Material from the surgical team, or patient, including cotton, talc, and hair have also been found inside shunts.

The amount of protein that we could remove from the catheters was too small to assay, but a large amount of soluble protein would not be expected if most of the debris were of cellular origin. It should be noted that plasma contains essentially the same proteins as CSF, but in a 500-fold greater concentration, and yet shows no tendency to precipitation. Therefore, the accumulation of large amounts of CSF protein within shunt catheters should not be expected.

It could be argued that as protein adsorption is a dynamic process, more protein might have been adsorbed onto the catheters at times when the CSF protein concentration was higher, and as the CSF protein concentration fell, the amount of adsorbed protein also became less. However, our results from the ventricular drains, which were in situ for a few days only, were no different from those of the other catheters, which had been in situ for much longer. Furthermore, the ventricular drains were inserted when the CSF protein was often higher. In addition, other work has indicated that the CSF protein concentration of patients who developed early shunt obstructions was not significantly higher than of the other patients.

There remains the possibility that not all of the adsorbed protein was removed, but scanning electron microscopy only disclosed a thin biofilm on the catheters, and further suggested that it might be derived from degenerating
cells. Other groups that have studied protein adsorption in vitro and in vivo have presented qualitative results only. Methods to quantify protein adsorption have usually involved the use of radiolabelled protein in vitro.

Protein adsorption to polymers is considered by some to continue for several hours after exposure, but others consider that it is at a maximum by 30 minutes. It is thought that successive layers are formed, and that the protein in the outermost (lumenal) layer is in dynamic equilibrium with the protein in solution, whereas the protein adjacent to the catheter wall is firmly adherent. Several mechanisms, including hydrogen bonding, ionic interaction, and calcium bridges between carbonyl groups, are thought to be involved in the adsorption process. Long term consolidation of the adsorbed protein is also considered a possibility.

One group found that the total amount of protein that adsorbed to silicone rubber was 74 µg/cm² at two hours, using a 20 mg/l albumin and fibrinogen solution. This is equivalent to protein adsorption on to the inner surface of a shunt catheter at a rate of 30 µg/cm², which is far too small to obstruct it. Another group measured the amount of adsorbed protein at only 2% of this value, using a different polymer and technique.

Several workers have reported that albumin adsorbs the most to hydrophobic catheters, although γ-globulin also adsors to a significant degree. Clotting proteins, if present, also adsorb preferentially to the polymer. In this study, albumin was detected on all catheters, and so this agrees with the published work. The adsorption of albumin is advantageous, as it improves the biocompatibility of silicone rubber and inhibits bacterial adhesion, which should help prevent shunt colonisation.

The identity of the two γ region bands was not confirmed. They might be previously unrecognised CSF constituents that had accumulated on the catheters, or might be breakdown products of a more common protein fraction. A bacterial origin is unlikely as they were present on both non-infected and infected catheters. Although their nature remains unknown, their low concentration would suggest that they are unlikely to be relevant in shunt malfunction.

Conclusions
We have been unable to show that protein accumulates on shunt catheters in sufficient amounts to cause obstruction, although a thin film is formed. The major protein in this film is albumin, and other studies have shown that this will improve the biocompatibility of the silicone rubber.

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