We propose that post-shunting recurrence of ischemic priapism is a postoperative thromboembolic complication, similar to cases of postoperative thrombosis of veins (femoral or popliteal) or arteries (cerebral or coronary). Therefore, perioperative anticoagulation should be given to men undergoing these shunting procedures to prevent this complication.

Two illustrative cases are presented herein.

**Case 1**

The patient is a 22-year-old man with a history of schizophrenia, controlled with oral medications (olanzapine and valproic acid). Over the past two years, he had eight episodes of ischemic priapism, lasting between 4-14 hours, which all resolved spontaneously.

The patient presented again to a local hospital with a chief complaint of painful erection for three days. The patient was immediately transferred to our center. A thorough history yielded only recent (and only intermittent) use of olanzapine as a possible causative factor for the ischemic priapism. He complained of severe, dull pain in his penis. On physical examination, his penis was rigid and tender. There were no signs of infection. His WBC count was normal.

He was taken to the operating room, as he declined bedside procedure in the emergency department. On exam, the tip of the rigid corporal body was easily palpable at a shallow depth from the surface of the glans. A marking pen was used to make two vertical incision lines about 1 cm long, 0.5 cm lateral to urethral meatus. Local anesthetic (0.25% bupivacaine) was injected into the subepithelial layer (not the underlying spongy tissue) of the glans overlying the planned, T-shunt sites (*Figure 1*). Bilateral T-shunts were then performed using a 10-blade scalpel, in quick succession. Bilateral corporal tunneling was performed using 22-Fr. straight urethral sounds as previously described (1-3). With the sound oriented slightly laterally, the sound was passed gently to the crura without injury to the urethra. After the sound was removed, there was immediate drainage, of thick dark viscous blood. The penis was milked until blood draining forth from the shunt sites turned into a bright-red color. The T-shunt sites were closed with running-locking 4-0 chromic sutures. Care was taken to place each suture shallowly within glans tissue, so as to minimize incorporation of deeper glans tissue at the shunt site. The penis was moderately edematous, but remained non-erect throughout a 10-minute observation period and through the end of the surgery. A Foley catheter was placed which drained clear yellow urine. Following transfer to the recovery room, the patient was discharged to the ward for observation.

The patient was found to have a partially erect phallus until the evening. However, the next morning, the patient awoke with a rigid erection. Based on our exam, the patient appeared to have early recurrence of ischemic priapism. He consented to our recommendation to return to the operating room for repeat T-shunt and tunneling. We also explained to him our recommendation of perioperative anticoagulation.

He was given subcutaneous heparin 5,000 units pre-operatively. In the operating room, approximately 15 hours after his previous surgery, we removed the sutures on the glans. Upon doing so, a small clot immediately presented itself below the suture line on each side (*Figure 2*). With repeat T-shunt and tunneling, there was drainage of very dark blood that was less viscous than what was drained at surgery 15 hours prior. There was no additional clot...
noted in the blood evacuated from the corporal bodies. As before, we milked the penis until the first return of fresh bright blood. We observed the penis for several minutes and noted no recurrence of priapism. We then proceeded to close the T-shunts with 4-0 chromic interrupted locking sutures. The penis remained moderately soft and not rigid through the end of the case. He was given 325 mg of aspirin and 40 mg of famotidine after recovering from anesthesia and instructed to take baby aspirin (81 mg) and famotidine (40 mg) daily for two weeks. One more subcutaneous heparin injection was given 12 hours after the initial dose.

For the next 24 hours, the penis remained partially erect, but not rigid. The patient reported no penile pain at rest, and was discharged approximately 30 hours after his second surgery. As numerous case reports have suggested that olanzapine can cause ischemic priapism, his psychiatrist was contacted before discharge. Close outpatient follow-up (within 24 hours) with his psychiatrist was arranged prior to discharge.

At follow-up 2-week later, the patient reported no recurrence of painful erection since discharge. He confirms that he continues to take aspirin 81 mg daily. He also confirms that he awakens each morning with a good erection (painless, about 90% rigidity, sufficient for penetration), and that these erections detumescence spontaneously. Erections recur intermittently during the day and evening. On exam, no gross swelling or ecchymosis was visible. His penis was fully engorged but soft and the mid-shaft could be easily compressed. Color duplex penile ultrasound was performed. Arterial flow was evident in each cavernous artery (Figure 3) and the glans-cavernous shunt was patent with detectable flow (Figure 4).

Case 2

The second case is a 40-year-old politician who had suffered from anxiety and insomnia for several months because of a personal financial problem. He was given trazodone for insomnia. After taking the first dose, he woke up the next morning with a rigid erection. He was
well known in the community and was embarrassed to seek medical treatment. He presented to a local hospital about 48 hours later. Aspiration of old blood and injection of a diluted phenylephrine solution failed to resolve the priapism. A Winter’s shunt was performed by a local urologist. The priapism was relieved only to recur about six hours later. He was taken to the operating room and a spongiosum-cavernosum shunt was performed. The penis became soft (but swollen) after the procedure. However, persistent painful erection occurred the next morning and he was referred to our emergency room for further management. Under local anesthesia a 10-blade scalpel was used to create a T-shunt bilaterally. This was followed by bilateral tunneling with a 22 Fr. straight female sound. The priapism subsided shortly after, and the patient was discharged home.

He returned the next day, again, with a rigid painful erection. We recommended our perioperative anticoagulation therapy and started him with 325 mg aspirin and subcutaneous injection of 5,000 units of heparin. He was brought to the operating room for a repeat T-shunt procedure. When the sutures on the glans were removed, blood clots were noted at the T-shunt site. Repeated T-shunt and tunneling was performed which resulted in a partial detumescence. Manual compression of the base of the penis dislodged a few blood clots followed by non-coagulated old blood. The penis remained at partial erection but compressible ten minutes after the wound on the glans was closed with continuous locking water-tight 4-0 chromic sutures to prevent postoperative bleeding. He was given one more dose of subcutaneous 5,000 units of heparin and instructed to take baby aspirin (81 mg) daily for two weeks after discharge. His partial erection decreased gradually and the penis was flaccid after about five days. He reported fully-regained erectile function without the need for phosphodiesterase-5 inhibitors about three months later.

**Comments**

In ischemic priapism that does not respond to alpha-adrenergic agonists, various shunting procedures have been developed to re-establish circulation of the corpora cavernosa and prevent necrosis of erectile tissues. However, early closure of the newly created shunt, resulting in recurrent priapism, is a common complication which leads to repeated shunting procedures, longer hospital stays, increased patient sufferings, and less than favorable outcomes. In fact, early recurrent priapism is the most common emergent phone consultation to our andrology clinic. In subsequent exploration/re-shunting, blood clots at the site of shunting can always be expelled, followed by un-coagulated “crank-case oil”-like old blood. It is well documented that the old blood inside the corpora cavernosa does not clot due to the abundance of endothelium-derived anticoagulating and fibrinolytic factors within the corpora cavernosa (4). However, the newly created shunt is not lined and protected by endothelium. In fact, the shunt cuts a new wound through the collagen-rich tunica albuginea. The collagen-activated platelets and fibrin begin to form a clot within minutes to seal off the shunt in a similar fashion as a clot forming at the site.
of an injured blood vessel wall (5) (Figure 5). Damage to the tunica albuginea and endothelial lining of the corpus cavernosum during creation of the shunt triggers the tissue factor to bind factor VIIa and activates the cascade of coagulation. In addition, exposed collagen of the tunica albuginea causes accumulation and activation of platelets and release of inorganic polyphosphate (PolyP), a powerful activator of thrombin. The final product is thrombosis of the shunt by blood clot made of a mixture of red blood cells (RBC), fibrin and platelets.

Figure 5 Proposed mechanism of clotting of shunt modified from the cell-based model of blood coagulation (5). Damage to the tunica albuginea and endothelial lining of the corpus cavernosum during creation of the shunt triggers the tissue factor to bind factor VIIa and activates the cascade of coagulation. In addition, exposed collagen of the tunica albuginea causes accumulation and activation of platelets and release of inorganic polyphosphate (PolyP), a powerful activator of thrombin. The final product is thrombosis of the shunt by blood clot made of a mixture of red blood cells (RBC), fibrin and platelets.

Figure 6 Diagram of cross sections of lateral longitudinal view of the penis showing complete thrombosis of a small glans-cavernosum shunt (A), and a wild-patent shunt in a penis received perioperative anticoagulation and a large caliber shunt (B).
ever present, and each poses a major threat to a patient’s life. Clinical guidelines have been established (and are constantly evolving) to prevent these complications in various surgeries. Likewise, premature closure of a priapism shunt is a postoperative thrombotic complication, and, given the resulting morbidity, deserves to be treated with the same precautions as other thrombo-embolic complications. Over time, we have witnessed many of these complications along with the resulting pain and suffering of our patients. However, in the past two years, we have also observed the success of perioperative anticoagulation therapy in preventing premature shunt closure, resulting in better outcomes. We, therefore, propose that perioperative anticoagulation should be an integral part of priapism shunting procedures.

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Footnote

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