Case Reports

Cold Agglutinin Autoantibodies in a Patient without a Visible Coronary Sinus Ostium: Strategies for Myocardial Protection without Using Retrograde Cardioplegia

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Abstract: The presence of cold agglutinins (CA) during cardiac surgery with cardiopulmonary bypass usually creates the need for an altered surgical plan. In this case, the CA were discovered after the initiation of bypass, limiting the time, and cardioplegia solutions that could be used in the new approach. The inability to cannulate the coronary sinus with a retrograde cardioplegia catheter excluded the standard approach to myocardial preservation with CA of using continuous warm blood. For this case, we used intermittent cold crystalloid delivered via the antegrade needle for the first half of the procedure and through the saphenous vein graft anastomosis during the aortic valve portion of the cross-clamp period. Keywords: antibody/antigen, cardioplegia, coronary sinus, hematology.

Cold agglutinins (CA) are circulating autoantibodies that react at cold temperatures with antigens on the surface of red blood cells leading to hemagglutination, microvascular thrombosis, and/or hemolysis (1). The CA activation may be due to an idiopathic cause or more commonly a secondary infective process (2). Cold agglutination phenomenon is rare in the clinical arena, but is of greater importance in cardiac surgery due to the routine use of hypothermic temperatures while on cardiopulmonary bypass (CPB) (2). The presence of high-titer CA during cardiac surgery requires careful and specialized planning and failure to avoid agglutination can have serious consequences (1). We present a case of acute, strong CA that were discovered after initiation of CPB, requiring a complex and evolving plan of action.

DESCRIPTION

A 67-year-old male presented with moderate-severe aortic stenosis, 80% stenosis of the left main coronary artery, and 99% stenosis of the right coronary artery for a coronary artery bypass grafting (CABG) and aortic valve replacement (AVR). Nine days prior to his CABG/AVR, the patient underwent a left carotid artery bypass operation to alleviate severe carotid artery stenosis.

After uneventful induction of general anesthesia, median sternotomy was performed, the left internal mammary was harvested as a pedicle, and the saphenous vein graft was harvested. In preparation for cannulation and CPB, the patient received 35,000 units of heparin with a resulting pre-CPB activated clotting time of 495 seconds. Subsequently, the ascending aorta and right atrium were cannulated and CPB was initiated. The bypass circuit consisted of a Sorin® Inspire 6F oxygenator with integrated arterial filter and reservoir (Sorin, Milan, Italy) and was primed with 1 L of Plasma-Lyte A® (Baxter, Deerfield, IL), 10,000 units of heparin, and 50 mEq of sodium bicarbonate.

A left ventricle vent was inserted through the right superior pulmonary vein, an ascending aortic antegrade-venting cardioplegia catheter was placed, and retrograde...
coronary sinus (CS) catheter insertion was attempted via a stab incision in the right atrium. Due to difficulty inserting the retrograde CS catheter, a large right atriotomy was made after performing bicaval cannulation. Under direct vision, we were unable to identify the CS and therefore retrograde cardioplegia could not be used. Review of the intraoperative transesophageal echocardiography (TEE) showed severe CS stenosis.

Promptly after initiation of CPB, the cardioplegia system was flushed and filled with patient blood and crystalloid cardioplegic solution (Plegisol™, Hospira, Lake Forest, IL) in a 4:1 mixture. As the blood cooled to the preset 4°C in a BCD Vanguard™ (Sorin) heat exchanger, it was noted that the blood rapidly separated into large clumps (Figure 1). Cold agglutinin autoantibodies were a suspected cause of the phenomenon and a sample of the patient’s blood was sent to the laboratory for confirmation. Thermal amplitude specificity was not available, so the blood was cooled to 4°C and 4+ agglutination was confirmed approximately 30 minutes after the sample was sent. Although the patient’s specific thermal amplitude was never known, further perioperative testing revealed 2+ agglutination at 10°C and no agglutination at room temperature. This was further confirmed by altering the temperature of the water running through the BCD Vanguard.

Once the diagnosis of CA was confirmed, the typical cold blood cardioplegia strategy needed to be altered before the cross clamp was applied. Typically, Plegisol™ solution is used for cardioplegia with a K⁺ concentration of 96 mEq/L for the induction dose and 26 mEq/L for subsequent maintenance doses at a 4:1 blood to cardioplegia ratio. In a typical case with a blood K⁺ concentration of 4 mEq/L, the resulting blood cardioplegia mixture had K⁺ concentrations of 22.4 and 8.4 mEq/L, respectively. However, in this case, the cardioplegia system was thoroughly flushed before delivery, and cardiac arrest was achieved with antegrade 4°C crystalloid-only cardioplegia (buffered Plegisol™ solution with a K⁺ concentration of 26 mEq/L).

Maintenance cold crystalloid doses were given antegrade with the cardioplegic solution (buffered Plegisol™ solution with unaltered, off-the-shelf, K⁺ concentration of 16 mEq/L) every 10 minutes during the anastomoses of the two distal grafts. Following aortotomy, handheld cardioplegia via the native coronary ostia was attempted, but sufficient flow was unattainable due to severe left main stenosis. Since the retrograde catheter placement was unsuccessful, cardioplegia was administered via the saphenous vein graft to the posterior descending coronary artery during the AVR.

Prior to cross-clamp removal, warm crystalloid-only cardioplegia was administered while monitoring myocardial temperature to prevent agglutination. Once the septal myocardium reached 30°C, perfusion of the myocardium was changed from warm crystalloid to warm blood. The electrocardiogram (EKG) and myocardial temperature were monitored throughout the cross-clamp period; no EKG activity or temperature above 10°C was observed before myocardial rewarming. The maximum potassium concentration in the blood during CPB was 6.4 mEq/L and decreased to normal with zero-balance ultrafiltration and injection of insulin and glucose prior to weaning from CPB (3). After CPB weaning, TEE revealed a successful valve replacement and preserved biventricular function.

The operation consisted of a CPB time of 169 minutes and aortic cross-clamp time of 87 minutes, during which 6.7 L of fluid was ultrafiltrated, 4.5 L of crystalloid cardioplegia and 1.2 L of crystalloid fluid were given, contributing to a total net fluid balance of ~400 mL. Due to a preoperative hemoglobin of 8.5 gm/dL, two units of packed red blood cells were transfused while on CPB. During the postoperative period, the patient’s temperature was kept above 36°C, and a fluid warmer was used for any cold infusions. The patient was extubated on postoperative day (POD) 1 and then required reintubation from POD 4 to 7 due to a persistent high oxygen requirement secondary to methicillin-resistant Staphylococcus aureus (MRSA) pneumonia.

The patient was discharged on POD 14. On POD 26, the patient returned for routine postoperative evaluation and cold agglutination testing was performed. At this visit, the cold agglutination testing was negative at 4°C.

Figure 1. Agglutinated clumps of red cells falling through the plasma after rapid separation in the cardioplegia heat exchanger.
Additionally, the patient's preoperative white blood cell count (WBC) was 10.2 K/cmm compared to 20.1 K/cmm at the rewarming period on CPB. The WBC slowly trended down to 10.5 K/cmm at discharge and was 6.4 at the postoperative visit.

COMMENT

A preoperative screen/diagnosis of CA allows for thorough planning and treatment, however, most institutions do not screen for CA (2). CA autoantibodies can be significantly reduced with plasmapheresis (in cases with a high thermal amplitude) or one could wait for the CA to resolve if the cause is presumed to be acute from an infection (1,2). The causes of CA are either primary/chronic, or more commonly, secondary to an infection (1). Secondary causes include *Mycoplasma pneumoniae*, influenza B, human immunodeficiency virus, mononucleosis, Legionnaires' disease, other viral infections, and malignant disorders (2,4). In fact, 55% of patients with *Mycoplasma pneumoniae* have cold agglutinin antibodies (5). Due to this patient’s high WBC count, bipap and high-flow nasal cannula requirement post carotid bypass, prolonged intubation post coronary artery bypass surgery, and negative CA screen at his postoperative visit we believe an atypical pneumonia may be the presumed cause.

Chronic CA disease typically appears later in life, with a peak incidence at around 70 years of age, with both sexes being affected. Primary cold agglutinin disease represents a spectrum of clonal lymphoproliferative bone marrow disorders, usually with morphological signs of lymphoma (5). In chronic CA cases, monoclonal antibodies, such as rituximab, eculizumab, and rituximab, can sometimes be successful in temporary or sustained remission from CA disease (6).

CA cases with intended systemic temperatures below the thermal amplitude, such as hemi-arch replacements will require very complex planning. Almost all CA cases will require atypical myocardial protection plans as a low myocardial temperature to decrease the oxygen demand of the myocardial tissue during the cross-clamp period is unachievable (7). Discovery of CA while on CPB leaves little time for planning and preparation of additional cardioplegic solutions.

As others have reported, the typical management of CA discovered perioperatively includes continuous retrograde warm blood and intermittent warm blood cardioplegia every 15-20 minutes during the cross-clamp period (1,8,9). In this case, cannulation of the CS via a retrograde catheter was not possible as the ostium was not present in the right atrium (a rare anomaly) (10). Therefore continuous retrograde warm blood or even intermittent retrograde cold crystalloid cardioplegia were not feasible (11). In addition, during the AVR portion of the procedure, antegrade delivery was not an option due to the aortotomy and direct coronary ostia perfusion was not possible due to the severe left main stenosis. Infusion of cardioplegia via the distal grafts was critical to myocardial preservation in our case.

Although crystalloid-only cardioplegia has been studied and accepted as an appropriate option, it requires special preparation of the appropriate cardioplegia solution. This could be simplified with off-the-shelf cardioplegic solutions such as Plegisol® (St. Thomas) or Bretschneider's histidine-tryptophan keto glutarate (Custodiol® HTK) (12–16). If these solutions are not available, the custom-adjusted cardioplegic solution could be prepared by the pharmacy with an appropriately lower potassium concentration. Although time is of the essence, especially if the heart is already arrested upon the discovery of CA, careful attention must be paid to the proper formulation of the new, custom cardioplegic solution.

Although the exact thermal amplitude was not known in this case, if it was known, moderate hypothermic blood cardioplegia could have been used as long as it was above the thermal amplitude and topical ice was not used. Additionally, for mitral valve replacements or isolated CABG operations, the case could be completed with induced ventricular fibrillation instead of aortic cross clamping, avoiding the need for cardioplegia (17,18). An isolated CABG could be performed with a beating heart technique using a partial cross-clamp to avoid the need for cardioplegia (19). Moreover, for a patient with a high thermal amplitude needing an AVR, the use of transcatheter aortic valve replacement (TAVR) may be indicated (20).

This case was unique in that during the aortic valve portion of the procedure, retrograde cardioplegia, antegrade cardioplegia, direct ostial cardioplegia, ventricular fibrillation, or partial cross clamp were not feasible myocardial protection strategies. This left the newly sewn saphenous vein graft to the posterior descending coronary artery as the only method of delivery for the cardioplegic solution.

Complications from failing to prevent CA activation include ischemia due to vascular occlusion, cerebral insults, as well as renal and hepatic dysfunction (21). Early detection of CA is therefore crucial for planning and to avoid these serious complications. While preoperative CA screening is done at a few centers, the incidence of a positive screen is thought to be around 3–4% (22,23). Since CA pre-screening is not the norm, prompt detection upon CPB initiation is crucial. This involves priming the cold cardioplegia unit with blood cardioplegia as soon as possible (24). Often this requires priming the cardioplegia line with the scrub nurse before the surgeon has inserted the antegrade or retrograde...
cardioplegia device to prolong the CA detection window before the cross clamp is applied. In this case, the cardioplegia line was flushed to the field promptly after CPB initiation, and the difficulties cannulating the CS allowed enough time for CA detection by the perfusionist.

In retrospect, it was noted that the arresting dose could have started with warm crystalloid only cardioplegia to wash out any residual blood in the coronaries before it was exposed to an agglutinating temperature. This approach was not chosen due to the predicted time it would take our heater cooler 3T™ (Sorin®) to cool the cardioplegia down after the initial warm delivery. Another additional precaution would be to add a way to capture the cold cardioplegia at the draining point to avoid interaction with any warm blood. Given the complex and varied routes of cardioplegia delivery that was not feasible in this operation.

Although CA are a rare and feared obstacle encountered during cardiac surgery, with proper detection and planning serious complications can be avoided (1,2,11,22–26).

REFERENCES