Perfusion Strategies for Bivalirudin Anticoagulation: AIIMS Protocol

Gaurav Sharma¹ Suruchi Hasija² Poonam Malhotra Kapoor²

¹Department of Perfusion Technology, All India Institute of Medical Sciences, New Delhi, India
²Department of Cardiac Anaesthesia and Critical Care, All India Institute of Medical Sciences, New Delhi, India

Abstract

Anticoagulation strategies for cardiac surgery are witnessing a change with the identification of serious limitations of heparin, including development of resistance in 3 to 13% of patients undergoing cardiac surgery and heparin-induced thrombocytopenia/thrombosis syndrome in 1 to 5.5% of patients. Heparin alternatives have a potential role in these scenarios. Bivalirudin, a reversible direct thrombin inhibitor, has an onset time of 2 to 4 minutes and half-life of 25 minutes, is eliminated mainly by a proteolytic mechanism, does not require antithrombin III for effect, and is nonimmunogenic. The considerations for extracorporeal circulation are peculiar with its use, and this article outlines the aspects of initiating, maintaining, and terminating cardiopulmonary bypass and extracorporeal membrane oxygenation with bivalirudin as the anticoagulant.

Keywords
► anticoagulation
► bivalirudin perfusion strategies

Introduction

Anticoagulation strategies for cardiac surgery are witnessing a change with the identification of serious limitations of heparin, including development of resistance in 3 to 13% of patients undergoing cardiac surgery and heparin-induced thrombocytopenia/thrombosis syndrome in 1 to 5.5% of patients.¹ Heparin alternatives have a potential role in these scenarios. Bivalirudin, a reversible direct thrombin inhibitor, has an onset time of 2 to 4 minutes and half-life of 25 minutes, is eliminated mainly by a proteolytic mechanism, does not require antithrombin III for effect, and is nonimmunogenic, and does not have an antidote. The considerations for extracorporeal circulation are peculiar with its use, but there are no standardized protocols. This article outlines the protocol followed at the authors' institution, highlighting the technical aspects of initiating, maintaining, and terminating cardiopulmonary bypass (CPB) and extracorporeal membrane oxygenation (ECMO) with bivalirudin as the anticoagulant.

Conduct of CPB with Bivalirudin as Anticoagulant

Pre-CPB

- Heparin-coated oxygenators are not used. Instead, heparin-free oxygenators are used.
- Heparin-coated tubings are not used. Instead, polyvinyl chloride tubings are used.
- Bivalirudin is added while priming the CPB circuit, instead of heparin.


© 2022. Official Publication of The Simulation Society (TSS), accredited by International Society of Cardiovascular Ultrasound (ISCU). All rights reserved.
This is an open access article published by Thieme under the terms of the Creative Commons Attribution-NonDerivative-NonCommercial-License, permitting copying and reproduction so long as the original work is given appropriate credit. Contents may not be used for commercial purposes, or adapted, remixed, transformed or built upon. (https://creativecommons.org/licenses/by-nc-nd/4.0/)
Thieme Medical and Scientific Publishers Pvt. Ltd., A-12, 2nd Floor, Sector 2, Noida-201301 UP, India
**Table 1** Bivalirudin dosage protocol

<table>
<thead>
<tr>
<th></th>
<th>Adults(^2,3)</th>
<th>Children older than 1 year(^4,5)</th>
<th>Children younger than 1 year(^6)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Flush solutions, mg/mL</td>
<td>0.1</td>
<td>0.1</td>
<td>0.1</td>
</tr>
<tr>
<td>Prime, mg</td>
<td>50</td>
<td>50</td>
<td>25</td>
</tr>
<tr>
<td>Pre-CPB</td>
<td>1 mg/kg followed by 2.5 mg/kg/h</td>
<td>1.7 mg/kg followed by 3 mg/kg/h</td>
<td>1 mg/kg followed by 2.2 mg/kg/h</td>
</tr>
<tr>
<td>CPB*</td>
<td>2.5 mg/kg/h</td>
<td>3.0 mg/kg/h</td>
<td>2.2 mg/kg/h</td>
</tr>
<tr>
<td>ECMO</td>
<td>Loading dose 0.1–1.0 mg/kg</td>
<td>Maintenance dose 0.3–0.5 mg/kg/h</td>
<td></td>
</tr>
</tbody>
</table>

Abbreviations: CPB, cardiopulmonary bypass; ECMO, extracorporeal membrane oxygenation.

*Target plasma concentration > 10–15 μg/mL.

- Blood is added to the prime just before CPB is commenced to ensure that bivalirudin levels are not depleted on exposure to blood.
- Activated clotting time (ACT) sample is drawn 5 minutes after administering bivalirudin.
- ACT of 400 s or 2.5 times baseline is ensured before cannulation and commencing CPB (**Table 1**).

**CPB**

- It is of paramount importance to avoid stagnation in all components of the CPB circuit. Stasis of blood rapidly depletes the local concentration of bivalirudin and triggers the coagulation process.\(^3,4\)
- ACT is monitored religiously every 30 minutes to ensure value above 400 s.
- To avoid chamber and chest cavity stasis, cardiotomy suckers are frequently used.\(^6\)
- The potential areas where blood can stagnate are the shunt lines, cardioplegia circuit, and hemofilter. The shunt lines containing blood, for example, from oxygenator or arterial filter to reservoir are ideal sites for clot formation if not in use for long periods. Therefore, these are flushed at regular intervals of 15 minutes to avoid stagnation.\(^4,5\)
- The choice of cardioplegia solution and circuit can be modified depending upon the surgical procedure. Use of sanguineous cardioplegia is a good option for surgeries that be completed in a shorter clamp time, whereas asanguineous cardioplegia covers for longer procedures.
- While using sanguineous cardioplegia, there are chances of clot formation as the circuit remains idle for a period varying between 20 and 60 minutes. The cardioplegia pump/circuit is potentially at risk for clot formation in the time period between the cardioplegia delivery schedule (20 minutes for St. Thomas’ solution and 60 minutes for del Nido solution).
- The other option is to use asanguineous cardioplegia such as Bretschneider’s (Custodiol) solution wherein the risk of clot formation is averted altogether. Crystalloid cardioplegia solutions give a safe cardiac arrest period of 90 minutes, which is usually sufficient for most cardiac surgeries. In case there is prolongation of arrest period, supplemental doses can be administered using a pressure bag and sparing the cardioplegia circuit.
- Hemofilter is an important and integral part of CPB circuit, which not only removes extra volume but also reduces the inflammatory mediators. During the periods of CPB when hemofiltration is not being performed, clot formation can be avoided by allowing blood to continuously flow through the hemofilter by clamping the effluent port/filtrate outlet line (**Fig. 1a**).\(^5\)
- Bivalirudin (molecular weight: 2,180 Da) can pass through hemofilter membrane (pore size: 60,000–65,000 Da). It is eliminated to the extent of 45 to 69% and its half-life decreases by 20%.\(^7\) Therefore, dose escalation may be needed when hemofiltration is being performed.

**Post-CPB**

- After weaning off CPB and stabilizing the patient hemodynamically, the circuit is kept on circulation by opening the recirculation line and connecting the arterial–venous loop.
- Bivalirudin infusion is continued in the circuit till the patient is shifted and stable in the intensive care unit. This is to ensure that the circuit remains clot-free in case the need for a subsequent CPB run arises.
- The volume in the venous reservoir may have to be built up adequately. To this, another 50-mg bolus of anticoagulant is added and the solution is circulated through the entire circuit, keeping all purge lines/shunt lines open.

**Conduct of ECMO with Bivalirudin as Anticoagulant**

- It is of paramount importance to avoid stagnation in all components of the ECMO circuit. Stasis of blood rapidly depletes the local concentration of bivalirudin and can initiate the coagulation process (**Fig. 2a,b**).
- Use of heparin-coated ECMO circuit is avoided.
- As ECMO is a closed circuit, proper cannulation is essential to avoid airlock/air embolism. The arterial and venous cannulas with inbuilt Luer-lock ports are used.
A loading dose is unnecessary unless it is in the subtherapeutic range and/or in the context of suspected thrombosis (►Table 2).8

Target ACT is 180 to 220 s and target activated partial thromboplastin time is 50 to 80 s.8

The use of three-way stopcocks is minimized as they are potential areas of initiation of clot formation. All the three ways are flushed at regular intervals or exchanged if required.

The shunt line clamp is released regularly.

Stagnation is avoided during hemofiltration by keeping blood flowing through it continuously (►Fig. 1b). The ACT is checked periodically if hemofiltration is in use. Thrombosis in the ECMO circuit/membrane can lead to increase in circuit pressure/resistance causing rupture of the membrane.

Stagnation of blood is also avoided in the cardiac chambers if there is lack of ejection/contractility.9

A shunt line with one-way valve between arterial cannula and venous cannula is placed to keep the circuit in circulation for trial off or standby conditions (►Fig. 3).

ECMO flow is increased whenever anticoagulation is reduced to minimize the risk of thrombus formation.

The pump flow rate is never reduced below the recommended flow rates for the given centrifugal pump.

A standby ECMO circuit is always made available.

### Table 2 Advantages and disadvantages of bivalirudin

<table>
<thead>
<tr>
<th><strong>Advantages</strong></th>
<th><strong>Disadvantages</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td>Safe and efficacious</td>
<td>Mandates increased vigilance</td>
</tr>
<tr>
<td>Predictable response</td>
<td>– Avoid stasis in all components of the CPB circuit</td>
</tr>
<tr>
<td>Not associated with:</td>
<td>– Looking out for clot formation</td>
</tr>
<tr>
<td>– Resistance</td>
<td>– Frequent anticoagulation monitoring</td>
</tr>
<tr>
<td>– Immunogenicity</td>
<td>Requires dose adjustment</td>
</tr>
<tr>
<td>– Less platelet dysfunction</td>
<td>– Renal impairment</td>
</tr>
<tr>
<td>– Inhibits free and clot-bound thrombin</td>
<td>– Hemofiltration</td>
</tr>
<tr>
<td>– Can be monitored with ACT plus</td>
<td>No reversal agent, extra time expended on surgical hemostasis</td>
</tr>
<tr>
<td></td>
<td>Cost</td>
</tr>
</tbody>
</table>

Abbreviations: ACT, activated clotting time; CPB, cardiopulmonary bypass.
Disclaimer
The protocol presented herein is followed at the authors’ institution and readers may adopt the practical suggestions as per their discretion. The ECMO circuit is improvised to make it less complicated and easy to manage by using fewer connectors, stopcocks, and avoiding potential sites of flow turbulence and blood stasis.

Conflict of Interest
None declared.

References
1 Kaplan JA, Augoustides JGT, Manecke GR Jr, Maus T, Reich DL. Kaplan’s Cardiac Anesthesia: For Cardiac and Noncardiac Surgery. 7th ed. Philadelphia, PA: Elsevier; 2017


