The Multiple Faces of Heparin: Opportunities in COVID-19 Infection and Beyond

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Anticoagulant and Nonanticoagulant Actions of Heparin

The fascinating multifactorial faces of the polysulfated glycosaminoglycan heparin were reported first by Doyon in 1912. He revealed heparin’s nonanticoagulant action when he investigated the effect of peptone on blood coagulation in canine livers. McLean and his mentor Howel observed an anticoagulant effect in the liver when they purified a thromboplastic component from various organs and named it heparin, according the Greek name of its origin (ηπαρ = hepar). These contradictory actions of heparin are still observed until today such as they can have disastrous thrombotic side effects in the form of heparin-induced thrombocytopenia.¹

The chemical composition of heparin was discovered by Chargaff and Olson in 1937. They found that the negatively charged heparin binding to positively charged protamine sulfate antagonized the anticoagulant property of heparin. Jorpes and Bergström identified a polysulfated glucosamine/glucopyranose and uronic acid of the heparin polysaccharide. The function of heparin still remains under investigation because the sulfated saccharides have a three-dimensional structure for several reasons such as it changes upon binding to proteins and cell surfaces.²

Heparin’s Multifaceted Structure

In this issue, Beurskens et al from the cardiovascular institute in Maastricht published a review on the pleiotropic anticoagulant and nonanticoagulant effects of heparin.³ They describe in detail the structure of heparin as a glycosaminoglycan. The chemical structure of heparin is commonly known by its anticoagulant activity toward factor Xa and thrombin, which is mostly related to its high affinity binding to antithrombin (AT). Low affinity binding of heparin to AT mediates anticoagulant as well as nonanticoagulant action. Furthermore, the structural composition of heparin varies between species and even tissues; commercially available forms of heparin are typically derived from porcine intestinal mucosa and rarely from bovine intestinal mucosa; from bovine lung it is not extracted any more for clinical use. The growing demand for heparins has pushed us to consider other sources, such as from the ovine intestinal mucosa, and bioengineered heparin.

Some additional structural aspects of heparin compared with Beurskens et al³ include a relationship between different disaccharide components and how these relationships can identify the origin of a heparin sample obtained by industrial extraction processes.⁴

• Different structures for as many as five pentasaccharide sequences for the AT binding (ATB) region of mucosal bovine and porcine heparins were identified.⁵
• Two molecules of trisulfated disaccharides of an 18-unit oligomer containing ATB at reducing end are required to maximize the effect of AT on the anionic charge density of thrombin.⁶
• The conformational flexibility of the iduronic acid chain components induces heparin’s pleiotropic properties.
• Shorter chains than 18 monomers containing AT sites are devoid of anticoagulant activity but have antithrombotic activity.

It is astonishing that heparin’s structure continues to surprise us today despite its use as a cornerstone therapy.
for thromboembolism for more than 50 years. This structural variation may be explained by heparin’s glycosylic structure; a synthetic full-length active polyanionic oligosaccharide sequence is not yet commercially available, unlike proteins consisting of amino acids. Furthermore, improving physicochemical methods can now better detect three-dimensional peculiarities after binding to proteins.\(^4\)

**Heparin’s Anticoagulant and Nonanticoagulant Actions**

Beurskens et al\(^3\) have precisely described the many antithrombotic and anticoagulant actions of heparin. Antithrombotic actions include release of tissue factor pathway inhibitor, modulation of fibrinolytic activity, binding of chemokines and cytokines, and activation of growth factors. Nonanticoagulant actions include inhibition of tumor growth, metastasis, inflammation, and neutrophil extracellular traps (\(\rightarrow\) Fig. 1). The authors conclude that the clinical benefits of heparin outweigh its many and potentially life-threatening side effects.

The multiple actions of heparin comprise the following processes:

- Heparin and other polyanions such as dermatan sulfate (located in the vessel wall) potentiate the antithrombotic activity of heparin cofactor II.
- Heparin inhibits protein Z (PZ) by potentiating a PZ inhibitor and thereby inhibiting blood coagulation.
- Heparin potentiates C1-esterase inhibitor action on the contact system by inhibiting kallikrein and factor XIIa.\(^2\)
- Heparin inhibits the naturally occurring inhibitor of heparanase.\(^7\)

**Anticoagulant Actions of Heparin in COVID-19**

Beurskens et al reviewed heparin’s actions before the coronavirus disease (COVID-19) pandemic caused by the severe acute respiratory syndrome coronavirus-2 (SARS-CoV-2).\(^8\) The most severely ill patients initially present with respiratory insufficiency that progresses to multiple organ dysfunction, involving septic-induced coagulopathy and eventually disseminated intravascular coagulation.\(^9\) The treatments of choice for disseminated intravascular coagulation are unfractionated heparin, low molecular weight heparin (LMWH), and fondaparinux.\(^10\) These reduce mortality and normalize coagulation marker levels, prothrombin time, activated partial thromboplastin time, fibrinogen levels, platelet count, and D-dimer levels.\(^11\) Twice daily administration of LMWH appears to have a more protective effect in COVID-19 patients than once daily dose, probably because of better rhychemeral coverage.\(^12\)

**Heparin’s Nonanticoagulant Antiviral Actions**

Unfractionated nonanticoagulant heparins are obtained by desulfation processes that modify the anionic charge of the AT domain, or by glycol split of heparin. Glycol split does not change the number and distribution of anionic charges and introduces flexible joints to the polymer chain, which increases the freedom of charge orientation toward the target proteins’ cationic sites. Another therapeutic application of heparin is its use as an inhibitor of viral adhesion.\(^13\) This is also supported by the observation that heparin disrupts the interaction of the SARS-CoV-2 surface protein Spike with its host cell receptors of various organs via its S1 receptor\(^14\) as follows:

- Angiotensin-converting enzyme 2, a metallopeptidase, was identified as one of the functional binding receptors allowing SARS-CoV to enter host cells.\(^15\)
- The immune changes associated with coagulation in COVID-19 patients may resemble the NETosis process observed during bacterial, fungal, and some viral infections.\(^16\)
- Lesions of the endothelium destroy the glycocalyx leading to microthrombotic reactions and extravascular fluid leaks. Heparin can enter the glycocalyx\(^17\) to mobilize syndecan pools\(^18\) and partly take over syndecan 1 function.\(^19\) This restores glycocalyx function in the vascular endothelium to prevent inflammation\(^20\) and reduce septic shock.\(^18\)

Fig. 1 Interplay of anticoagulant and nonanticoagulant actions of heparin and heparin-like compounds on biological functions.
Autopsies of patients who died of COVID-19 revealed microvascular thrombosis in many organs such as lung, liver, and heart, as well as neurological effects such as severe acute hearing loss which indicate viral sepsis.

Heparins achieve their multiple nonanticoagulant actions by inhibiting heparanase, which is elevated in malignancies, chronic inflammation, atherosclerosis, and Alzheimer’s disease (► Fig. 2). Advances in the field are presented at annual symposia; the 28th symposium will be postponed until 2021 because of the COVID-19 pandemic. The anticoagulant and nonanticoagulant actions of heparin make it a promising candidate for the treatment of COVID-19.

Extended Anticoagulation and COVID-19

Ideally, heparins should be administered to COVID-19 patients as early as possible. Before hospital admittance, anticoagulation therapy should be in the form of indirect-acting vitamin-K antagonists (VKA) or direct-acting oral anticoagulants (DOACs) in patients with nonvalvular atrial fibrillation and thromboembolism. Upon hospitalization, anticoagulation therapy should be switched as soon as possible to heparin/LMWH to ensure beneficial effects and to reduce interactions of VKA and DOACs during polypharmaceutical treatment of COVID-19.

Thrombosis and Haemostasis
References