Innate immunity activation on biomaterial surfaces: A mechanistic model and coping strategies

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ABSTRACT

When an artificial biomaterial (e.g., a stent or implantable pump) is exposed to blood, plasma proteins immediately adhere to the surface, creating a new interface between the biomaterial and the blood. The recognition proteins within the complement and contact activation/coagulation cascade systems of the blood will be bound to, or inserted into, this protein film and generate different mediators that will activate polymorphonuclear leukocytes and monocytes, as well as platelets. Under clinical conditions, the ultimate outcome of these processes may be thrombotic and inflammatory reactions, and consequently the composition and conformation of the proteins in the initial layer formed on the surface will to a large extent determine the outcome of a treatment involving the biomaterial, affecting both the functionality of the material and the patient's life quality. This review presents models of biomaterial-induced activation processes and describes various strategies to attenuate potential adverse reactions by conjugating bioactive molecules to surfaces or by introducing nanostructures.

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1. Introduction

1.1. Hemocompatibility

Medicine today utilizes a wide range of biomaterials, most of which make contact with blood either permanently or transiently. In the United States alone, it is estimated that more than 25 million patients have some kind of implanted device [1]. Even more patients are treated with non-implanted and temporary biomaterials (including 400 million catheters, 25 million renal dialyzers, and more than 2 million stents annually) [2]. In 2007, the combined applications in drug delivery and medical devices have been estimated to generate a market of $200 billion in the US alone [3]. Despite considerable progress in biomaterial engineering and clinical development, many materials and procedures are still associated with undesirable side effects. For example, hemodialysis can contribute to systemic inflammation and accelerated arteriosclerosis, and treatment with stents is associated with thrombosis. These adverse reactions are initiated by contact between the biomaterial and the defense systems in the blood, primarily the cascade systems, which leads to cellular activation. Thus, the elusive goal of blood compatibility has not yet been attained [4].

Under physiologic conditions, direct contact with the blood is restricted to the intact endothelial cell lining of the blood vessels, the only truly hemocompatible surface found in nature. Activation of complement and clotting is triggered by any disruption of this surface or by the introduction of foreign materials, non-blood cells, or microorganisms into the circulation. The plasma cascades comprising the complement, coagulation, contact activation (or kallikrein/kinin), and fibrinolysis systems all act according to similar principles; and interactions between these systems, both direct and indirect, (i.e. cell-mediated) have long been known to occur. Furthermore, there is tight crosstalk between these cascade systems and the platelets and leukocytes during the induction of clotting and inflammation [5–7]. In addition, the endothelial cells play active roles not only in propagating an inflammatory/thrombotic event [6] but also by providing platelet inhibitory compounds [8].

1.2. Complement

The complement system is a primary contributor to the innate immune system of the host, clearing the body of foreign cells and organisms through direct lysis or by recruiting leukocytes that promote phagocytosis. The cascade consists of an intricate network of plasma proteins and cell surface-bound receptors and regulators. Its activation involves four steps: i) recognition of non-self surface patterns via initiation of different pathways; the classical and lectin pathways (CP and LP) are induced by antigen–antibody complexes and by certain carbohydrates, respectively, and the alternative pathway (AP), may be triggered directly by foreign surfaces (e.g. by man-made biomaterials); ii) activation of complement component C3 into C3a and opsonizing C3b by two multi-molecular enzyme complexes called C3 convertases; iii) initiation of an amplification loop by the AP, which leads to the vast majority of all C3 activation, because surface-deposited C3b initiates the formation of more AP convertase complexes (C3bBb); iv) generation of convertases that are able to activate component C5 into the potent anaphylatoxin C5a and the fragment C5b, which may induce formation of the terminal complement complexes (TCC or sc5b-9). The anaphylatoxins (C3a and C5a) activate and recruit phagocytes and other immune cells, while target-bound C3 fragments facilitate binding to and activation of the recruited cells [9].

In vivo, the complement system is controlled by several soluble and membrane-bound regulators that protect self-cells against damage caused by autologous complement activation products. Most of the modulators are members of the regulators of complement activation (RCA) superfamily, which act at the level of the convertases. The plasma proteins Factor H (regulator of the AP) and C4b-binding protein (C4BP, regulator of the CP), as well as the cell-surface-bound protein membrane cofactor protein (MCP), decay acceleration factor (DAF), and complement receptor 1 (CR1 or CD35) all belong to this family. Both C4BP and Factor H not only attenuate complement activation in circulation but also recognize specific pattern (e.g., glycosaminoglycans [GAGs] and sialic acid) on host cells and thereby support complement inhibition on surfaces (i.e., self-recognition) [9,10].

1.3. Crosstalk between blood cascade systems and cells in inflammation

It has been known for several decades that activation products of the contact activation system (Factor Xla [FXlαa] and kallikrein), as well as thrombin and plasmin, are able to cleave purified complement component or fragments thereof in vitro [11–15]. Recently, these early observations have been confirmed and extended, and FXla, FXa, and FⅩⅤa have been added to the list of proteases that potentially are able to bypass convertases and directly generate C3a and C5a, respectively [16]. In addition, thrombin-mediated generation of C5a has been demonstrated to take place in C3-knockout mice, which cannot form C5 convertases and thus are unable to activate C5 by conventional mechanisms [17].

A reciprocal connection in which complement activation would lead to coagulation activation, has also been described in the case of C5a-mediated upregulation of tissue factor (TF), the potent initiator of the extrinsic pathway (= the TF pathway) of coagulation, on both endothelial cells [18] and circulating polymorphonuclear leukocytes (PMNs) [19]. Furthermore, it has been demonstrated that complement activation occurring in vivo during the hemodialysis of patients with end-stage renal disease leads to the generation of C5a and expression of functionally active TF on PMNs, thereby resulting in a procoagulative state that may contribute to the increased risk of thrombosis in these patients [20].

Platelet activation during thrombotic events is intimately associated with the activation of complement and the contact system, which in turn leads to inflammation. Chondroitin sulfate A (CS-A), released from alpha granules during platelet activation, is a potent mediator of crosstalk between platelets and the complement system. Thrombin receptor activated platelets are strong promoters of inflammation since the released CS-A activates complement in the fluid phase and generates anaphylatoxins that induce leukocyte activation [21–23]. In addition, platelet activation leads to the activation of the contact system enzymes FXⅩⅤa and FXⅩla, which are specifically inhibited by antithrombin (AT) rather than by C1INH, as is the case when contact activation is induced by material surfaces [24,25].

2. Biomaterials

2.1. Biocompatibility

The term “biocompatibility” refers to the “ability of a material to perform with an appropriate host response in a specific application” [26]. Most biomaterials come in contact with whole blood, either continuously or during implantation. Consequently, they will be exposed to and identified by the recognition molecules of the different cascade systems: C1q, mannos-binding lectin (MLB), and properdin of the complement system; FXⅩ and high molecular weight kinogen (HMWK) of the contact activation system, and FⅩⅦ and TF of the coagulation system.

This initial contact leads to the generation of potent mediators: the anaphylatoxins C3a and C5a, and the lytic sc5b-9 complex (complement system), bradykinin (contact activation system), and thrombin (coagulation system). These mediators trigger leukocytes (PMNs and monocytes) and platelets, leading to inflammatory and thrombotic
reactions. The processes that may manifest locally and directed against the biomaterial, or in severe cases, systemically and cause whole body inflammation that may be detrimental or even fatal to the patient (Fig. 1).

2.2. Yesterday's biomaterials

Few inventions have shaped medicine in such a dramatic way as biomaterials, dating back to the use of glass eyes [3,27] and the application of gold in dentistry 2000 years ago [28]. A first revolution in the evolution of biomaterials was triggered by the advent of synthetic polymers in the early 20th century, which allowed reproducible manufacturing of materials with distinct characteristics. While originally adapted for medical applications from other sources (e.g., textiles, commodity plastics), it became clear that such polymers have to be carefully tailored to optimize their performance. Many of these early successes were the result of serendipity rather than design. Engineered implants employing common and material borrowed from other fields, developed through collaborations of physicians and engineers, have taken advantage of advances in materials science (albeit from other fields). Commonly used biomaterials from this era include dacron and parachute cloth for vascular implants, titanium alloy for dental and orthopedic implants, cobalt–chromium–molybdenum for orthopedic implants, and ultrahigh molecular weight (UHMW) polyethylene bearing surfaces for total joint replacements, heart valves, and pacemakers.

A point of interest is that one of the first clinical complications reported of complement activation on biomaterials surface was made in relation with hemodialysis using cellulose-derivatized membranes. The relatively large areas of membranes caused a massive complement activation leading to increased levels of C3a and C5a in patient blood and granulocyte aggregation [29]. In another early report, other extracorporeal treatment procedures such as nylon fiber filtration leukapheresis, were found to induce profound complement activation [30]. Since then, much more careful selections of blood contacting polymers have been made.

2.3. Today's biomaterials

Concurrent with the progress occurring in materials science, rapid developments in molecular biology, micro-manufacturing, and nanotechnology have produced a second, ongoing revolution that has introduced biomaterials to hitherto-unimagined fields. Today, we define biomaterials as any "substances other than food and drugs contained in therapeutic or diagnostic systems that are in contact with tissue or biological fluids" [27]. Their application ranges from drug delivery systems and implantable devices (e.g., insulin pumps) to extracorporeal circuits used during cardio-pulmonary bypass surgery [3,27,31]. Today's biomaterials consist of bioengineered implants using bioengineered materials and some modified and new polymeric devices, with few examples on the market but many under development. Cutting-edge examples include tissue-engineered implants designed to re-grow rather than replace tissues, artificial skin, cartilage cell procedures, resorbable bone repair cements, and genetically engineered "biological" components.

Despite this plethora of available biomaterials, their effective use is still challenging because they stimulate both application-directed and potentially adverse reactions by the human body; after all, biomaterials are foreign objects that can induce in vivo defense systems. Given their enormous impact and potential, the urgent need for developing polymers with improved biocompatibility and methods for testing their effects on the body has now been recognized [27]. The elucidation, prevention, and active modulation of adverse reactions mediated by immune and contact-systems are considered highly important aspects of the development of tomorrow's biomaterials.

3. A model of complement activation on a biomaterial surface

During the inevitable exposure of biomaterials to blood, either transiently during administration or implantation or continuously during their lifetime in the body, their artificial surfaces immediately become covered by a film of plasma proteins, that is essentially a monolayer [32]. This adsorption can be described as a recognition phase, a passive process that may or may not include conformational changes in individual proteins within this layer (Fig. 2A,B). Examples of proteins that appear particularly prone to undergo conformational changes upon binding to such surface layers are complement...
component C3 [33] and IgG [34,35], both of which can induce activation of the complement system on a biomaterial surface via the AP or CP, respectively. Other examples are adsorbed FXII, which triggers contact activation [36], and fibrinogen [37,38], which binds to GPIIb/IIIa on platelets, thereby inducing their activation. Deposition of this primary protein layer triggers the activation of the complement, coagulation, and contact systems in ways that are dependent on the composition and conformation of the adsorbed proteins (Fig. 1). The recognition molecules of these cascade systems may bind to deposited proteins or become incorporated into the initial protein film. Because of the absence of specific regulators on the biomaterial surface and a lack of self-recognition, activation of each cascade system typically leads to a rapid amplification of the respective response.

Continuous complement activation leads to the generation and accumulating deposition of C3b and its degradation fragments (iC3b, C3d) on top of the initially adsorbed protein layer and the concomitant release of anaphylatoxins to an extent that is dependent on the conformational exposure of acceptor sites for C3 (Fig. 2C) [32]. Enhancement through the AP amplification loop (Fig. 2D) may finally lead to a total concealment of the initial protein layer by C3 activation fragments. The generated C3a and C5a anaphylatoxins act as strong chemo-attractants that recruit PMNs and monocytes to the site of biomaterial-induced complement activation. The sequence of events is summarized in Fig. 3. In addition, anaphylatoxins exert strong pro-inflammatory effects, which can lead to acute and/or chronic systemic inflammation. Whereas opsonization of foreign surfaces by C3b and iC3b usually facilitates the phagocytic removal of non-self particles via recognition by complement receptors such as CR3 (CD11b/CD18) on activated leukocytes, the large size of biomaterial devices for clinical use often prevents such uptake; consequently, the unresolved activation may lead to a change in the status of recruited immune cells, as has been exemplified by the increased fusion of macrophages to foreign body giant cells [39].

In addition, the activation of the contact activation/coagulation systems ultimately generates thrombin, which is a powerful platelet activator. The thrombin-activated platelets release CS-A, which activates fluid phase complement, further amplifying the ongoing inflammatory reaction [21,23]. Together with the C5a-mediated induction of TF on PMNs and monocytes [19,20,40], these events lead to a marked increase in coagulopathy. Contact system activation also produces the potent vasoactive peptide bradykinin (Fig. 1). In summary, the complement-related surface osorption and release of anaphylatoxins and the subsequent recruitment and activation of leukocyte populations, in combination with the thrombin-mediated activation of platelets, result in inflammatory and thrombotic reactions that can be detrimental to the biomaterial and/or the patient. Thus, the outcome of a medical treatment is to a large extent determined by the composition and conformation of the layer of plasma proteins that are initially formed on the bio-artificial surface.

4. Impact of matrix surfaces on biomaterial-induced complement activation

Solid surface in general such as polymer surfaces, metal surfaces or ceramic surfaces, adsorb blood or tissue proteins as a general phenomenon [41]. Protein adsorption usually takes place as rapid initial adsorption in the first milliseconds until a monomolecular layer of proteins has been formatted. After the formation of a more or less stable monolayer, the adsorption is inhibited. The patterns of individual proteins adsorbed are influenced by the concentration of the proteins in the biological fluids (e.g. blood) as well as the chemistry of the biomaterial surfaces [42]. Activation of the plasma cascade systems, (complement, coagulation/contact activation) is also initiated by direct protein interactions with the biomaterial surfaces, but in contrast to the rapid initial protein adsorption, the effects of the contact activation systems prolong for a longer time and involve cell engagement at the surface [43].

Factors that affect the amount, composition, and conformation of proteins within the initial layer include the hydrophobicity/hydrophilicity of a surface, as well as its charge and the distribution of charged groups. In general, proteins are more prone to undergo substantial conformational changes when binding to hydrophobic than to hydrophilic surfaces [44]. This higher level of binding then results in a higher packing density of the proteins deposited on hydrophobic surfaces [44]. One protein that has been extensively studied is complement component C3, which is known to undergo profound conformational changes when binding to hydrophobic surfaces, as demonstrated using monoclonal antibodies specific for neo-epitopes in denatured or biologically activated C3 [45,46]. In more recent studies, it has been demonstrated that complement is more readily activated on hydrophobic compared to hydrophilic surfaces [47,48].

Numerous reports have demonstrated that surfaces coated with polyethylene glycol (PEG) generally feature low nonspecific protein adsorption, [49,50] and therefore a decreased activation of the coagulation system and subsequent platelet and cellular adhesion [51]; however, activation of the complement system was still substantial on PEG-coated surfaces [52,53] (Table 1). The protein-repellent nature of PEG has been attributed to numerous factors, including surface hydrophilicity and steric repulsion [54,55]. The molecular weight, and surface density of the polymers chains influence the protein binding, and understanding and modifying these factors is the subject of substantial research worldwide [56–61].

Another, well documented, strategy to design inert or “nonfouling” surfaces with low protein binding involves the conjugation of poly(2-methacryloyloxyethyl phosphorylcholine) e.g. [62,63]. In addition, recently, a surface coated with poly(carboxybetaine acrylamide) was reported to show greatly reduced protein binding when exposed to undiluted human blood serum or plasma [64].
A factor that should be taken into consideration when evaluating the complement-activating potential of an artificial surface is the possibility that complement activation (recognition, convertase assembly, and deposition of C3 fragments) may take place transiently but that these compounds later detach from the surface [43]. This phenomenon has been reported to occur on hydrophilic surfaces [43]. Consequently, such surfaces would appear as low complement activators when the evaluation is restricted to fragments on the material surface, yet may indeed be strong activators regarding fluid-phase activation products such as C3a/C5a and sC5b-9. In contrast, a heavily charged surface can induce substantial complement activation but subsequently adsorb the highly cationic compounds C3a and C5a thereby reducing their levels in the circulation; thus, the activation potential of such a material would be underestimated if these activation markers were to be measured only in the fluid phase. In such a case, a more accurate estimate would be obtained by eluting the anaphylatoxins from the material surface [65]. A comparatively high-level binding of complement-initiators such as IgG and C1q can be counteracted by a simultaneous high-level binding of inhibitors such as C1INH, resulting in a lower activation than on a surface with binding identically high amounts of activators but lower amounts of inhibitors [65]. The binding and activating properties of the original surface can also be further modified by low-energy plasma treatment [46,66]. A recent example is a study by Andersen et al., who demonstrated that the modification of a medical device consisting of silicone rubber with plasma-polymerized vinyl pyrrolidone (pPV) coating can strongly decrease the surface activation of the blood complement system [67] (Table 1).

### 5. Controlling biomaterial-induced complement activation

#### 5.1. Active shielding (binding intact biomolecules)

Heparin coatings have been extensively used to render biomaterials blood-compatible, with regard to coagulation, contact system, and complement activation (Fig. 4A, left). The accepted hypothesis is that the inhibition is achieved by acquiring regulators such as AT and Factor H through direct binding to heparin coated surfaces [68]. However, the coagulation system is inhibited by heparin at much lower concentrations than is the complement system [69], so the concentration of surface-bound heparin when optimized to inhibit activation of the coagulation system causes insufficient inhibition of complement (Table 1). Complement activation may be further attenuated by higher heparin surface concentrations, but this effect is not the result of increased binding of Factor H; thus, it appears that Factor H is not the only, or even the main, regulator of complement on heparin surfaces [70]. Furthermore, since complement activation at surfaces is far from obliterated, it is clear that improved and more specific methods must be developed to inhibit complement activation on biomaterials [70–73]. In addition, heparin is known to interact with a plethora of plasma proteins and may cause undesired effects because of its “broadband” specificity.

Conjugation of a surface with biologically active, naturally occurring RCAs is a potential approach to lowering the complement activation on the material surfaces. This concept has been tested by covalently binding human purified Factor H to a model biomaterial, polystyrene, using two different linkers, N-succinimidyl 3-(2-pyridyldithio) propionate (SPDP) and PEG (Fig. 4A, middle, right). In the case of both surfaces, Factor H, when bound in native form, as determined by quartz crystal balance with dissipation (QCM-D), was able to completely abrogate complement activation upon exposure to human blood [52,74]. Offering a promising alternative to the use of immunosuppressive and cytotoxic drugs on stents, the Factor H–PEG surface is currently being prepared for human clinical trials to prevent inflammation, coagulation, and vascular tissue damage (Table 1).

#### 5.2. Autoprotection

Surfaces lacking regulators or self-recognition patterns (microbial intruders as well as artificial biomaterials) will likely trigger amplification of complement response. As described in Section 5.1, heparin coating of biomaterials with the aim to actively adsorb Factor H and thereby inhibit complement activation shows varying results, in some cases insufficient efficacy [68,70,71,75]. A more targeted approach that selectively recruits Factor H is therefore considered.

![Fig. 4. Examples of surface modification strategies designed to cope with innate immunity-related recognition of biomaterial surfaces.](image-url)

**Table 1** Examples of surface modification procedures, effects on the hemocompatibility, and examples of biomedical devices with each surface modification.

<table>
<thead>
<tr>
<th>PEG a</th>
<th>Low</th>
<th>Low</th>
<th>Varying</th>
<th>Low</th>
<th>Experimental (drug delivery systems b)</th>
</tr>
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<tbody>
<tr>
<td>Low energy plasma</td>
<td>Varying</td>
<td>–</td>
<td>Moderate</td>
<td>Low</td>
<td>Experimental (e.g. oxygenators), central venous catheters, stents</td>
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<tr>
<td>Heparin</td>
<td>Low</td>
<td>–</td>
<td>–</td>
<td>Low</td>
<td>Stents</td>
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<td>RCA c</td>
<td>–</td>
<td>–</td>
<td>Low</td>
<td>–</td>
<td>Experimental</td>
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<tr>
<td>Anti-RCA peptide</td>
<td>–</td>
<td>–</td>
<td>Low</td>
<td>–</td>
<td>Drug eluting contact lenses</td>
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<tr>
<td>Molecular imprint</td>
<td>–</td>
<td>–</td>
<td>Low</td>
<td>–</td>
<td>Dental titanium implants</td>
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<tr>
<td>Nanostructure</td>
<td>–</td>
<td>–</td>
<td>Low</td>
<td>Low</td>
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a Polyethylene glycol.  
b Discussed in Chapter 7 in this issue of Advanced Drug Delivery Reviews.  
c Little or no data regarding systematic investigation of these parameters is available.

d Regulators of complement activation.

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important for studying complement-induced effects but may also directly lead to clinically applicable products. An obvious choice would be to utilize antibodies (either intact, or Fab, or single chain fragments) to target surfaces (Fig. 4B, left). However, such coatings with large natural structures would be rather complex and costly to produce and may therefore be difficult to translate into clinical applications. Alternative sources of regulator-recruiting entities are therefore desired.

Intriguingly, some human pathogens expose RCA-capturing molecules as part of their immune evasion strategy [76]; for example, M proteins of Streptococcus pyogenes bind C4BP [77-79], Neisseria meningitidis expresses a Factor H-binding protein that mimics glycan patterns [80-82], and staphylococcal proteins (Efb, Sbl) have recently been implicated in the enhancement of Factor H binding [83,84]. Whereas coating with such capturing intact proteins would appear less feasible because of size and immunogenicity concerns, M protein-derived peptides have recently been shown to recruit C4BP to model polystyrene surfaces and reduce complement activation [85] (Fig. 4B, middle).

Despite these promising proof-of-principle studies, those M protein-derived peptides do not lend themselves to cost-effective synthesis because they are still some 50 amino acids long. Also, it is imperative to target the AP amplification loop, preferably by exploiting its main inhibitor Factor H, because this pathway plays a pivotal role in biomaterial-induced complement activation, as illustrated by the fact that it contributes >80% of the C5 activation in a model in which complement activation is initiated by the CP [86].

With the aim of developing a small-molecule, Factor H-specific capturing compound, we recently screened variable cysteine-constrained phage-displayed peptide libraries and identified several small peptides with high Factor H-capturing activity. One of these peptides was found to retain the functional integrity of factor H by capturing at a non-regulatory area of the regulator. Indeed, coating of a model biomaterial (polystyrene) with this peptide recruited high levels of factor H efficiently inhibit complement activation by the AP (Fig. 4B, middle) [87]. Such peptides are therefore evaluated and optimized for potential clinical applications (Table 1).

5.3. Molecular imprinting

The use of functional synthetic materials with a predetermined molecular-level structure as biomaterials is a fascinating possibility and offers a potential alternative to approaches based upon immobilized biomolecules, with their inherent instability. Two fundamentally different strategies to achieve this end are the topic of current research: The first involves the use of molecular imprinting techniques (MIPs) for developing autoregulatory biomaterial surfaces that provide a surface pattern capable of selective recruiting RCAs (Fig. 4B, right). The second strategy involves the use of nanostructured surfaces with dimensions that produce steric constraints, thereby inhibiting cascade recognition events (see Section 5.4 and Fig. 4C).

The MIP approach [88] involves the formation of cavities in a synthetic polymer matrix that are structurally and functionally complementary to a template molecule/entity. The ability of MIPs to selectively recognize and bind a template structure in the presence of closely related chemical species has led to attempts to apply them to a wide range of biomedical and biotechnological applications. These antibody combining-site mimics have demonstrated binding affinities and cross-reactivity profiles comparable to their biological counterparts and have even been employed as substitutes for biological antibodies in clinical diagnostic assays [89]. While the molecular imprinting process in many ways parallels the immune system’s production of antibodies, it also exhibits significant contrasting features and offers certain advantages, for example, a lack of hapten-conjugation protocols and situations involving non-immunogenic substances pose no problem. Moreover, materials formed using MIPs are stable under extreme conditions of pH, temperature, and organic solvent exposure, quite unlike their biological counterparts [90]. Accordingly, these materials have great potential for basic research and for the biotechnology and pharmaceutical industries as potential biomaterials. Indeed, recent efforts aimed at developing molecularly imprinted contact lenses as vehicles for the controlled release of medication [91], e.g., beta blockers such as timolol for the treatment of glaucoma, have proven useful [92] (Table 1).

The first reports of imprinted polymer surfaces, being developed as biomaterials, have recently appeared. Generally, the template surfaces are prepared by attaching specific biomolecules of interest to a glass surface (the template surface). The template surface is then placed in contact with a suitably derivatized surface, and a water-soluble monomer mixture, e.g., one based on acrylates or suitable acrylamides [93], is allowed to polymerize between the two surfaces. After separation of the imprinted surface from the template, acceptor sites are revealed on the imprint surface (Fig. 4B, right). We have recently demonstrated that prototype imprints of heparin are significantly less prone to activate complement than are control polymer surfaces [94].

5.4. Nanostructure

The synthesis of nanostructured surfaces for applications in tissue engineering and drug delivery is a rapidly emerging field [95-97]. In a recent study, Ferraz et al. demonstrated that complement activation on aluminum surfaces is strictly dependent on pore diameter; they found that activation on surfaces with a pore diameter of 20 nm was significantly lower than on surfaces with 200 nm pores, despite the fact that the total surface of the former material was much greater [98]. In order for complement activation to occur, a number of large proteins need to interact at a surface; that is, the subunits of the AP C3 convertase (C3bBb) need to associate with and cleave nearby C3 molecules. Structures that are too narrow may not allow for such interactions to take place at normal rates (Fig. 4C, left). In contrast, on surfaces with more expansive structure, convertase assembly and cleavage will take place both inside and between the pores; in this case, the effect by introducing the nanostructure is to increase the accessible surface area (Fig. 4D, right; Table 1).

Similarly, in a recent study, it was demonstrated that synthetic polymers with a similar charge but different pore sizes differ in terms of their protein binding and complement activation, with lower values being obtained for polymers with small pores than for those with larger ones [65].

5.5. Soluble inhibitors

In addition to modifying biomaterials to shield them from unwanted reactions, it is possible to use soluble complement-targeting drugs to suppress the amplification of the complement response. This approach might be particularly suitable in situations with repetitive yet temporally limited periods of exposure to artificial surfaces (e.g., hemodialysis) or at the initial stages of implantation, to allow better embedding. A variety of complement-targeted drugs are currently on the market or in clinical or pre-clinical trials [99,100]. The first complement inhibitor to be licensed as an orphan drug, was eculizumab, a humanized anti-C5 antibody which is used for treatment of the rare disorder paroxysmal nocturnal hemoglobinuria [101]. However, in view of the key involvement of the AP in biomaterial-induced complement activation, inhibition at the level of C3 appears to be most promising. RCA-based therapeutics have been of recent interest as a result of the development of chimeras between the regulatory domains of Factor H and the C3d-binding domains of CR2 [102] for use in targeting regulators to sites of activation. However, the cost of such protein therapeutics is rather high, and these chimeras only target the AP amplification step.
In contrast, Compstatin is a peptidic complement inhibitor that interacts with C3/C5b and thereby inhibits the activation of both the CP/ LP and AP; the safety of this compound has recently been established in a Phase I clinical trial for age-related macular degeneration, and it has also been successfully used in a variety of disease models [103]. Most importantly, Compstatin analogs have recently been tested in a model of hemodialysis and found to efficiently suppress the filter-induced activation of complement and neutrophils and to reduce the expression of TF [20]. The use of novel Compstatin analogs with increased inhibitory potency for biomaterial-related applications is therefore considered promising [104].

6. Conclusions

Overall, considerably more success has been achieved in reducing the thrombogenicity of bio-artificial surfaces than in controlling complement activation: For example, surfaces coated with different forms of heparin or PEG are associated with low or negligible activation of coagulation and subsequent platelet loss. Thus, there are numerous surfaces that have low thrombogenicity available that still bear substantial complement-activating capacity. Some of these materials will no doubt be a suitable starting material for exploring the various modification procedures for disarming complement activation that have been described in Sections 5.2–5.4 and summarized in Table 1. Table 1 also gives examples of surface modification procedures used in different medical devices. The ultimate goal of this approach is to create a hybrid surface that combines the inherent coagulation-inert properties of the original surface with specific complement-autoregulation, thus minimizing the risk of short-circuiting of the systems via platelet derived Cs-A, CsA and TF as discussed in Section 3. Such a biomaterial would show superior blood compatibility and fewer detrimental side effects (with regard to both the patient and the biomaterial), as compared to the materials available in the clinic today.

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