Therapeutic Targeting of the Complement System: From Rare Diseases to Pandemics

Peter Garred, Andrea J. Tenner, and Tom E. Mollnes

Laboratory of Molecular Medicine, Department of Clinical Immunology, Rigshospitalet, Copenhagen, Denmark, and Faculty of Health and Medical Sciences, University of Copenhagen, Copenhagen, Denmark (P.G.); Departments of Molecular Biology and Biochemistry, Neurobiology and Behavior, and Pathology and Laboratory Medicine, University of California, Irvine, California (A.J.T.); and Research Laboratory, Nordland Hospital, Bodø, Norway, Faculty of Health Sciences, K.G. Jebsen TREC, University of Tromsø, Tromsø, Norway (T.E.M.); Centre of Molecular Inflammation Research, Norwegian University of Science and Technology, Trondheim, Norway (T.E.M.); and Department of Immunology, Oslo University Hospital and University of Oslo, Oslo, Norway (T.E.M.)

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Address correspondence to: Tom E. Mollnes, Research Laboratory, Nordland Hospital, N-8092 Bodø, Norway. E-mail: t.e.mollnes@gmail.com

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ABBREVIATIONS: AAV, anti-neutrophilic cytoplasmic antibodies–associated vasculitis; aHUS, atypical HUS; AMD, age-related macular degeneration; APS, antiphospholipid syndrome; CAPS, catastrophic APS; C3aR, C3a receptor; C5aR1, C5a receptor 1; C5aR2, C5a receptor 2; C4BP, C4b-binding protein; C3G, C3 glomerulopathy; C3GN, C3 glomerulonephritis; CL, collectin; CNS, central nervous system; COVID-19, coronavirus disease 2019; CR, complement receptor; CRP, C-reactive protein; DAF, decay-accelerating factor; DAMP, damage-associated molecular pattern; DDD, dense deposit disease; Fc, fragment crystallizable; FDA, Food and Drug Administration; Gd-IgA1, galactose-deficient IgA1; GPI, glycosyl phosphatidylinositol; HUS, hemolytic-uremic syndrome; iC3b, inactive C3b; IL, interleukin; mAb, monoclonal antibody; MAP-1, mannose-binding lectin/ficolin/collectin associated protein 1; MASP, MBL-associated serine protease; MBL, mannose-binding lectin; 3MC syndrome, Malpuech, Carnevale, Michels, and Mingarelli syndrome; MCP, membrane cofactor protein; MS, multiple sclerosis; NeF, nephritic factor; NLRP3, nucleotide-binding oligomerization domain-like receptor 3; PAMP, pathogen-associated molecular pattern; PAR, protease-activated receptor; PNH, paroxysmal nocturnal hemoglobinuria; PTX3, pentraxin 3; sC5b-9, soluble C5b-9; sCR1, soluble complement receptor 1; SLE, systemic lupus erythematosus; SNP, single-nucleotide polymorphism; SP, surfactant protein; TCC, terminal complement complex; TLR, Toll-like receptor; VEGF, vascular endothelial growth factor.
**Abstract**—The complement system was discovered at the end of the 19th century as a heat-labile plasma component that “complemented” the antibodies in killing microbes, hence the name “complement.” Complement is also part of the innate immune system, protecting the host by recognition of pathogen-associated molecular patterns. However, complement is multifunctional far beyond infectious defense. It contributes to organ development, such as sculpting neuron synapses, promoting tissue regeneration and repair, and rapidly engaging and synergizing with a number of processes, including hemostasis leading to thromboinflammation. Complement is a double-edged sword. Although it usually protects the host, it may cause tissue damage when dysregulated or overactivated, such as in the systemic inflammatory reaction seen in trauma and sepsis and severe coronavirus disease 2019 (COVID-19). Damage-associated molecular patterns generated during ischemia-reperfusion injuries (myocardial infarction, stroke, and transplant dysfunction) and in chronic neurologic and rheumatic disease activate complement, thereby increasing damaging inflammation. Despite the long list of diseases with potential for ameliorating complement modulation, only a few rare diseases are approved for clinical treatment targeting complement. Those currently being efficiently treated include paroxysmal nocturnal hemoglobinuria, atypical hemolytic-uremic syndrome, myasthenia gravis, and neuromyelitis optica spectrum disorders. Rare diseases, unfortunately, preclude robust clinical trials. The increasing evidence for complement as a pathogenetic driver in many more common diseases suggests an opportunity for future complement therapy, which, however, requires robust clinical trials; one ongoing example is COVID-19 disease. The current review aims to discuss complement in disease pathogenesis and discuss future pharmacological strategies to treat these diseases with complement-targeted therapies.

**Significance Statement**—The complement system is the host’s defense friend by protecting it from invading pathogens, promoting tissue repair, and maintaining homeostasis. Complement is a double-edged sword, since when dysregulated or overactivated it becomes the host’s enemy, leading to tissue damage, organ failure, and, in worst case, death. A number of acute and chronic diseases are candidates for pharmacological treatment to avoid complement-dependent damage, ranging from the well established treatment for rare diseases to possible future treatment of large patient groups like the pandemic coronavirus disease 2019.

## I. Introduction

The complement system has been essentially ignored and/or unknown in the clinic, and treatment of diseases with dysfunction of complement has been limited. There are several reasons for this. First, the willingness to consider this cascade as a component of disease initiation or progression has been virtually absent since most clinicians find it too complicated. Second, the relevance of the system in human diseases has been unclear. Third, lack of drugs has been a limitation for conducting clinical trials. The status for the complement system as late as in the 1980s was described in a summary after the Complement Meeting in the Royal Society in London in 1984, published in Immunology Today (now Trends in Immunology). Mike Hobart (1984) concluded, “Many immunologists hold that complement is baffling or irrelevant or, most conveniently, both, but a recent meeting emphasized that complement is interesting and that it may be important, even only as an elegant model system.”

In recent years there has been a “clinical complement revolution,” and we have learned that complement indeed is important, and although it is an elegant biochemical system, it is more than that. Modulation of the system has already come to clinical use, and the medical community would see gains in patient outcomes with a better awareness and understanding of the complement system (Mastellos et al., 2017; Ricklin et al., 2017, 2018; Gialeli et al., 2018; Harris et al., 2018; Tomlinson and Thurman, 2018; Wong and Kavanagh, 2018; Bordron et al., 2020). The hemostatic system with the coagulation and fibrinolytic cascades is well known with the consequences of bleeding or thrombosis when deficiency or dysregulation occurs. The complement cascade works along with the same principles, and phylogenetically the plasma cascades originated from a common ancestor.

In this review we aim to give an introduction to the complement system, including its primary functions. However, complement dysfunction and how we can approach diseases wherein complement participates in the pathogenesis will be highlighted. Thus, complement is a double-edged sword from being our protective friend on the one side to our enemy in the case that it gets out of control and does harm to the host. A thorough understanding of the interacting parts will pave the way for future successful pharmacological approaches to suppress excessive complement-mediated detrimental processes while not inducing long-term suppression of the beneficial effects of complement. This review will not cover a detailed list of ongoing trials and all possible drugs to be developed but rather highlight the general approaches on how to meet the challenges and alternatives of when and how to treat complement dysfunction. For those interested in a complete overview of ongoing trials and the corresponding drugs we recommend using https://clinicaltrials.gov/ and “complement” as search key. This list is complete and continuously updated in contrast to tables printed in articles that are recently outdated.

### A. An Overview of the Complement System

#### 1. Discovery of the System

The complement system was discovered during the 1880s–1890s by several
scientists, including Paul Ehrlich, George Nuttall, Hans Buchner, and, notably, Jules Bordet (Buchner, 1891; Ehrlich, 1899; Skarnes and Watson, 1957; Nesargikar et al., 2012). It was discovered as a heat-labile factor in serum that contributed to bacterial killing and first named “alexin.” Later it was termed “complement” based on Bordet’s classic experiments from 1895 in which he distinguished two factors responsible for serum killing of a Gram negative bacterium: one was a heat-stable factor representing the antibodies against...
The emergence of the more rapid DNA-sequencing techniques in the 1980s, advances in NMR and X-ray crystallography at the turn of the century, and, later, the use of cryogenic electron microscopy enabled further molecular understanding of this multiple-component recognition of danger and fully capable of inducing lytic activity was not fully appreciated until the second half of the 20th century. The biochemistry of C3 (Muller-Eberhard et al., 1960) and the three initial pathways (Pillemer et al., 1954; Pillemer, 1955; Lepow et al., 1963; Daha et al., 1976), the assembly of the common terminal C5b-9 complex, and the detection of C3 in injured tissue (Lachmann et al., 1962) were largely defined at that time because of the creative use and sometimes necessary brute force of technological advances in protein chemistry (Reid et al., 1972), biochemistry, and monoclonal antibody technology, which are briefly chronicled in Sim et al. (2016). The nature of the membrane attack complex (Podack et al., 1980) was intensively studied, and after 3 decades it is still debated whether it acts by leaky patches (Esser, 1991) or by a physical hole (Bhakdi and Tranum-Jensen, 1991). The biochemical identification and functional characterization of many of the complement regulator proteins (Lachmann and Muller-Eberhard, 1968) and receptors (Fearon, 1980), the use of animal models for in vivo mechanistic studies, and the identification of individuals genetically deficient in these components drove a plethora of novel insights on the critical importance of this system in human health and disease.

B. The Complement System Approach in the Clinic: The Therapeutic Era

1. The Revolution of Complement Research for the Last 3 Decades. It is now appreciated that there are over 40 proteins that can be counted as part of the complement system, including the canonical components, receptors for activation-generated cleavage products, and regulators of the cascade (Fig. 1). The influence of complement fragments (e.g., C3d, C3a, and C5a) on the type and extent of the adaptive response became evident, as reviewed in Luque et al. (2019), Reis et al. (2019), and Lo and Woodruff (2020). Novel complement control proteins are still being discovered, often with tissue-specific expression (Cong et al., 2020), and are reviewed in Gialeli et al. (2018). However, the domain protein structures have similar features (Forneris et al., 2016). The evasion by microbes of complement-mediated killing has demonstrated the constant battle as these pathogens evolved but also provide examples of how such regulation can be harnessed for application in mitigating excesses of complement activation. The consequences of an activated complement system with the vasculature, including but also beyond coagulation, lung tissue, brain, and the gut, are just beginning to be more fully addressed (Huber-Lang et al., 2018).
cascading system. The structures of the globular head domain of C1q and C3 (Gaboriaud et al., 2003; Janssen et al., 2005) and how complement is activated by IgG hexamers (Diebolder et al., 2014) were practically breathtaking in the impact they would have on the field. Combined with other sophisticated protein chemistry approaches, these advances enabled initial rational drug design for these pivotal proteins (Fredslund et al., 2008; Schuster et al., 2008; Skjoedt et al., 2012; Schmidt et al., 2016; Papanastasiou et al., 2017).

The impact that the complement system has on human health and disease and the advantages of precision medicine became even more evident with the recognition that distinct combinations of genetic variants leading to more or less effective activators (such as factor B, C3) and more or less effective inhibitors (such as factor H), which are collectively referred to as a “complotype,” confer differential sensitivity to infection and autoimmunity (Harris et al., 2012). Indeed, the identification of variants of factor H as being the major genetic risk factor in age-related macular degeneration (AMD) in 2005 (Daiger, 2005; Hageman et al., 2005) was a major stimulus in the investigation of complement in the role of neurodegenerative diseases of the elderly. In subsequent investigations, genetic variants in other complement components were also linked to AMD and led to clinical trials of targeted complement functions (Park et al., 2019). Genetic polymorphisms also can critically affect responses to complement drugs, such as that reported for C5 inhibition in the treatment of paroxysmal nocturnal hemoglobinuria (PNH) and discussed below (Nishimura et al., 2014). Even more recently, the identification of intracellular complement activities, the “complosome,” has provided intriguing evidence of far-reaching consequences of complement components on cellular metabolism and, subsequently, on both innate and adaptive immune responses (as reviewed in Kemper and Kohl (2018) and West et al. (2020)). The impact of this intracellular complement system on the complement therapeutic landscape remains to be determined.

Although the liver was long thought to be the site of complement protein production [reviewed in Perlmutter and Colten (1986)], it became increasingly apparent that various complement components can be differentially induced in a variety of cell types and tissues (Minutti et al., 2017; Kemper and Kohl, 2018), including the brain (Singhrao et al., 1999). For example, complement components can be induced in the central nervous system (CNS) resident neurons, astrocytes, oligodendrocytes, cerebrovascular smooth muscle cells, and microglia during development or induced by injury or aging [reviewed in Tenner (2020)]. These findings have been confirmed recently by single-cell RNA sequencing (Zhou et al., 2020).

In tissues, as evident in the central nervous system, there appears to be transcriptional control to the sequence of induction of complement proteins depending on the type and level of signaling received by the cells. C1q can be synthesized in the absence of the C1 serine proteases C1r and C1s in peripheral myeloid cells (Bensa et al., 1983) and is rapidly upregulated in response to injury (Lee et al., 2000). The addition of interferon-γ, perhaps as a signal of increased damage, was required to detect C1r and C1s production by macrophages (Bensa et al., 1983). It then became clear that the C1q molecule itself independent of C1 or other complement activities contributed to critical functions for both homeostasis, such as clearance of apoptotic cells, neuronal blebs, and cellular debris, and sculpting the adaptive immune system through induction of cytokines in myeloid cells that could subsequently facilitate a limit on adaptive responses [e.g., to self peptides ingested by phagocytes, as reviewed in Thielen et al. (2017)].

One of the most unexpected and paradigm-shifting discoveries in the past 2 decades has been the identification of the role of the classical complement pathway components C1 through C3 in synapse pruning (refinement) in the nervous system. Although this was first reported as mediating the ingestion of weak or unnecessary synapses during the development of retinal circuits (Stevens et al., 2007), it is now known to contribute to synaptic plasticity in adult brain, including most recently the process of forgetting (Wang et al., 2020). This pruning involves microglial engulfment of inactive C3b (iC3b)-tagged synaptic material (Stevens et al., 2007; Hong et al., 2016; Dejanovic et al., 2018) via the complement receptor (CR) 3 (Schafer et al., 2012). In addition, several groups have provided evidence for detrimental complement-mediated synapse pruning in animal models of aging, Alzheimer disease, multiple sclerosis, and other disorders that display cognitive or behavioral impairments (Stevens et al., 2007; Hong et al., 2016; Lui et al., 2016; Sekar et al., 2016; Vasek et al., 2016; Werneburg et al., 2020) and more recently in motor neuron disease (Vukojeic et al., 2019). Importantly for translation to the human condition, the accumulation of C1q/C3b-tagged synapses and decreased synaptic density are also found in multiple human disorders, such as tauopathies, Alzheimer disease, and West Nile Virus–induced cognitive loss (Lui et al., 2016; Vasek et al., 2016; Wu and Sun, 2019). Surface-exposed phosphatidyl serine or annexin V (Györffy et al., 2018), decreased mitochondrial functions (Györffy et al., 2020), and loss of CD47 (Lehrman et al., 2016) have been associated with enhanced C1q binding and synapse engulfment in developmental or injury models similar to characteristics influencing the clearance of apoptotic cells.

Awareness and consideration of the above homeostatic, neuronal, and other reparative functions (Bossi et al., 2014; Thielen et al., 2017) are critical when selecting a target to pharmacologically inhibit harmful
downstream complement activities, particularly in the case of chronic versus acute treatment.

2. Complement from Bench to Bedside: Today’s Indications. Currently, when this review was written, there are four diseases for which complement-targeted drugs are approved for routine clinical use by the US Food and Drug Administration (FDA) and the European Medicines Agency. The “complement revolution” into the clinic started when eculizumab (Soliris), a mAb-blocking cleavage of C5, was shown to be very efficient in protecting from hemolysis in PNH in 2007 (Brodsky et al., 2008; Parker, 2009), and this was later followed by treatment of atypical hemolytic-uremic syndrome (aHUS) (Tschemi et al., 2011; Loirat et al., 2016). Recently it was also approved for the neurologic diseases generalized myasthenia gravis (Dhillon, 2018) and neuromyelitis optica spectrum disorders (Selmaj and Selmaj, 2019). For details of these rare diseases, see separate sections below.

Eculizumab has been the only approved complement inhibitor for routine use until recently when ravulizumab (Ultomiris) and Zilucoplan were introduced. Ravulizumab is the second generation of eculizumab; in other words, some minor amino acid modifications were made in the eculizumab molecule, thereby increasing the half-life substantially so treatment intervals could be increased from every 2nd to every 8th week. It has the same binding site to C5 as eculizumab (i.e., blocking its cleavage and thereby preventing the release of C5a and formation of C5b-9) (Kulasekararaj et al., 2019; Lee et al., 2019; McKeage, 2019; Stern and Connell, 2019; Lee and Kulasekararaj, 2020).

Zilucoplan is an entirely different drug structurally from the antibody mentioned above but with the same principal function. It is a synthetic, macrocyclic peptide inhibitor for subcutaneous self-administration with principally the same function as eculizumab, blocking the cleavage of C5 (Beecher et al., 2019; Albazli et al., 2020; Howard et al., 2020). A phase 2 randomized, double blind, placebo-controlled, multicenter clinical trial demonstrated that Zilucoplan administration yielded rapid, meaningful, and sustained improvements over 12 weeks in a broad population of patients with moderate-to-severe acetylcholine-receptor-antibody–positive generalized myasthenia gravis (Howard et al., 2020). A phase 3 trial (Recovery After an Initial Schizophrenia Episode study) investigating the safety, tolerability, and efficacy of Zilucoplan in subjects with generalized myasthenia gravis (https://clinicaltrials.gov/ct2/show/NCT04115293) is currently ongoing. RA 101295, a close analog of Zilucoplan (RA101495), has been used in animal studies and has been shown to increase survival in baboon Escherichia coli sepsis (Keshari et al., 2017).

Below we refer to different complement therapeutic drug groups under development and in trials. We emphasize, however, that C1-inhibitor, a regulator of the classical and lectin complement pathways, frequently is listed among therapeutic complement inhibitors. However, it should not be regarded as a specific complement drug since it is a broad serine protease inhibitor with regulatory functions in several plasma cascade systems. Thus, patients deficient in C1-inhibitor do not have a pure complement pathophysiological phenotype but rather a disturbance in the bradykinin system. Still, it is included when the complement system is discussed because the complement laboratories do the diagnostics. The indication for use should be conditions with “pan-cascade” disturbances and not specific complement dysregulation.

II. Complement Deficiencies and Loss- or Gain-of-Function Mutations

Complement deficiencies can be genetic or acquired (Sjöholm et al., 2006; Botto et al., 2009; Skattum et al., 2011). Homozygous genetic deficiencies generally result in undetectable protein levels or function in plasma, whereas heterozygous deficiencies typically present with approximately half of normal levels. Without genetic analysis, they can be difficult to discriminate from an acquired deficiency, which can present with any level of decreased concentration, although very rarely a complete absence. Acquired deficiencies often affect several components. Most frequently they are caused by increased consumption due to in vivo activation and thus consumption of the components or due to liver failure, as most of the blood complement components are produced in the liver. A valuable tool to distinguish between these causes of reduced components is to measure complement activation products (Harboe et al., 2011). These will normally be elevated during in vivo activation and low, normal, or decreased when synthesis is reduced, such as by liver damage. Screening for complement deficiencies, particularly for total genetic deficiencies, is performed by hemolytic assays, which gradually have been replaced by ELISA assays (e.g., the Wieslab Total Complement Screen assay in which all the complement pathways are screened for) (Seelen et al., 2005). The principles and interpretation of these assays will be described below (see section VI and Fig. 7).

Genetic mutations with the protein present may be loss-of-function or gain-of-function mutations, whereas the synthetic rate and concentration of the protein in plasma can be normal. If the mutation leads to activation and consumption of certain proteins (e.g., C3), the serum concentration is frequently reduced. Some functional assays exist for certain proteins, but today genetic tests are used for identification of clinically relevant mutations.

A. Classical Pathway Deficiencies

1. The Components of the Classical Pathway, Their Activation, and Regulation. The classic pathway is
shown in the upper left part of Fig. 1. The classical pathway consists of the trimolecular protein complex C1qRβ2, C4, C2, C1-inhibitor, and C4b-binding protein (C4BP). C1q is normally activated by binding to the Fc part of antibodies after they bind to its antigen. However, antibodies are not the only targets for C1q. Pentraxins like C-reactive protein (CRP), long pentraxin 3 (PTX3), serum amyloid P, and fibrillar amyloid P can also bind and activate C1q in an antibody-independent manner (McGrath et al., 2006; Biró et al., 2007; Du Clos and Mold, 2011; Doni et al., 2012). C1r and C1s are proenzyme serine proteases triggered by C1q bound to an activator to be activated and cleave C4 and subsequently C2. C1qRβ2 has catalytic autoactivity, which is kept under strict control by C1-inhibitor (Ziccardi, 1982). C4BP is the second regulator in the classical pathway. It serves a cofactor for factor I and thereby controls and inactivates C4b (Sjöberg et al., 2009) (Fig. 1).

Receptors for classical pathway components and activation products are mainly limited to the different C1q receptors (Ghebrehiwet and Peerschke, 2014). It remains to be elucidated whether these are therapeutic targets. No specific receptor for C4a has been described, but protease-activated receptors (PARs) 1 and 4 were recently shown to bind C4a (Wang et al., 2017). PARs are a subfamily of related G protein–coupled receptors that are activated by cleavage of part of their extracellular domain.

2. Deficiencies of the Classical Pathway Components. Deficiencies of the classical pathway are relatively rare; the phenotype varies from being healthy to presenting with a severe disease, which is usually autoimmune or infectious (Truedsson, 2015). Low C1q may be genetic or due to consumption, as seen in immune-complex diseases and acquired angioedema. Genetic deficiencies are highly associated with systemic lupus erythematosus (SLE)-like disease, which might be severe particularly when affecting the kidneys. Most of the genetic C1-deficient individuals develop autoimmune disease (Kirschfink et al., 1993), but it should be noted that only a very small percentage of patients with SLE have genetic C1 deficiency because it is extremely rare.

C4 deficiency may have a similar phenotype as C1 deficiencies, but it is less severe. C4 has two isoforms encoded from separate genes (C4A and C4B) and up to six copies per genome (Yang et al., 2007). The C4 isoforms are very homologous and differ with only four amino acids in their sequence. Deficiency of the isoforms is very common and may be seen in between 1% and 10% (Szilágyi and Fust, 2008). However, total C4 deficiency due to lack of both C4A and C4B is extremely rare.

Total C2 deficiency is the common classical component deficiency (frequency 1:10–20000). Half of the individuals are healthy, but some develop severe infections or autoimmune disease (Trapp et al., 1987; Yang et al., 2007). Complete deficiency of C4BP has so far not been described in humans, but nonsynonymous alterations in its sequence have been found in hemolytic-uremic syndrome and recurrent pregnancy loss (Ermert and Blom, 2016).

3. Disturbances of the Classical Pathway and Therapeutic Targets. Substitution therapy exists for C1-inhibitor deficiency with several purified C1-inhibitor concentrates available. However, bradykinin-receptor antagonists like icatibant have gradually replaced C1-inhibitor. There are no other purified components available for clinical therapy in the classical pathway nor in the rest of the complement system. Plasma infusions have occasionally been used as the source for a defective of the deficient component. For example, one patient who was C2-deficient with SLE affecting the kidneys was treated for years with plasma infusions that apparently had a good effect (Steinsson et al., 1989). However, plasma infusions are not typically used in complement deficiencies. It contains proteins from the whole plasma cascade and enzymes as well as regulators, and caution should be taken not to bring “oil to the fire.” Thus, diseases related to classical pathway dysfunction are treated according to symptoms: immunosuppressive drugs for autoimmune diseases, vaccines and antibiotics in case of infections, and general care to detect early symptoms requiring treatment.

A completely different treatment approach is required when in a normal individual with the classical components intact, and the classical pathway is activated by an external structure. If this activation is fully classical pathway–specific and does not involve initial lectin or alternative pathway, inhibition of the system at the level of C1 would be a rational option. Targets could then be C1q, C1r, or C1s, although chronic indications targeting C1q should be avoided since it has far more biologic functions than activation of the complement system, as explained above. The prototypical example of such a purely classical pathway disease is the cold agglutinin syndrome, wherein IgM autoantibodies lyse red cells purely through a classical complement-mediated mechanism (Berentsen, 2018). Studies using blockers of C1s are discussed below under the disease chapter.

B. Lectin Pathway Deficiencies

1. The Components of the Lectin Pathway, Their Activation, and Regulation. The lectin pathway is shown at the upper middle part of Fig. 1. The lectin pathway consists of two classes of soluble pattern-recognition molecules named “collectins” and “ficolins,” a set of activating enzymes termed “mannose-binding lectin/ficolin/collectin associated serine proteases” (MASPs), and different regulators (Garred et al., 2016). All the recognition molecules in the lectin pathway are composed of a structure to C1q from
classical pathway containing polypeptide chains having a C-terminal globular recognition domain and an N-terminal collagen-like structure assembled into a triple helix structure. The triple helix structure is further assembled into macromolecules with a tulip-like appearance in electron microscopy (Lu et al., 1993). The collectin protein family comprises the prototypic mannose-binding lectin (MBL) (also termed mannan-binding lectin), collectin (CL)-10 (also termed liver 1 or CL-L1), and CL-11 (also termed collectin kidney 1 or CL-K1). CL-10 and CL-11 exist as homomorphic proteins as well as heteromeric complexes between CL-10 and CL-11 polypeptide chains CL10/CL-11 (also called CL-LK) (Henriksen et al., 2013). In addition, the lung surfactant protein (SP)-A and SP-D also belong to the collectin family. However, they do not activate the complement system (Tenner et al., 1989), as reviewed in Murugaiah et al. (2020). The protein CL-12 (also called collectin placent 1 or CL-P1) may be included in the family (Hansen et al., 2016). It is a transmembrane protein that correspondingly is found as a fluid analog. This analog does not activate the lectin pathway but appears to serve as a docking station for the alternative pathway molecule properdin (Ma et al., 2015; Zhang et al., 2020).

The ficolin protein family comprises ficolin-1 (M-ficolin), ficolin-2 (L-ficolin), and ficolin-3 (H-ficolin) (Endo et al., 2015). As for CL-10 and CL-11, ficolin-2 and ficolin-3 may also be found as heterocomplexes in the circulation, but the stoichiometry for these complexes is at present unknown (Jarlhelt et al., 2020). The main functional difference between the collectins and ficolins is defined by the C-terminal globular recognition domain, which for the collectins is a calcium-dependent carbohydrate-binding domain, whereas for the ficolins it is a fibrinogen-like binding domain.

The activating enzymes of the lectin pathway comprise a set of serine protease named the MASPs after the first discovery of their interaction with the MBL molecule. The MASP family is comprised of five different proteins arising from two different genes (Yongqing et al., 2012). MASP-1 and MASP-3 are enzymes, whereas mannose-binding lectin/ficolin/collectin associated protein 1 (MAP-1) (also termed MAP-44) is a truncated lectin pathway regulator without enzymatic activity (Matsushita and Fujita, 1992; Sato et al., 1994; Degn et al., 2009; Skjoedt et al., 2010). They are derived from the MAP1 gene by alternative splicing, whereas the MASP-2 enzyme and the truncated protein MAP-2 [also termed MAP-19 or small MAP (sMAP)] are derived from the MAP2 gene (Thiel et al., 1997; Stover et al., 1999).

All the MASPs and MAPs are found in the circulation as calcium-dependent homodimers predominantly associated with recognition molecules. MASP-1 is a multifunctional protease that may activate MASP-2 and cleave C2 (Héja et al., 2012). However, it also influences the coagulation and kinin systems from multiple angles (Ekdahl et al., 2016). MASP-3 has been shown to be crucial in the activation of the alternative pathway by cleaving the proform of factor D to the active form initiating its enzymatic activity (Dobó et al., 2016b; Hayashi et al., 2019). Apart from profactor D, the substrate specificity of MASP-3 is not well determined, but indeed insulin-like growth factor–binding protein-5 is the first and at present the only other protein that MASP-3 has been shown to cleave except for profactor D (Cortesio and Jiang, 2006). The predominant substrate of MASP-2 is C4, and this is followed by C2, which is analogous to C1s (Rossi et al., 2001). Thus, both MASP-1 and MASP-2 appear to be necessary to create a robust C3 convertase.

Recently, it has been shown that MASP-2 under certain circumstances might induce direct activation of C3 into C3a and C3b, creating a C4 and C4 bypass mechanism (Yaseen et al., 2017). This interaction might be involved in enhancing the effect of the alternative pathway amplification loop. This noncanonical activation mechanism might be of importance under several pathophysiological circumstances. The function of the noncatalytic splice variant MAP-2 also appears to have a regulatory function in the lectin pathway but to a lesser degree than MAP-1 (Rossi et al., 2001).

C1-inhibitor inhibits the enzymatic activity of MASP-1 and MASP-2 (Ambrus et al., 2003). However, a contribution of antithrombin in the presence of heparin has also been reported (Presanis et al., 2004). The molecular mechanism behind MASP-3 regulation is at present unknown. MBL, ficolin-1, and ficolin-2 interact with CRP, serum amyloid P, and PTX3, whereas this does not appear to be the case for ficolin-3 (Ma et al., 2017). C4BP is also a lectin pathway regulator serving as a cofactor for factor I in the cleavage and control of C4 and to a lesser degree C3, and it also has a decay-accelerating function on the C3 convertase (Blom et al., 2004). No exclusive lectin pathway receptors have been described, and the suggested receptors are limited to the different C1q receptors (Ghebrehiwet et al., 2019). None of these have firmly been established as real receptors but at present act more as binding partners. Recently the endocytic collagen receptor urokinase plasminogen activator receptor–associated protein has been shown being important for sequestration of some collectins in tissues (MBL and SP-D) (Jürgensen et al., 2019).

2. Deficiencies of the Lectin Pathway Components. Deficiencies of certain lectin pathway components are relatively common, whereas others are rare (Degn et al., 2011; Goicoechea de Jorge et al., 2018). The phenotypes vary from being healthy to presenting with a severe disease usually related to a tendency for increased infection, increased severity of different diseases without being causally related, or, surprisingly, embryonic development disturbances. The most common deficiency of the lectin pathway is due to low concentrations
or dysfunction of MBL, which is seen in 5%–7% of the Caucasian population (Garred et al., 2006). Although the relevance of MBL as a single disease-causing factor is still a matter of debate, decreased MBL levels are considered to be an aggravating factor in the pathophysiology of diseases, such as cystic fibrosis (Garred et al., 1999; Dorfman et al., 2008; Degn et al., 2011) and other chronic lung diseases not related to cystic fibrosis (Chalmers et al., 2013). Common variable immunodeficiency is another example of a chronic illness wherein low levels of functional MBL appear to aggravate the disease course (Andersen et al., 2005; Fenvang et al., 2005; Litzman et al., 2008).

Low levels of MBL may also have a negative impact on kidney graft survival (Bay et al., 2013; Golshayan et al., 2016; Czerewaty et al., 2019) and on long-term outcomes of cardiovascular diseases (Ohlenschlaeger et al., 2004; Vengen et al., 2012). However, this might be counterbalanced by an acute proinflammatory function of MBL since high levels of MBL have also been shown to be associated with adverse outcome in more acute situations as well (Fumagalli et al., 2017). It has been proposed that MBL association may follow a U-shaped curve since both low and high levels might be associated with disease (Troelsen et al., 2010). This has recently been elegantly demonstrated in a type 2 diabetes population cohort in that both low- and high-serum MBL (as predicted from the genetic polymorphisms) were associated with poor cardiovascular outcome and mortality (Gedebjerg et al., 2020). A similar phenomenon has been observed for MBL in patients admitted to intensive care units because of systemic inflammation and sepsis (Hellemann et al., 2007). Nevertheless, this makes interpretation of MBL disease associations difficult and requires that these studies be carefully planned and performed.

Several single-nucleotide polymorphisms (SNPs) have been identified in the promoter and exons of the MBL2 gene encoding MBL in humans that explain the high variability observed in MBL serum levels. Most of that variation can be attributed to three different SNPs (i.e., alleles B, C, and D) within the MBL2 exon 1 and SNPs in the promoter region of the MBL2 gene, which gives rise to a complex haplotype system that determines the MBL serum concentration (Garred et al., 2006). However, the serum concentration may vary considerably between individuals carrying the same haplotype. Even in homozygotes for the structural variants, MBL protein might be detected. However, this variant protein is dysfunctional with low binding avidity to ligands and cannot activate complement (Garred et al., 1999; Garred et al., 2003; Larsen et al., 2004).

CL-10 and CL-11, which are encoded by COLEC10 and COLEC11, respectively, are two highly homologous collectins that mostly circulate in the plasma as high-molecular-weight CL-10/CL-11 heterocomplexes (Henriksen et al., 2013; Bayarri-Olmos et al., 2015, 2018). They display an identical domain distribution and a 47% sequence homology at the amino acid level, but whereas CL-11 is a known activator of the lectin pathway (Bayarri-Olmos et al., 2018), the biologic role of CL-10 on its own remains obscure. However, the heterocomplexes between CL-10 and CL-11 activate complement (Henriksen et al., 2013). Recently, exome sequencing studies in patients with Malpuech, Carnevale, Michels, and Mingarelli syndrome (3MC syndrome) revealed that several of them carry COLEC10 and COLEC11 mutations as homozygotes or compound heterozygotes, which result in complete CL-10 or CL-11 deficiencies (Rooryck et al., 2011; Munye et al., 2017). Originally described as four separate disorders, the 3MC syndrome is an ultra-rare congenital disorder characterized by mental retardation, growth deficiency, and physical abnormalities, such as cleft lip, hypertelorism, eyelid drooping, and skeletal malformations (Titomanlio et al., 2005). It has been suggested that CL-10 and CL-11 are essential in the development of craniofacial structures by regulating neural crest cell migration and maintaining cell adhesion (Gajek et al., 2020).

For the three ficolins, only ficolin-3 deficiency, which is caused by a rare frameshift mutation in the FCN3 gene, has been described in humans with variable clinical manifestations mainly related to a tendency for increased infection, autoimmunity, and neurologic complications (Munthe-Fog et al., 2009; Schlabach et al., 2011; Michalski et al., 2012; Troldborg et al., 2019; Babaha et al., 2020; Dadfar et al., 2020). However, polymorphisms in the FCN1 and FCN2 genes associated with different serum levels of ficolin-1 and ficolin-2, respectively and their functions have been associated with different infectious and autoimmune conditions, but no clear consensus exists about these associations (Garred et al., 2016).

MASP deficiencies are rare (Degn et al., 2011). MASP-2 deficiency was the first MASP deficiency to be described and has been associated with recurrent infections and chronic inflammatory diseases (Stengaard-Pedersen et al., 2003; Bibert et al., 2019). However, it appears not to be indispensable since deficiencies are observed in healthy individuals and blood donors (García-Laorden et al., 2020). Mutations in the MASP1 gene have also been reported in families with 3MC syndrome (Sirmaci et al., 2010; Graul-Neumann et al., 2018; Basdemirci et al., 2019). The large majority of MASP genetic variants are located in exons coding for the specific serine protease domain of MASP-3 (Gajek et al., 2020). All described variants except for one destabilize the protein structure leading to intracellular degradation or decreased protein expression (Gajek et al., 2020). Regardless of their location in the MASP1 gene, all mutations obliterate MASP-3 activity, thereby reinforcing the notion that impaired
activity of MASP-3 activity is central to the development of the 3MC syndrome.

Both MASP-1 and MASP-3 display a wide variety of activities, which may be for the different consequences of defective MASPs. MASP-1 is known to interact with systems spanning well beyond the lectin pathway, such as coagulation and kallikrein-kinin as well as inflammatory and cellular processes, including the activation of PAR (Dobó et al., 2016a). Originally thought to be mediated by MASP-1, MASP-3 has recently been shown to be a principal profactor-D activator in blood in addition to its association with the 3MC syndrome (Iwaki et al., 2011; Dobó et al., 2016b; Oroszlán et al., 2016; Gajek et al., 2020). In the latter, the molecular substrate specificity has not been solved yet. So far substrate specificity of MASP-2 appears to be narrower than that of MASP-1 and MASP-3, but MASP-2 may also cleave prothrombin into active thrombin in addition to its role in the complement system (Dobó et al., 2016a).

3. Disturbances of the Lectin Pathway and Therapeutic Targets. MBL deficiency has been regarded as an attractive pharmaceutical treatment possibility, particularly in the situation of treatment-induced neutropenia combined with MBL deficiency (Neth et al., 2001; Peterslund et al., 2001; Eisen and Minchinton, 2003; Jensenius et al., 2003). Substitution therapy both with plasma purified and MBL produced by recombinant technique has been tried in smaller scale (Valdimarsson et al., 1998; Garred et al., 2002; Petersen et al., 2006; Bang et al., 2008; Brouwer et al., 2009), but more extensive studies have not been initiated so far. Pharmaceutical regulation of the activity in the lectin pathway has gained interest over the last few years since lectin pathway activation mediated via MASP-2 has been shown to associate with several inflammatory diseases (Dobó et al., 2016a); suitability of substitution therapy for lectin pathway deficiencies remains to be determined.

An inhibitory anti–MASP-2 antibody has been suggested to be effective in pilot trials in IgA nephropathy, SLE nephritis, PNH, aHUS, and bone marrow transplantation–associated thrombotic microangiopathy (https://www.omeros.com). This has led to the initiation of phase 3 trials, which await completion. Also, inhibition of MASP-3 appears to be an attractive target in alternative pathway-related diseases, but so far results from clinical trials have not been published. Thus, it may be anticipated that pharmacological manipulation of the lectin pathway might be a feasible target in the future.

C. Alternative Pathway Deficiencies

1. The Components of the Alternative Pathway, Their Activation, and Regulation. The alternative pathway is shown at the upper right part of Fig. 1. The alternative pathway is the amplification loop of the complement system, enhancing the activation potential of both the classical and lectin pathways by 80%–90% and leading to an efficient activation of the terminal C5–C9 pathway (Harboe et al., 2004; Harboe et al., 2006). It consists of the central component C3; factor B; factor D, which is the only protease circulating in active form in plasma; and the regulatory protein’s factor H, which is the analog to the classical pathway C4BP, which is a cofactor for factor I in cleavage and inactivation of C3b. Properdin is the only regulatory protein in the complement system that enhances activation, and its role in complement activation was recently revised (Harboe and Mollnes, 2008). It binds to the alternative pathway convertase C3bBb giving C3bBbP, with an extended half-life of the convertase enabling more C3 cleavage. The role of properdin as a recognition molecule is under debate (O’Flynn et al., 2014; Harboe et al., 2017). In addition to these fluid factors, several membrane receptors act as regulators of the alternative pathway: CR1 (CD35) binds C3b, C4b, and, to a lesser extent, C1q and is a cofactor for factor I. Membrane cofactor protein (MCP) (also known as CD46), decay-accelerating factor (DAF) (also CD55), and CR1g also regulate the extent of C3b decay acceleration and/or cofactor activity (Fig. 1).

C3 is activated by several mechanisms: First, the internal thioester spontaneously hydrolyzes resulting in the formation of C3(H2O), which binds factor B, which is subsequently cleaved and activated by factor D, which then cleaves C3 to C3b and C3a. This was called C3 “tick-over,” as described in 1980 (Tack et al., 1980), and the concept was revisited recently (Fromell et al., 2020). Second, C3 is activated by the classical/lectin C3 convertase C4bC2b (Bohlon et al., 2019). Finally, once C3b is covalently linked to foreign surfaces, it can bind to factor B, which is in turn cleaved by factor D. This surface attached the alternative pathway convertase C3bBb and can continue to cleave more C3, depositing more C3b on the foreign surface. Such continued activity is not supported on self surfaces because it is inhibited by the regulators mentioned above. For example, factor H competes with factor B and prevents further cleavage of C3 and deposition of C3b on host cells. In general, factor H is the most important regulatory protein in the alternative pathway. This is particularly due to the protein not only inhibiting the C3 convertase in the fluid phase but also because it binds to the endothelium and other cells to protect from host surface activation (Jokiranta et al., 2005; Heinon et al., 2007). This is a particularly important mechanism when it comes to the pathogenesis of complement and to the therapeutic utility of factor H, factor H fragments as a mediator of complement inhibition in a number of diseases.

2. Deficiencies of the Alternative Pathway Components. Deficiencies of the alternative pathway are the most important with respect to development of severe human diseases. In particular, mutations in
factor H and, secondly, in other regulators, including factor I and CD46, can lead to severe diseases due to uncontrolled turnover of C3 and consequent excessive activation of the whole system (Meri, 2013) (Fig. 2D). Similarly, gain of function of C3 and factor B give the same phenotype as the loss of function in the regulatory proteins since these also result in uncontrolled C3 activation (Fig. 2D). Rare genetic deficiency of C3 results in serious infections and kidney diseases (Singer et al., 1994; Reis et al., 2006). One patient who was factor B-deficient has been described (Slade et al., 2013), and a few cases of human factor-D deficiencies have been characterized wherein the patients suffer from infectious diseases, particularly Neisseria, and kidney disease (Singer et al., 1994; Reis et al., 2006). One patient who was factor B-deficient has been described (Slade et al., 2013), and a few cases of human factor-D deficiencies have been characterized wherein the patients suffer from infectious diseases, particularly Neisseria, and kidney disease (Singer et al., 1994; Reis et al., 2006). In contrast, genetic properdin deficiencies are well known with different penetrance and with increased susceptibility for infections, particularly for neisserial species since these bacteria are prone to being killed by C5b-9 (Sjöholm et al., 1988; Schejbel et al., 2009). They often require C5b-9 or a higher level of C3 opsonization to be killed.

3. Disturbances of the Alternative Pathway and Therapeutic Targets. As with the classical and lectin pathways, there are no specific substitution therapies for deficient alternative components. Plasma infusion is frequently contraindicated, as it brings “oil to the fire” in cases with genetic deficiency, inducing antibody responses to the exogenously provided component, which is seen as a foreign protein. Plasma exchange may be an option if there are nephritic factors (NeFs) present, which are autoantibodies to the convertases that stabilize them and act phenotypically like a factor-H mutation. However, this is an unsatisfactory therapy, and other treatments exist, including general immunosuppression or immunotherapy, such as rituximab to limit antibody production. However, recent years of experience have suggested complement inhibition to be the treatment of choice in most of the disturbances of the alternative pathway. It will be less demanding and give fewer adverse effects than the current therapy.

Targets to be inhibited in the alternative pathway to reduce overactivation are C3, factor D, factor B, and properdin. C3 blocks the whole cascade and not only the alternative pathway, as it is the common molecule at which the three initial pathways merge and thus would be the most potent inhibition of the system. Factor D is the rate-limiting molecule of the alternative pathway and requires small amount of drug in contrast to factor B, which is present in large amounts. Inhibition of properdin reduces the alternative activation, as it is a positive regulator.

There are several receptors that bind C3 fragments. The C3a molecule and its receptor C3aR are both therapeutic targets (Ames et al., 2001; Lohman et al., 2017; Ahmad et al., 2020). C3a is generated as a result of
activation of any of the three initiation pathways upon cleavage of C3. It binds to a G protein–coupled seven-span transmembrane receptor C3aR. The consequences of this receptor signaling can be either proinflammatory or anti-inflammatory and vary depending on cell type. For example, C3a induces the release of histamine from mast cells and enhances proinflammatory cytokine production in macrophages but suppresses neutrophil migration to inflamed tissues [as reviewed in Coulthard and Woodruff (2015)]. In addition to the C3aR cell-type specificity, some responses require costimulation with other receptors, such as Toll-like receptors or C5aR1. To further complicate the system, it has been shown that a peptide cleavage fragment of VGF, TLQP-21, suppresses microglial activation via C3aR (El Gaamouch et al., 2020) and thus may be particularly influential in the context of Alzheimer disease because VGF has been identified as an important regulatory protein in Alzheimer disease (Beckmann et al., 2020). There is also evidence for an additional (and perhaps counter-) receptor for C3a, as enhanced expression of C3a in brain of a C3aR knockout mouse provided protection in a mouse model of endotoxic (lipopolysaccharide) shock (Boos et al., 2005). These dual activities are a challenge when trying to apply therapeutic targeting strategies in both acute and chronic disorders.

The other C3 fragment receptors serve various functions. Specifically, CR1 (CD35) binding C3b and CR1g binding C3b and iC3b protect the host cells against complement attack (He et al., 2008) and can facilitate phagocytosis of opsonized pathogens. CR2 (CD21) binds C3d covalently linked to antigen, resulting in enhanced B-cell activation upon antigen recognition by the B-cell surface immunoglobulin. CR3 and CR4 (CD11b/CD18 and CD11c/CD18) are integrins binding iC3b and serve as efficient phagocytosis receptors for microbes, cells, and particles opsonized by iC3b. There are soluble forms of several of these receptors, and interestingly some of them are possible targets for inhibition of C3 cleavage, such as sCR1 (Weisman et al., 1990), CR1g (Katschke et al., 2007), CD55 (Spitzer et al., 2004), and CD46 (Christiansen et al., 1996).

D. Terminal Pathway Deficiencies

1. The Components of the Terminal Pathway, Their Activation, and Regulation. The terminal pathway is shown at middle lower part of Fig. 1. When the C5 convertases from the classical or lectin pathway (C4b2bC3b) or the alternative pathway (C3b2BbP) have been formed, the initiation of the terminal pathway starts with the last cleavage in the cascade, with the C5 molecule being cleaved to C5a and C5b. In contrast with C3 and C4, C5 has no internal thioester and therefore cannot be covalently bound to the membrane. However, a binding site for C6 is exposed on the cleaved C5b, and the bimolecular C5b6 binds C7. The hydrophilic single components change to an amphiphilic state the C5b67 complex is able to bind to a nearby membrane. This usually occurs on the surface where the activation has started, but since C5b-6 is water-soluble and not always bound immediately to the activating surface, it can move to another cell membrane and attack this cell instead after binding C7 and subsequently assemble C8 and C9. This phenomenon is called “bystander lysis” (Lachmann and Thompson, 1970). Although the cell is not necessarily lysed, it is attached and may be activated. Of particular importance in this regard is that some complement components, in particular C7, are produced locally in the tissue (Würzner et al., 1994) and can bind C5b6 and initiate the terminal assembly (Würzner, 2000).

The C5b-9 terminal complement complex (TCC) exists in two forms. If inserted completely into the cell membrane it is termed the membrane attack complex. Notably, there are no membrane receptors for this complex. It is inserted because the induced amphiphilic structure results in a lipophilic state such that the complex can penetrate and make a hole in the membrane. The main membrane regulator of C5b-9 is the CD59 molecule that is able to bind C8 and C9 blocking pore formation and lysis of self cells (Meri, 1994). If the cell, which is typically a nucleated metabolically active cell, is protected from lysis, it may be activated instead and release inflammatory mediators in a process called sublytic attack, which then contributes to tissue inflammation and, if excessive, tissue destruction (Morgan and Campbell, 1985; Zhang et al., 2014b).

If C5b-7 is not bound to a surface but remains in plasma or other fluids, the sC5b-9 complex accumulates. Soluble regulators are required to cover the lipophilic sites and keep the whole complex hydrophilic and soluble. The regulatory proteins vitronectin and clusterin are attached after C7 is bound such that the C8 and C9 cannot incorporate into a membrane. There is no receptor for sC5b-9, but it has been postulated that it can bind to the vitronectin receptor via the vitronectin molecule part of the complex (Bieseker, 1990). It has no known biologic function but is a very valuable tool to measure the degree of complement activation (Harboe et al., 2011).

C5a is the most potent inflammatory mediator of the complement system (Manthey et al., 2009; Wood et al., 2018). It is rapidly inactivated by carboxypeptidases cutting off the terminal arginine and thus termed C5adesArg. C5a has two membrane receptors, the traditional highly proinflammatory C5aR1 (CD88) and the later discovered C5aR2, which seems to have partly opposing effects of C5aR1, keeping this inflammation under some control (Laursen et al., 2012; Yan and Gao, 2012; Wood et al., 2018; Li et al., 2019).

2. Deficiencies of the Terminal Pathway Components. C5-C9 deficiencies are rare and typically associated with Neisseria infections, whereas association with other infections or autoimmune diseases is debated.
(Würzner et al., 1992; Tedesco et al., 1993). A high incidence of C9 deficiencies has been documented in Japan (Fukumori et al., 1989), which is of particular interest since the incidence of Neisseria is low in Japan. Consequences of C5 deficiency are of particular importance because a gradually larger patient population is treated with a complement C5 blocker, as will be detailed below. These patients have the same phenotype as genetic C5 deficiencies if treated over a long period. Except for Neisseria, these patients rarely suffer any other complications (Scheijbel et al., 2013; Rondeau et al., 2019). CD59 deficiency occurs in two different forms. When there is a genetic defect in the CD59 gene, a separate phenotypical syndrome with many dysfunctions can be seen (Ben-Zeev et al., 2015), such as paroxysmal nocturnal hemoglobinuria (Yamashina et al., 1990). This is also seen when both CD55 and CD59 are lacking on the red cell membrane due to a defect in the glycosyl phosphatidyl inositol (GPI) anchor (Kinoshita, 2018).

3. Disturbances of the Terminal Pathway and Therapeutic Targets. Except for the consequences of the terminal components described above, dysregulation and imbalance of the terminal pathway are normally due to an imbalance in the earlier steps of the cascade, particularly the alternative pathway. This leads to excessive cleavage of C5 with subsequent potent inflammatory reaction and organ damage. Thus, C5, C5a, and the C5a receptors are important therapeutic targets for upstream imbalance and overactivation of the system. Although in most cases it is C5a that is the most pathogenic mediator, C5b-9 may contribute in addition. In such cases the rationale is to block C5 cleavage. If C5a is the only mediator, a C5a/C5aR axis blocker would be ideal to keep C5b-9 intact to protect against Neisseria infection. However, in certain instances C5b-9 is the crucial mediator, such as in PNH, and in such a case, targeting a component downstream to C5 to protect the assembly of C5b-9 but leaving C5a free for normal defense could be an actual approach.

E. Therapeutic Approach to Treat Complement Deficiencies or to Enhance Its Efficacy

1. Substitution of Purified Components and Plasma. Complement deficiencies cannot be treated with substitution of the completely missing component because of lack of access and the caveats mentioned above. Specific complement proteins have not been purified or recombinantly produced on a large scale for clinical treatment. The two components that have been proposed for large-scale production for treatment are MBL (Laursen, 2003) and factor H (Michelfelder et al., 2017). Indications for MBL treatment are questionable because of lack of diseases with known specific need or benefit; this is in contrast to factor-H treatment, which might be highly desired. The cost of production would most likely be relatively high, as the diseases requiring factor-H substitution are relatively rare. A recombinant mini–factor H has been constructed and conjugated to CR2 (TT30), but this is not primarily to replace factor H but to inhibit activation where C3d is bound in the tissue, and the treatment will thus be target-specific as will be discussed below.

2. Prophylactic and Symptomatic Treatment. Cases of deficiencies with risk of infections should be vaccinated, particularly against pneumococci and Neisseria, and active infections should be treated with antibiotics. Patients with properdin deficiency leading to insufficient terminal pathway activity should be vaccinated with the latest generation of vaccines since they often get infected with rare species. This is also the case for patients treated with a C5 blocker. They should have antibiotics at home and should be able to call a professional healthcare provider 24/7 if they get symptoms. Interestingly, a question arises for the terminal pathway deficiencies: if the pathogenic strain is serum-sensitive, will the vaccine work if the response depends on C5b-9 formation only? The answer is most likely not, since the C5 is inactive; thus, the response will depend on the Fc-mediated phagocytosis (Jodele et al., 2016; Reher et al., 2018; Gäckler et al., 2020).

3. Liver Transplantation. Most of the complement components are produced in the liver, including the crucially important regulatory factor H. Loss of function of factor H is relatively rare, but when occurring it often leads to severe diseases due to an extreme overactivation of the alternative pathway and subsequent activation of the terminal pathway. The prototypical example is the form of thrombotic microangiopathy, aHUS. This is one of the four FDA-approved diseases for C5 inhibition with eculizumab. The effects and results are excellent (Fakhouri et al., 2016; Wong and Kavanagh, 2018; Gonzalez Suarez et al., 2019), but still the factor H is mutated in the patient. In a few cases in which these patients have had both kidney and hepatic failure, a combined transplantation was performed (Jalanko et al., 2008; Tran et al., 2014; Nayagam et al., 2020). The patients got a new liver producing normal factor H and a new kidney that was not attacked by the host’s previous overactivated complement. This will certainly be a limited offer to very few patients, but it is the only treatment to permanently replace the defective component with a normal one until genetic therapy might be a reality.

4. Triggering of Complement Activation in Individuals with a Normal Complement System: An Anticancer Approach. Often the clinical problem is that the imbalance in complement homeostasis is enhanced activation of complement, which needs to be attenuated. However, there is one particular condition we should not neglect, and that is patients treated with therapeutic antibodies. There are many of such antibodies, but the prototype is lymphoma treated with rituximab or recent similar antibodies that bind CD20+ B-cells. This
strategy has changed the treatment of lymphoma substantially in the last few decades, and important treatment modifications have been proposed (Beurskens et al., 2012). Notably, cancer cells tend to increase their number of surface complement inhibitors, including CD59, as it is a protective mechanism for the cells to survive complement attack. In recent years, a new approach has developed to block the complement inhibitors in combination with the antitumor antibodies, so-called bispecific antibodies, to increase the sensitivity of these cells to be killed by complement. This principle has not reached the clinic yet, and this review will not go further into this field of complement, but some important contributions are cited here (Maio et al., 1998; Jurianz et al., 1999; Fishelson and Kirschfink, 2019).

5. Gene Therapy. Even though it is a very attractive approach so far, gene therapy to modulate complement function has not reached the clinic. Gene silencing using RNA interference delivery has been attempted to reduce concentration of C5 using the compound ALN-CC5 for treatment of aHUS in phase 1 and phase 2 trials (https://clinicaltrials.gov/ct2/show/NCT03303313). No results have been published so far. However, preclinical studies of ALN-CC5 and efficacy of C5 silencing in rat models of myasthenia gravis support further clinical development of the RNA interference approach as a potential therapeutic complement-mediated disorder (Kusner et al., 2019). Another approach is to increase the local concentration of factor I by gene therapy, thus downregulating excessive activation of the complement system. This strategy is presently being tested in human phase 1 and phase 2 trials for local treatment of dry age-related macular degeneration (https://www.gyroscopetx.com). It must be assumed that different forms of gene therapies with different targets will emerge as attractive candidates for the treatment of complement-mediated diseases when this area is developed further.

III. Role of Complement in Disease Pathophysiology

Normally complement is a system in a fine-tuned balance between inhibition and amplification resulting in homeostasis (Fig. 2B). It has, however, a substantial potential to get out of control if there is a shift in this balance (Fig. 2A). Deficiencies or loss-of-function mutations of the ordinary complement components may lead to infections and autoimmune diseases (i.e., the balance is shifted to hypoactivation; Fig. 2C). However, the main reasons for acute complement dysfunction in the clinic, which can be life-threatening within hours or days, is an imbalance and dysfunction related to enhanced activation of the system (Fig. 2D). Thus, this is the main task for the future therapeutic approached—in other words, to dampen the activation by complement inhibitors thereby re-establishing the balance and homeostasis.

A. Consequences of a Dysfunctional Complement System

1. Decreased Regulatory Activity and Increased Activation Potential. It is crucially important to keep the complement system in balance. This is the case for any of the blood cascade systems. They are circulating in a preactivated form ready to act immediately, within seconds, as soon as a danger is sensed. Importantly, they should act locally—like coagulation making a clot to stop bleeding in a cut in a finger—and complement to stop and kill an intruder in the ear. The cascades can be compared with undetonated bombs: Normally they do no harm, but if they explode systemically, as in sepsis, they might kill the host (Fig. 2A). The cascades are built up by approximately as many ordinary activation components as regulatory inhibitors to keep the system in check.

This scenario is comparable to playing football. There are 11 players on each team, with each taking care of the other team’s players; when a red card is given and one of the players has to leave, there is an unbalanced shift to one side. Thus, if in a cascade there is one defective inhibitor or one overactive component, the imbalance will occur with the same consequence (i.e., with a shift to the activation side). For the complement system a typical example is that a loss of function in factor H in one patient will frequently give the same phenotype as a gain of function of C3, typically leading to, for example, aHUS (Fig. 2D).

The aims of complement inhibition are thus to bring balance to the system both in acute and chronic diseases. In a systemic inflammation, such as with trauma and sepsis, it is important to immediately avoid the overactivation and explosion of the bomb, whereas in chronic inflammation it is important to over time reduce the complement activation and the subsequent tissue damage. Both cases have the aim of attenuating irreversible tissue damage and organ failure.

2. Pathophysiologic Role of an Uncontrolled Overactivated System. In pathophysiologic conditions it should be emphasized that it is the innate immune system that reacts immediately after danger appears, whether through external pathogen-associated molecular patterns (PAMPs) or internal damage-associated molecular patterns (DAMPs). Thus, the complement system will react within seconds to minutes and generate C5a. As mentioned above, the anaphylactic peptide C5a in excess can be the most potent activator of an inflammatory response. C5a via C5aR1 signaling synergizes with Toll-like receptors (TLRs), TLR cofactor CD14 (Barratt-Due et al., 2017), NFκB activation, and the nucleotide-binding oligomerization domain-like receptor 3 (NLRP3) inflammasome activation leading to proinflammatory cytokine production, such as IL-1β.
and tumor necrosis factor (Zhang et al., 2014a). C5a was found to increase the release of IL-1β and IL-6 when added to Aβ amyloid-primed human monocytes in vitro (O’Barr and Cooper, 2000), and C5aR1 antagonists have been found to reduce proinflammatory cytokines like IL-1β in periodontal disease (Hajishengallis and Lambris, 2012; Recknagel et al., 2012). Importantly, C5a will start the further downstream sequelae, including reactive oxygen species generation as well as platelet activation and a number of other inflammatory mediators (Barratt-Due et al., 2017). This a major reason for blocking the upstream mediators, including C5a in particular, instead of one of all the hundreds of downstream inflammatory mediators generated as a result of C5a responses (Ward, 2004; Panayiotou et al., 2019; Li et al., 2020). In addition, it is possible that in the CNS, as in the periphery, C5a via C5aR1 signaling synergizes with TLR/NFκB and the NLRP3 inflammasome activity to trigger potent detrimental inflammatory responses (Yang et al., 2020).

If unopposed or overwhelming, this can lead to the cytokine storm that systematically, as in sepsis and severe trauma, leads to shock and lethality or, if local, will generate tissue damage proportional to the imbalance of reactions. Suppressing the NLRP3 signaling cascade or the downstream cytokine response with the important IL-1β has long been suggested as a potential therapeutic strategy for diseases with a detrimental inflammatory component. However, use of a C5aR1 antagonist would eliminate the synergistic response while avoiding systemic direct suppression of critical NLRP3, thereby limiting or eliminating susceptibility to inflection.

In pathophysiologic conditions it should be emphasized that it is the innate immune system that reacts immediately after danger appears, whether through external PAMPs or internal DAMPs. Thus, the complement system will react within seconds to minutes and generate C5a. Thus, C5a is one of the most upstream mediators released and signals together with other upstream recognition receptors (e.g., TLR cofactor CD14) (Barratt-Due et al., 2017). Importantly, C5a will induce further downstream inflammation, including reactive oxygen species production, and release of the whole network of cytokines from monocytes and macrophages, platelet activation, and a number of other inflammatory mediators (Barratt-Due et al., 2012). This a major reason for blocking mediators like C5a in particular instead of one of all the hundreds of downstream inflammatory mediators generated as a result of C5a responses (Ward, 2004; Panayiotou et al., 2019; Li et al., 2020).

3. Complement Attack on Host Cells when They Lack Regulators. A unique property of cascade systems is their continuous low-grade activation under physiologic conditions. This has the advantage that they are ready to react immediately; as compared with driving a car, they do not need to start the engine first (Barratt-Due et al., 2012). This also implies that they will attack the host’s own cells unless the cells are well protected by

**Fig. 3.** Loss of host cell regulators leads to attack by the host’s own complement. (A) A normal erythrocyte protected from complement by DAF (CD55) and CD59. (B) An erythrocyte from a patient with PNH, which is caused by a clonal deletion in enzyme enabling GPI anchors, which is responsible for binding both DAF (CD55) and CD15 to the surface, resulting in cell lysis. (C) PNH cell from a patient treated with eculizumab, a monoclonal antibody blocking cleavage of C5. The cell is protected from C5b-9 attack. FB, factor B.
inhibitory regulators. CR1 binds C3b as ligand and is a cofactor for factor I that cleaves C3b to iC3b, which can no longer form the C3 convertase C3bBb. On red cells the large amount of CR1 binds C3b opsonized particles, such as immune complexes, and traffics them to the liver for clearance (Cornacoff et al., 1983; Brekke et al., 2019).

Genetic deficiency of CD59 results in insertion of the C5b-9 complex into the membrane and is associated with a complex syndrome, including hemolysis seen in PNH (Tabib et al., 2017). CD55 is also a protective regulator of C3b activity on the cell surface, and a rare deficiency of this protein has shown pathophysiology mainly in the gastrointestinal tract (Ozen et al., 2017).

The most prominent disease with lack of complement inhibitors on its blood cells is PNH (Fig. 3). In contrast to those mention above, this is not a genetically inherited defect but a somatic mutation in the bone marrow of an enzyme required for membrane association of complement regulators. The Phosphatidylinositol N-acetylglucosaminyltransferase subunit A (PIG-A) or subunit T (PIG-T) genes are mutated for the GPI anchor (Takeda et al., 1993; Krawitz et al., 2013). More than 20 proteins that do not have a transmembrane signaling part are bound to the membrane via this GPI anchor, including the two important complement proteins CD55 (inhibiting at the C3 level) and CD59 (inhibiting the C8/C9 insertion into the membrane) (Fig. 3). The phenotype of the disease is red blood cell hemolysis to a severe degree, which often requires blood transfusions, and frequently thrombosis (Hill et al., 2013). Particularly serious are those developing in the gastrointestinal area, such as the Budd-Chiari (De-la-Iglesia et al., 2016). The details of treatment of PNH are given below.

4. Complement Attack in Tissue Damage and Common Pathophysiological Mechanisms to Treat a Broad Panel of Diseases. The main role of innate immunity, complement included, is to induce inflammation, both to protect from pathogenic intruders and repair damaged self. The mechanisms by which the system does this are principally the same (Chen and Nunez, 2010; Denk et al., 2012; Schaefer, 2014). With pathogenic invasion, the role for complement is to sense the conserved molecular patterns on the microbes (PAMPs) through the pattern-recognition molecules. During internal damage, typically when ischemia reperfusion occurs, a number of DAMPs are exposed, reacting with naturally IgM molecules, which can activate both the classical and the lectin pathway (Zhang et al., 2006b) (Fig. 1). The physiologic consequences of inflammation are virtually identical whether PAMPs or DAMPs have initiated them (Fig. 4). Increases in blood flow due to the damage gives the classical heat, redness, swelling, and pain. Later, reduced function is added to this list (Fig. 4, last item).

An example is a patient presenting with these signs and symptoms. They might look quite similar, and this could be due to sterile inflammation, such as tennis elbow; rheumatoid arthritis; or fracture without

Fig. 4. Signs and symptoms of inflammation. The classic four characteristics seen in any inflammation: calor, rubor, tumor, and dolor. Later, function loss was also added to these. Reprinted with permission from Lawrence T, Willoughby DA, and Gilroy DW (2002) Anti-inflammatory lipid mediators and insights into the resolution of inflammation. Nature Reviews Immunology 2:787–795. Printed with permission from Nat Rev Immunol.
complications—or it could be due to nonsterile (infectious) inflammation like septic arthritis or complicated fracture with infection. The response is principally the same. The mechanisms behind the inflammation might be similar, and the treatment will certainly be different dependent on the diagnosis. It should be emphasized, however, that all these conditions might benefit from inhibition of the innate immune system, complement inhibition included. It remains to be shown how many diseases will benefit from complement inhibition alone or in combination with regulation of other innate immune receptors as detailed in other parts of this review.

A notable idea emerging from these data with complement inhibition alone or combined with other innate immune system molecules is how far this strategy can be used for diseases with different etiology but with a common complement-induced pathogenesis. Surveying the literature, it seems like the list of possible candidates for complement inhibition might be very long, whereas very few diseases in which complement plays no role have been documented and confirmed.

IV. Therapeutic Complement Inhibition

A. Diseases with a Potential for Complement Therapeutic Modulation

1. Introduction: The Search for Complement as Pathophysiologic Mediator. To select a certain disease as a possible candidate for complement inhibition, a number of criteria should be fulfilled. From clinical evaluation of the disease, plasma should be measured for complement activation products. Not all diseases with increased local activation release activation products to plasma, so biopsies of the suspected organ(s) are needed to see whether there is activated complement deposited in the tissue. If this is the case, in vitro experiments should be done with human material, and if confirmed, this should be followed up with animal experiments in a reliable animal model. Then the possible mechanism of activation should be studied, but that is a difficult task. If not possible, inhibition of possible pathogenic targets should be investigated (Fig. 1). Then the relevant inhibitors should be selected and investigated both in ex vivo human models and in animal models. Finally, different modes of the preparation and the actual routes of administration should be decided (see below). The pharmacological kinetics, half-life, and, not least, the cost must also be considered. It is crucially important to dissect the pathophysiology and the pathogenic mechanism before deciding the therapeutic strategy. Evaluation of efficacy versus adverse effects is obviously the most important one and will be revealed during clinical trials. It is equally important to evaluate costs versus the benefits and possibly adverse effects of a certain treatment.

2. Neurologic Diseases. The prevalence of neurodegenerative diseases worldwide is accelerating. The number of individuals with dementia alone is estimated at 50 million worldwide, creating a great financial as well as personal burden. There is increasing evidence for a role for complement in Alzheimer disease (Hong et al., 2016; Hernandez et al., 2017; Shi et al., 2017) and other neurodegenerative and neurologic disorders, such as amyotrophic lateral sclerosis (Lee et al., 2018; Parker et al., 2019), epilepsy (Chu et al., 2010; Schartz et al., 2018), stroke (Alawieh et al., 2015), traumatic brain injury (Sewell et al., 2004), multiple sclerosis, and schizophrenia (Sekar et al., 2016). Some of these indicators are genetic, such as schizophrenia, and may be related to aberrant deficient or excessive synaptic pruning, but many others are based on the presence of complement activation products in the affected region of the brain or spinal cords. An influx of complement components as a result of the disruption of the blood brain barrier, such as in stroke or traumatic brain injury, results in acute complement activation in the presence of tissue debris and other DAMPs. The resulting activation fragments or inappropriate opsonization of cells will then result in cascading increases in inflammation and further cell damage/death and loss of neuronal function. However, as discussed above, it is now well established that all components of the complement system can be synthesized in the brain (Reichwald et al., 2009) and thus are present in the brain in the absence of blood brain barrier breakdown. Although often undetected in the young uninjured brain, components are induced in what appears to be a measured response to levels of injury (Fonseca et al., 2011).

In animal models, the effect of genetic ablation on neurodegeneration has contributed to the understanding of the role of complement in these diseases. C1q, C3, CR3, C3aR, and C5aR1 knockouts have provided clear evidence of the roles of the early classical cascade on synaptic pruning. However, since elimination of C1q and C3 would also prevent C3a and C5a generation, it is likely that the neuroinflammation induced by an ever-increasing cycle of glial activation, particularly via C5aR1, neurotoxicity, and damage with further complement activation overwhelms the neuronal resilience to perturbation and thereby leads to loss of cognitive and other neurologic functions. For example, in the Arctic model of Alzheimer disease (Cheng et al., 2007; Fonseca et al., 2011) that was crossed to C5aR1 knockout mice, behavior deficits and loss of neuronal integrity were prevented in the absence of C5aR1 even in the continued presence of amyloid plaques. When microglia were isolated and RNA was sequenced at increasing ages, proinflammatory gene expression was lower in the C5aR1-deficient Arctic mice, whereas genes involved in phagocytosis and lysosomal function (for clearance and homeostatic functions) were increased in
the Arctic mouse but even more so in the Arctic C5aR1 knockout mouse (Hernandez et al., 2017). These findings suggest preventing a C5aR1-induced program of gene expression lower in inflammation while allowing phagocytosis and clearance of neuronal blebs, apoptotic cells, and other debris to occur and be enhanced by the opsonic functions resulting from the unobstructed early components of complement (C1q/C1 and C3).

In amyotrophic lateral sclerosis, Parkinson disease, and Huntington disease, current investigations also focus on determining the contribution of C5a signaling through C5aR1 (Tenner et al., 2018). Amyotrophic lateral sclerosis is characterized by the progressive death of motor neurons in cortex, brain stem, and spinal cord, resulting in muscle weakness, paralysis, and death. Complement activation fragments are elevated in blood and in post-mortem spinal cord, suggesting a contribution to the pathogenesis and/or progression of this disease (Sta et al., 2011; Mantovani et al., 2014). In mouse models, although deletion of C1q, C3, or C4 did not alter the course of the disease, genetic deletion or pharmacologic inhibition of C5aR1 did reduce infiltration of inflammatory cells and nerve loss while improving muscle function and survival (Woodruff et al., 2014; Lee et al., 2017).

CNS synthesis of complement components is induced and maintained for extended periods of time in animal models of stroke (Huang et al., 1999; Pavlovski et al., 2012) and traumatic brain injury (Bellander et al., 2001). In extensive analyses of the different sequelae in an occlusion model of stroke—acute, subacute, and long-term—Tomlinson and colleagues have demonstrated that a fusion molecule consisting of a single-chain monoclonal antibody directed toward a stroke-induced modified annexin IV linked to the murine CR1 ortholog Cr1-related protein Y prevented opsonic C3b/iC3b deposition on neurons and prevented synaptic uptake (Alawieh et al., 2020) and neuronal phagocytosis by microglia (Alawieh et al., 2018a). In addition, the inhibition of complement prevented microglial activation, perhaps because of reduced generation of C5a, which otherwise persisted chronically in this model.

The result was improved long-term motor function as well as cognitive ability (Alawieh et al., 2018a). As with the sequelae of events in stroke or reperfusion injury, traumatic brain injury presents with both acute and chronic neuropathology. Using a fusion construct that targets complement activation sites with CR2 (binds the covalently bound C3d activation fragment) and the regulatory protein factor H as a treatment as much as 12 hours after contusion, lesion and scarring volume were reduced, and motor and cognitive skills improved (Leinhase et al., 2006; Alawieh et al., 2018b).

In multiple sclerosis (MS) and other disorders, the evidence for detrimental aberrant synaptic pruning and activation-induced neuroinflammation has been increasing as animal models have been developed, and technological advances allow precision characterization of disease progression. In the case of two murine models of MS, delivery of adeno-associated virus encoding Crry resulted in decreased synaptic loss and visual impairment (Werneburg et al., 2020). In addition, tissue and biomarkers have become available to demonstrate complement deposition correlates with cognitive decline in humans (Vasek et al., 2016; Wu et al., 2019).

Thus, the search for dysfunction-modifying treatments for both neurodegenerative disease and neurologic disorders in which complement imbalance or even complement mediated slowly accumulating hits to normal processes is accelerating. Every branch point in the complement cascades, every activating protein, and every regulator protein can be considered an opportunity for precision pharmacologic inhibition. Of course, different points of inhibition may be optimal for different disorders, particularly those involving a compromised blood brain barrier caused by stroke, traumatic brain injury, and multiple sclerosis. In addition, it is clear that therapies for acute neurologic injury will have different considerations than those of chronic degenerative disorders. Fortunately, there is a growing number of therapeutics specific for different complement components in various stages of drug development and clinical trials and additionally at the preclinical stage as discussed below.

3. Eye Diseases. A couple of serious diseases of the eye seem to be closely related to a disturbed complement system regulation. The most frequent is AMD, which is the most common cause of blindness in the Western world (Troutbeck et al., 2012; Park et al., 2019). Several genetic mutations of complement genes have been associated with AMD (van Lookeren Campagne et al., 2016), and many of these are associated with other complement dysfunction diseases like aHUS. It is therefore a challenge to further search for the reason why some of the genetic defects in the same proteins can lead to such different phenotypes. The current treatment of selected patients with the wet form of AMD is injection of antibodies that block VEGF. Although the mice do not have traditional AMD, it is possible to induce a choroidal neovascularization process that precedes development of retinitis conditions, including AMD (Liu et al., 2011). In this model they showed that complement activation with generation of C5b-9 came much earlier than VEGF generation and that complement blockade prevented VEGF production. Thus, complement inhibitors should continue to be investigated to stop the initial pathogenic factors leading to this debilitating disease.

In addition to AMD, a couple of other eye diseases have been related to complement activation. If neuro-myelitis optica spectrum disorder, which is close to a neurologic disease, can be considered an eye disease, its serious consequence is blindness. Previously it was regarded a part of MS, but now several entities have
been described, including the one caused by autoantibodies to aquaporin 4. This particular form has been shown to be highly complement-dependent and is now the latest of the four diseases approved by the FDA for routine treatment by eculizumab with so far good results (Frampton, 2020). Autoimmune uveoretinitis has also been efficiently treated experimentally with complement inhibitors (Chen et al., 2010; Copland et al., 2010), and, probably most surprisingly, glaucoma has come on the list of candidates for treatment because of promising experimental results (Howell et al., 2013; Reinehr et al., 2019).

4. Kidney Diseases. For a long time, it has been recognized that the pathogenesis of many kidney diseases is closely associated with complement activation (Thurman, 2020). More than 50 years ago, it was shown that the complement system is activated in the glomeruli of patients with immune-complex glomerulonephritis (Lachmann et al., 1962; Verroust et al., 1974). Over the last 15–20 years it has also become clear that the complement system is not only important to the pathogenesis of many nonimmune complex–mediated kidney diseases but also the primary driver in diseases, such as atypical hemolytic-uremic syndrome and C3 glomerulopathy (C3G) (Bu et al., 2014; Xiao et al., 2014). Certain kidney diseases are, therefore, a prime objective for complement-targeted therapy. Some examples of those that are expected to benefit from complement therapy are discussed below.

Hemolytic-uremic syndrome (HUS) is an ultra-rare disease defined by the triad of mechanical hemolytic anemia, thrombocytopenia, and acute kidney injury (Avila Bernabeu et al., 2020). Typical forms of HUS are related to infection by Shiga toxin–producing E. coli, whereas the atypical form of HUS termed aHUS is due to defects in alternative complement activation pathway (Avila Bernabeu et al., 2020). In 50%–60% of aHUS cases, genetic variants in proteins regulating the alternative pathway of complement are found, but it is a clear consensus that absence of molecular finding does not exclude an alternative pathway deregulation. The most commonly affected genes are factor H, factor I MCP (CD46), and thrombomodulin, but acquired autoantibodies like antifactor H in association with inherited deletion of particularly the factor H–related one to three genes might also be a cause of aHUS. Conversely, gain-of-function mutations in factor B or C3 are also a cause of aHUS because of increased alternative pathway.

![Figure 5](image-url)
activity surpassing the capacity of regulators of the complement activation. There are pathogenic genetic variants not related to the complement system, such as diacylglycerol kinase ε, which is an endothelial cell and podocyte protein causing aHUS in small children. So far aHUS is the only approved complement-targeted indication for kidney diseases.

C3G is a rare group of kidney diseases with a poor long-term outcome (Ahmad and Bomback, 2020). It can be further divided into two forms: C3 glomerulonephritis (C3GN) and dense deposit disease (DDD). Both diseases are characterized by C3 glomerular deposits detected by immunofluorescence. Fig. 5B shows deposition of C5b-9 in a glomerulus from a DDD case. C3GN and DDD are difficult to distinguish from each other based on light microscope and immunofluorescence investigation. However, in electron microscopy that is mesangial and/or subendothelial, intramembranous and subepithelial deposits are seen in C3GN, whereas dense osmiophilic deposits are present along the glomerular basement membranes and in the mesangium in DDD. Both C3GN and DDD are distinguished from immune complex–mediated glomerulonephritis by the lack of immunoglobulin staining in immune fluorescence, and both are characterized by dysregulation of the alternative pathway, thus being mainly complement-driven. A major subset of C3G arises from so-called NeFs, which as mentioned above are autoantibodies stabilizing the convertases and have been characterized recently as C4, C3, and C5 NeFs, which react with the different convertases surpassing the effect of complement regulatory proteins. Finally, C3 glomerulopathy can be due to genetic mutations, affecting genes that encode for complement components or regulators. The most important seem to involve factor H–, factor I– and factor H–related proteins. The C3Gs are candidates for complement therapy, but the heterogeneity of the etiology behind the diseases implies that the therapeutic effect may vary (Nester and Smith, 2016).

IgA nephropathy is the most common form of glomerulopathies worldwide (Hassler, 2020). Its clinical presentation varies, but it often includes proteinuria and hematuria. The disease is associated with aberrant O-glycosylation of mucosal IgA1 with galactose-deficient IgA1 (Gd-IgA1), which plays a pivotal role in the progression of this nephropathy. Although there are genetic, ethnic, and environmental factors involved in the pathogenesis of IgA nephropathy, the exact etiology remains unknown. A multihit model has been suggested to explain the cause of the IgA nephropathy: An increased serum concentration of Gd-IgA1 (hit 1) can be recognized by overproduced variable heavy-chain autoantibodies, such as IgG or IgA (hit 2), inducing the formation of circulating pathogenic immune complexes (hit 3). These immune complexes will finally deposit on the renal glomeruli (hit 4). This deposition triggers complement and cell activation, cytokine release, and extracellular matrix expansion (Tortajada et al., 2019). Hence, IgA nephropathy is considered as a unique autoimmune disease whereby the pathogenesis may be driven by an autoantigen (Gd-IgA1) eliciting an autoantibody (IgG antiglycan) response.

Over the past years, increasing information from findings in kidney biopsies has emerged on the involvement of complement activation in both progression of native IgA nephropathy and after a renal allograft. Several lectin and alternative pathway components have been shown to codeposit in the glomerular deposits in these diseases (Floege and Daha, 2018). Components from the alternative pathway, such as C3, properdin, or factor H, or from the lectin pathway, such as MBL alone or in combination with MASPs, ficolin-2, C4d or C4BP, are a consistent finding in kidney biopsies (Roos et al., 2006). Nonetheless, no strong involvement of the classical pathway has been associated yet (Suzuki et al., 2011). Besides the glomerular deposition of complement components, abnormal circulating levels, or genetic variations in C3, MBL, factor H, MASP-3, and factor H–related protein-5 have been associated with the severity of the prognosis. Based on this, there is a strong momentum to focus on intervention of complement activation in IgA nephropathy, as several studies have been initiated or are in the planning phase currently. For instance, an oral C5aR inhibitor, an oral factor B inhibitor, or an MASP-2 inhibitor are a few examples of phase-II/III ongoing studies on IgA nephropathy. Thus, IgA nephropathy may be one of the kidney diseases that is a candidate for complement inhibition treatment.

It is pertinent to mention that almost all kidney diseases appear to have strong complement component related to their pathophysiology, which is outside the scope of this review but might be a target for complement therapy. A kidney disease in which complement target therapy has advanced to a stage wherein we may anticipate that it will be incorporated into the treatment arsenal is accompanied by necrotizing crescentic glomerulonephritis (Trivioli and Vaglio, 2020). This disease is discussed further under the systemic autoimmune chapter item 6 in the section below.

5. Hematologic Diseases. As presented already, PNH is the prototypical example of a disease that is one of the very few and “pure” complement-mediated diseases. It was the first complement disease to be approved for routine clinical therapy, and during the last 15 years, innumerable patients have been efficiently treated with very good effects and with minor adverse effects (Kelly et al., 2011; Hillmen et al., 2013; Loschi et al., 2016; Ninomiya et al., 2016). Importantly, patients with PNH suffer thrombosis as one of the main complications and reasons for morbidity and mortality and when treated with the complement inhibitor eculizumab do not suffer from these serious thromboses (Macrae et al., 2020). In some cases, there is residual lysis despite C5 inhibition, and it has been proposed
that the C3 opsonization might be the reason for extravascular lysis, and that inhibition of C3 is an alternative to prevent this lysis (Risitano et al., 2019).

Hemolytic anemias vary with respect to the efficacy of complement inhibition. The “warm type” has a considerable IgM-mediated extravascular hemolysis at 37°C, which is less affected by complement and more Fe-dependent, whereas the cold agglutinin disease caused by IgM autoantibodies is a pure complement disease with intravascular complement-mediated lysis at low temperature (Berentsen, 2018). Since the lysis is selectively classical pathway-initiated, blocking of C1s has been shown to efficiently reduce the lysis (Shi et al., 2014). An alternative is to block C5 cleavage (Röth et al., 2009), as is used for treating PNH, since C5b-9 is the effector molecule. A future possibility would be to block a component downstream of C5 to leave C5a active while blocking lysis.

Thrombotic microangiopathy is a serious complication in patients undergoing stem cell transplantation. There are data indicating that complement is important in the pathogenesis of this condition and that complement inhibition might be a promising treatment, as seen in a study of a large pediatric cohort (Jodele et al., 2020).

Finally, it should be mentioned that accidental ABO mismatch blood transfusions occur still, although fortunately this is very rare. With this, however, the situation might be life-threatening. Most likely the complement system is the driver by intravascular lysis by the IgM anti-A and anti-B antibodies. If the reaction is severe, the patient is brought immediately to the intensive care unit for treatment. It is possible that an ampulla of a complement inhibitor should be included in the acute kit. It works immediately with virtually no risk.

6. Autoimmune Diseases. The typical autoimmune diseases like rheumatoid arthritis and SLE have a complex etiology and pathophysiology with many branches of both the innate and adaptive immunity involved. Although experimental studies have documented a clear role for complement in rheumatoid arthritis (Kahn et al., 2003; Mehta et al., 2015), the clinical application of complement therapeutics is still in the future (Holors and Banda, 2018). The same is the case for SLE (Barilla-Labarca et al., 2013). This is not surprising and is consistent with the multiple factors playing a role for disease progression. Probably there might be a place for certain subgroups, particularly those with kidney affliction as a supplement to the already established therapy.

There are, however, a group of autoimmune diseases with systemic manifestations that have a more simple and narrow pathophysiology—for example, the anti-neutrophil cytoplasmic antibodies—associated vasculitis (AAV). These are autoantibodies against proteins in the neutrophil granulocytes (e.g., proteinase 3 and myeloperoxidase). Experimental studies in mice have shown that AAV is largely driven by the C5a-C5aR1 complement axis (Dick et al., 2018), and clinical studies using a C5aR1 inhibitor are promising (Jayne et al., 2017). Another systemic disease with a single autoantibody target is the antiphospholipid syndrome (APS) frequently seen in women with both venous and arterial thromboses as well as spontaneous abortion as main complications. Traditional treatment has been heparin to prevent thromboses. However, like with AAV, there is increasing evidence that complement is of importance (Salmon and Girardi, 2004; Meroni et al., 2016). Particularly when this condition exacerbates to a severe catastrophic APS (CAPS), a systemic life-threatening condition, complement inhibitor therapy has been very efficient (Tinti et al., 2019) and was most likely lifesaving in a patient with recurrent CAPS who repeatedly responded on complement inhibition (Barratt-Due et al., 2016). Based on what we know about the pathophysiology and what is published, we suggest that AAV and CAPS are reasonably good candidates on the list of diseases to be treated with complement inhibitors in the future.

However, the great limitation with most of the studies presenting results on complement inhibition is that the number of cases in most of these are too small to perform reliable, randomized controlled trials. Thus, the use of the inhibitors is to a great extent off-label and based on case reports or small uncontrolled cohorts. Furthermore, there is a high risk of bias in the literature since a number of negative results are not published both because the authors find them less important and because editors tend to prefer positive results when it comes to a decision when pressure is on journal space.

7. Transplantation. From a complement point of view, there are three important conditions related to transplantation that should be taken into account. The first and most basic condition is ischemia-reperfusion injury. An important experimental observation was that mice within the same genetic strain were given heart transplants and a slow, long-lasting rejection was observed without any adaptive immune responses, as reviewed in Parolari et al. (2002). This could be due to an extreme overactivation of complement by exposure to DAMPs that could be recognized by the innate immune system over time. This has important consequences for preserving the organ under conditions that inhibit ischemia-reperfusion injury. A complement inhibitor would be one such candidate.

The second and very serious complication for kidney transplantation is the acute antibody-mediated rejection, which frequently leads to organ failure (Montgomery et al., 2018). This effector mechanism is mainly from the classical pathway activation of complement, and the deposition of C4d in the peritubular capillaries (Fig. 5C) is one of the Banff criteria for the diagnosis. Complement inhibition has been used off-
label with many successful outcomes (O’Neill and Pierson, 2017; Haas, 2019). Thus, acute antibody-mediated rejection might be a future candidate for complement inhibition because the kidney may be saved. However, a question to be resolved is whether the therapy can be gradually tapered or has to be continued.

The third issue is the chronic rejection, which is more complex. Antibodies are involved, but T-cells are as well, and the relative roles of these are uncertain. In this case complement inhibition should be an option, wherein it would more likely be a supplement to traditional and new therapy.

8. Systemic Inflammation: Trauma Sepsis and the Pandemic Coronavirus Disease 2019. So far, complement inhibition with success has been limited to the rare diseases. One of the reasons for this is that these rare diseases often have a pathophysiology that has been shown to be mainly complement-driven, and a single treatment with a complement inhibitor can be sufficient to keep the disease under full control. When moving to trauma and sepsis, the pathophysiology is much more complex, although complement definitely plays an essential role. Complement is an upstream actor as first-line sensor of danger and thus may accentuate the inflammatory explosion. However, there are other first-line sensors as well (e.g., the Toll-like receptors), so combined inhibition of several of these sensors might be necessary, as we have shown for combining complement and CD14 (Huber-Lang et al., 2014). It has been speculated that inhibition of CD14 could be a possible treatment of coronavirus disease 2019 (COVID-19) (Martin et al., 2020). The challenge for future strategy of complement inhibition in sepsis will be to define subgroups of patients in which complement activation is dominating. This can be combined with other upstream bottleneck sensor molecules, whereas inhibition of single downstream mediators like cytokines is less likely to succeed because of the vast amounts of them, as in accordance with the many studies that have failed by blocking single downstream mediators.

The role of innate immunity with focus on complement in the pathophysiology in trauma and its potential for therapy is somewhat comparable with sepsis, although initially the DAMPs are most important. Secondary infection is, however, frequently seen with development of a full-blown sepsis. A comprehensive review on complement in trauma was recently published (Huber-Lang et al., 2018). We therefore do not cover trauma in this review. Similarly, a recent review on sepsis and complement was published (Mollnes and Huber-Lang, 2020). Both of these reviews concluded that trauma and sepsis are both complex syndromes that need more research to dissect the role of complement and to enable correct stratification and inclusion of patients that are of high risk of getting excessive activation of complement. Once complement has exploded, it may be too late to start treatment (Fig. 2A).

COVID-19 disease is a pandemic dramatic condition that this year has changed the world in many ways. Since complement is a main sensor of danger, a number of groups have started to work on complement and COVID-19 disease. Coronavirus 2 (severe acute respiratory syndrome–coronavirus 2) can lead to life-threatening pneumonia and multiple organ failure, which is termed coronavirus disease 2019. Both bacterial and viral pneumonia have been associated with complement activation and respiratory failure (Langlois and Gawryl, 1988; Wang et al., 2015). Importantly, the coronaviruses severe acute respiratory syndrome and Middle East respiratory syndrome both were shown to potently activate complement associated with respiratory failure (Gralinski et al., 2018; Jiang et al., 2018). One of the first observations of increased complement activation in patients with COVID-19 was an increase in the activation products C5a and sC5b-9, with a prolonged activation in the latter (Cugno et al., 2020). A role for the C5a-C5aR1 axis was further supported by detection of C5a in bronchoalveolar fluid as well as inhibition of lung injury in a human C5aR1 knock mouse model (Carvelli et al., 2020). In a clinical study of 39 patients, five different complement activation products from all pathways were measured in patients who were hospitalized. All activation products were consistently elevated in all patients (Holter et al., 2020), and second, C5b-9 was correlated with respiratory function. Interestingly, antibody titers were significantly correlated with respiratory function as well, although to a less extent. It is uncertain to what extent the classical pathway versus the lectin pathway contributed to C4 activation, but most likely they were both involved. Additionally, the C3 convertase C3bBbP was significantly elevated, implying that a therapeutic approach for COVID-19 should be broad and cover all pathways (i.e., C3 or C5). The interaction between the complement system and the coronaviruses has been reviewed in Java et al. (2020) and Noris et al. (2020).

It has been speculated that pathophysiology of COVID-19 is similar to thrombotic microangiopathy, involving both complement and hemostatic dysfunctions (Chauhan et al., 2020; Conway and Pryzdial, 2020; Fletcher-Sanderson and Bellander, 2020; Gavriilaki and Brodsky, 2020; Java et al., 2020; Jodele and Kohl, 2020; Mangalmurti and Hunter, 2020; Merrill et al., 2020; Ramlall et al., 2020; Song and FitzGerald, 2020). Thus, it is reasonable to suggest that complement might be a target for treatment in COVID-19, but so far treatment has been limited to case reports, in which most show inhibition at the level of C5 (Potlukova and Kralikova, 2008; Kulasekararaj et al., 2020; Laurence et al., 2020) and a single report uses a C3 inhibitor (Mastaglio et al., 2020). Although the data are promising, from an open-labeled randomized trial with 15
patients treated with an inhibitory anti-C5a antibody and 15 controls (Vlaar et al., 2020), it remains to be shown in larger randomized control trials whether complement inhibition is an option for treatment in COVID-19 (Lo et al., 2020).

V. Mode of Complement Inhibition

A. Targets to Be Inhibited

1. Introduction. There are numerous components, fragments, and receptors that are candidates for inhibition in the complement system. The question of which targets are the best to inhibit is a nonsensical question. It will depend on a number of factors that we have described in the sections above. First, what is the pathophysiological mechanism of the disease? Some diseases are virtually completely complement-dependent, such as PNH and the cold agglutinin syndrome, whereas others are partly dependent on complement from strongly to slightly, and there probably are no conditions wherein immune inflammation is involved when complement is not involved at all. The indications for treating a condition will therefore mainly depend on the extent on the scale to which the complement is involved. Second, there is a crucial difference between chronic life-long diseases wherein the patient normally is at home and acute life-threatening conditions wherein the patients are in hospital or even at the intensive care unit under continuous monitoring and covered with antibiotics, etc., and needs inhibition just for a few days or weeks. Third, the cost of complement inhibition until now has been extremely high and limited to rare diseases. New and less expensive drugs are approaching the market, and for healthcare cost implications, this also has to be taken into consideration. A number of other factors will be of importance when discussing this essential item in the future.

Below we give some selected target molecules from the cascade that we suggest will be the most relevant in the future. This might change, as the research field is rapidly progressing. We need to emphasize that the only drug for complement inhibition that has been available for routine use is an engineered humanized monoclonal antibody targeting the cleavage of C5. Therefore, the experience of blocking other targets was until recently missing. The targets we have selected as most relevant are illustrated in Fig. 1 with an asterisk.

2. Classical and Lectin Pathway. Immediately one would think that C1q could be a good candidate. However, we argue that C1q is a molecule involved in so many other processes than complement activation that it would not be an obvious good candidate. C1r and C1s are the serine proteases that are activated when the classical pathway is triggered, and, so far, there is some evidence that C1s might be a good candidate. However, this is a good rationale only if the classical pathway is triggered. The main concern of blocking the classical pathway activation is that the antibody-mediated effects of complement are lost.

In the lectin pathway MBL has been the most studied protein and might be a candidate in certain cases, but recently MASP-2 has also been proposed as an exciting candidate to block the lectin pathway. As in accordance with the C1q mentioned above, MBL is a recognition molecule with many functions, and MASP-2 might be a good candidate also because the lectin pathway is recognized by a number of other molecules, and the activation route goes mainly through MASP-2. The main concern of blocking the lectin pathway is that a substantial part of the recognition of danger function of complement is lost.

3. C3 and the Alternative Pathway. The ultimate and most robust molecule to inhibit is C3 since it blocks the whole system from C3 and downstream. This is because all three initial pathways merge at C3, and C3 is the main molecule of the alternative pathway amplification loop. Thus, if the intention is to target one molecule to block most of the system, C3 is the molecule of choice. Thus, there is a marked potential to slow down all complement-mediated functions by blocking C3.

For specific blocking of the alternative pathway there are three candidate molecules. Factor B is the main component but requires relatively large doses because of its high serum concentration. Although factor D is the only serine protease that is circulating in active form in plasma, it is also the rate-limiting step in the alternative pathway and requires relatively small doses of a drug. Blocking both factors B and D will inhibit the alternative completely, as will virtually be the case also for inhibiting properdin. Notably, in contrast with blocking C3, the cascade will not be fully blocked by the other components, since classical and lectin pathway activation will generate some C5 convertases, and a certain terminal pathway activation will occur. Thus, the most effective blocking of the system using one target would be to block C3. Nevertheless, blocking of C3a or the C3aR might also be an alternative option to block the anaphylatoxin activity of C3 cleavage, but this has not been tried in clinical trials.

The main concern of blocking C3 and the alternative components is the reduced opsonization and probably risk of increased infection. This should, however, be no concern during an acute condition, in which the patient is monitored and covered with antibiotics. This short-time treatment will also not be a concern for a child. C3 is important for speeding up the B-cell antibody production, but short-time blocking of C3 would hardly influence this effect. Another concern regarding C3 is the abundance of C3 in the circulation and also that it is produced locally in many tissues, each of which could make inhibition difficult.

4. Terminal Pathway. Blocking of C5 has been used clinically for more than a decade. Thus, C5 is a main
candidate also in the future. Additional targets to block are C5a, which will block both C5aR1 and C5aR2 effects. Alternatively, C5aR1 is a very good candidate because this receptor is known to signal production of substantial proinflammatory mediators. There is evidence that C5aR2 counteracts C5aR1, but lack of specific reagents to block C5aR2 are so far missing. The last target in the terminal pathway is to specifically block the assembly of C5b-9, thus preventing membrane attack complex formation but still having C5a released. This could be obtained by blocking, for example, C6 or C7. After C8 has bound, the leakage process through the membrane starts.

B. Reagents to Be Used

1. Monoclonal Antibodies and Their Derivatives.

Over the last 3 decades, monoclonal antibodies have made a dramatic transformation from scientific tools to powerful human therapeutics (Buss et al., 2012). To overcome the inherent immunogenicity and reduced effector function of murine monoclonal antibodies in man, chimeric mouse-human antibodies were developed. To further improve monoclonal antibody properties, humanized mAbs (suffix: -zumab) that are used for anti-C5 inhibition were developed by grafting just the murine hypervariable regions onto a human antibody framework, resulting in a molecule that is more than 95% human. These types of antibodies appear to overcome most of the inherent immunogenic problems of murine and chimeric monoclonal antibodies.

The advent of in vitro phage display technology and the generation of various transgenic mouse strains expressing human variable domains enabled the generation of fully human mAbs (suffix: -umab). Both humanized and fully human monoclonal antibodies have significantly reduced immunogenic potential and show properties like human endogenous IgG (Nelson et al., 2010). The disadvantage of using this technique is that it is quite laborious and expensive to manufacture in large scale.

As discussed above, an anti-C5 monoclonal antibody that blocks the cleavage of C5 and thus the generation of proinflammatory C5a and membranolytic C5b-9 has been the first complement-targeted monoclonal antibody therapeutic and has been tremendously effective for many patients with PNH and aHUS (Legendre et al., 2013). More recently (2017) the US FDA approved anti-C5 use in refractory generalized myasthenia gravis (Howard et al., 2017; Dhillon, 2018) and also in neuro-myelitis optica spectrum disorder (Pittock et al., 2019). Two caveats of this otherwise successful treatment are that first, there is a small population of patients (3% of the Japanese population) that have a C5 polymorphism that is not recognized by the anti-C5 antibody eculizumab (Nishimura et al., 2014). Multiple other anti-C5 antibodies, as referred to above, that can provide both improvements on eculizumab and broader patient efficacy are in phase I and II trials at this time (Mastellos et al., 2019; Röth et al., 2020). Thus far, a lack of C5a generation and thus C5aR1 signaling have not been harmful for extended clinical use of the C5 inhibitor antibody. An anti-C5a monoclonal antibody has been shown to be well tolerated in humans (Giamarellos-Bourboulis et al., 2020) and is being proposed as a complement therapeutic (e.g., in COVID-19), but it remains to be shown whether this will be an efficient therapeutic in the future (Vlaar et al., 2020).

In development for some time, strategies available to suppress deleterious overactivation of the complement cascade include the use of a specific monoclonal antibody to target the injured or diseased site fused to an inhibitor of complement activation. Predominantly, the complement inhibitor targets deposited C3b to suppress the amplification process and thus the generation of excessive downstream detrimental complement activation products. As mentioned above, an example of this is B4/Cr1-related protein Y, which has been shown to be effective in stroke and reperfusion injury in animal studies.

2. Small Molecular Peptides and Peptidomimetics.

Protein-protein interactions are well recognized as mediators of a plethora of processes in biologic systems and are vitally important in the progression of many disease states (Robertson and Spring, 2018). These interactions can be inhibited but also stabilized by peptides and peptidomimetics and thus are a potentially tool for drug discovery. Indeed, more than 60 different types of peptides have been approved for clinic use. On the other hand, chemical structures that mimic the effect of peptides are also an attractive approach. Peptidomimetics can respond to peptide limitations by displaying higher metabolic stability, good bioavailability, and enhanced receptor affinity and selectivity. Various synthetic strategies have been developed over the years to modulate the conformational flexibility and the peptide character of peptidomimetic compounds.

Both strategies have advantages over antibodies generally in cost but, more importantly, the potential access to tissue, including the CNS. In animal models of Alzheimer disease (Fonseca et al., 2009), Huntington disease (Woodruff et al., 2006), immune complex–associated blood brain barrier disruption (Mahajan et al., 2015), stroke (Garrett et al., 2009), and traumatic brain injury (Sewell et al., 2004), C5aR1 antagonists have suppressed pathology, inflammation (Fig. 5A), and functional loss. Treatment with PMX53 also reduced the time length of a seizure and the likelihood of a second seizure in a kindling model of epilepsy, suggesting that blocking C5aR1 can have antiepileptic effects (Benson et al., 2015).

A detailed comparison of the available peptide and small-molecule C5aR1 antagonists has recently been published by Woodruff et al. (Li et al., 2020) as well as thorough pharmacokinetics of the two commonly used
antagonists PMX53 and PMX205 that will facilitate additional testing in mouse models (Li et al., 2020). Drug-receptor co-crystallization has identified drug-binding sites and resulting conformational changes (Liu et al., 2018). Most encouraging is that this approach has found to be safe in human trials (Vergunst et al., 2007; Garrett et al., 2009; Bekker et al., 2016; Jayne et al., 2017; Liu et al., 2018). Indeed, a small-molecule C5aR1 antagonist (Avacopan; CCX168) has completed a successful phase 3 trial for AAV and is under review by the US FDA for approval for oral treatment of this disease. Targeting C5aR1 leaves the beneficial functions of other complement components, such as C1q, C3b, and C5b-9, intact while having positive effects on pathology and function, including cognitive behavior in models of neurodegenerative disorders (Fonseca et al., 2009; Benson et al., 2015; Hernandez et al., 2017).

APL-2 and AMY-101 are two derivatives of a family of cyclic peptides that inhibit C3 amplification and are currently in clinical trials as either combination therapy with eculizamab or as a monotherapy, as suppression of C3 cleavage would also prevent membrane attack complex formation. Peptides and chemical peptidomimetics are small molecules usually with short half-lives. The advantage is that they may be useful for diseases in organs with membranes limiting passage of large molecules and for short-time need of treatment, whereas the disadvantage would be long-time systemic treatment. The latter could be solved by conjugating them to proteins increasing the half-life, as described under item 4 below.

3. Aptamers. Aptamers are single-stranded DNA or, more frequently, RNA molecules that can bind specifically to a target and neutralize the function of a protein. The molecular mass is between 5 and 15 kDa, and the half-life is in the range of a few hours. They are potential reagents to inhibit complement activation, and an aptamer blocking C5a (Müller-Redetzky et al., 2017) has also been developed to block C5a (Yatime et al., 2015; Hyzewicz et al., 2017) and C5aR1 (Kumar, 2018). A mouse study showed protection against pneumococcal sepsis using an aptamer blocking C5a (Müller-Redetzky et al., 2020), but the aptamers have not yet reached the clinic.

4. Recombinant Proteins and Conjugates. Recombinant proteins are frequently made from the host’s own complement regulatory proteins. CR1 is co-factor for factor I and a very efficient inhibitor of C3 by enhancing inactivation of C3b. It is present in large amounts on a number of cells, including red cells. It exists in small amounts in a soluble form, sCR1 (TP10), which lacks the transmembrane portion. A breakthrough in the field of complement inhibition was when sCR1 was produced recombinantly and could be used experimentally, wherein it was shown to reduce the size of myocardial infarction in mice (Weisman et al., 1990). When conjugated to the E-selectin ligand sialyl Lewis x (sCR1/sialyl Lewis x; TP20), the effect was increased when treating mice with experimental stroke (Huang et al., 1999). TP10 was used in some clinical trials with benefits in subgroups (Li et al., 2006). Large recombinant molecules are expensive to produce, and TP10 never came to routine clinical use. Other regulators like CD55, CD46, and CD59 have also been made recombinantly as possible drugs and tested experimentally (Verbakel et al., 2000).

Another approach is to take only a small part that is the active domain in the protein and conjugate this to a specific targeting molecule, which leads to binding to the actual site that needs to be treated. TT30 consists of a small part of factor H, “mini-factor H,” which contains the active C3b-binding site of factor H (Nichols et al., 2015) fused to CR2, which binds C3d deposited in the tissue (Risitano et al., 2012). This is a prototypic reagent for targeting tissues where complement is already ongoing and C3d is deposited (Fridkis-Hareli et al., 2011; Rich et al., 2016).

Thus, this will bind to any tissue where C3d is exposed and reduce the amount of free inhibitor in the fluid phase, which is a major advantage to preserving the activity of complement in the fluid phase. Another similar approach is to conjugate the inhibitor to a tissue-specific protein targeting (e.g., the synovium, to treat arthritis) (Macor et al., 2012). Finally, a small fragment of CR1 has been conjugated to an endothelial cell membrane–binding reagent (Microcept, ATP070) that will target damaged endothelium, typically after ischemia/reperfusion and is thus not organ-specific and does not depend on ongoing complement activation but is intended to prevent activation whether it is a whole body or local damage (Souza et al., 2005).

In other cases, it is important to keep the inhibitor in the fluid phase, such as in trauma, sepsis, or other systemic inflammatory diseases. Then a small inhibitor can be conjugated to an inert, nontargeting molecule just for the purpose of increasing its half-life (e.g., an Fc fragment), and the half-life can be increased from a few minutes to days (Hepburn et al., 2008).

C. Routes of Distribution

1. Locally. Depending on the disease, its location and accessibility, and the structure of the drug, there are several alternatives for local treatment. First, a cream could be administered, such as, for example, for the highly complement-dependent disease periodontitis (Hajishengallis and Lambris, 2012), wherein inhibition both at C3 and C5 level has been shown effective (Abe et al., 2012; Kajikawa et al., 2017); toothpaste could be an alternative. Liniment might be an alternative to treat psoriasis, as it is postulated to be partly complement-dependent (Schonthaler et al., 2013). Complement is important for several eye diseases (see separate section), and, depending on the
drug, local injection might be required, but with small molecules that might penetrate, eye drops are obviously simple and uncomplicated. Obvious advantages with local treatment are that it is low-cost and, except for injection into the eye, the treatment can be done by the patient or a relative.

Local application can also be an alternative for systemic treatment. Subcutaneous injection is already well established for continuous release for long-time treatment and can be established at home. Finally, in some cases, particularly in children, nasal spray might be an alternative and obviously the simplest one.

2. Intravenously. Until now, intravenous treatment has been the usual way of administration. Systemic adverse effects might occur, and in most cases, the patient needs to go to the hospital. Thus, the costs are substantially higher than the local alternatives (i.e., subcutaneous injection).

3. Orally. The main goal for any producer of a complement inhibitor is to succeed with a pill. Great efforts should be done to obtain this endpoint, as it is definitely the best solution. Promising preclinical data exist, but oral administration has not come to routine use yet. The problems with short half-life and bioavailability must be resolved to produce the effective “pill.”

VI. Assays for Diagnostics and for Treatment Follow-up

A. Principles of Complement Assays

1. Detection of Complement Activation. There are a number of different assays to evaluate the complement system (for review, see Mollnes et al. (2007) and Harboe et al. (2011)). Here we will only describe the two assay types that are most important to understand for those who are treating patients with complement inhibitors in clinical routine. The first is an assay that detects the degree of complement activation in vivo. A number of assays that detect specific activation products are available, but the TCC is a valuable biomarker since it indicates that C5 is activated and that both C5a and C5b-9 are formed. This can be detected in plasma as sC5b-9 in a simple ELISA based on a monoclonal antibody reacting with a neoepitope exposed in C9 when C9 is incorporated in the complex but not present in the native C9 molecule (Fig. 6). The advantage of using this assay is that this complex has a relatively long half-life (1 hour) as compared with the C5a molecule with a very short half-life (1 minute). Furthermore, it will detect activation of the whole cascade to its very end. It is critically important that the blood samples are obtained using EDTA and that plasma is collected and snap-frozen to −70°C (Frazer-Abel, 2018; Proházska et al., 2018).

Increased levels of sC5b-9 in plasma indicate a disease process wherein complement is involved, and the condition may be a possible candidate for complement inhibition. However, complement might be activated locally in organs without increased activation products in plasma (i.e., a normal sC5b-9 level does not exclude complement activation in tissue). Thus, biopsies may be the ultimate test for diagnosis in which activation product can easily be detected with fluorescence microscopy, as illustrated in Fig. 5B. Plasma sC5b-9 might also be helpful for evaluation of the effect of therapy in a patient with originally high levels before treatment is started. The ultimate test for such an evaluation is, however, the screening for total complement activity, as described below.

2. Screening for Total Complement Activity. This group of tests measures the functional activity of the whole complement cascade. Previously, they were based on a complement hemolytic assay, but this has now been gradually replaced by more stable and sensitive enzyme immunoassays, like the total complement screen assay (also called Wieslisa) (Fig. 7). In contrast with the sC5b-9 assay described above, this assay tests complement function in vitro and must therefore be done in serum (fresh or freshly stored at −70°C). The detection principle is as for the sC5b-9 ELISA, but the design is different. The wells are coated with specific activators of the three pathways, and if all components are present and active, the C5b-9 complex will assemble in the well and can be detected with the same anti-C9 neoepitope antibody. If the complement system is normal, it will be detected as 100% activity in the wells (Fig. 7A).

If a terminal component or C3 is genetically defective, the C5b-9 complex will not be formed, as for example in a patient deficient in C5 (Fig. 7B). Here all the three pathways will show 0% activity since the detection antibody will not recognize a C9 neoepitope. If the screen is negative in one or more of the assays, further testing has to be done to find the deficient component. If
a patient is receiving a complement inhibitor (e.g., a C5 blocker), this will give the same result as for the patient who is C5 genetic deficient if all C5 molecules are blocked (Fig. 7C), and the activity will be 0%. For testing the effect of a C5 inhibitor, it is not necessary to use all the pathway assays. Either the classical or the alternative assay can be used to follow the patient, as the activity will gradually increase from 0% to 100% when C5 activity starts to increase, and finally all of the inhibitor is gone (Fig. 7, C and D). The half-life can vary substantially dependent upon which disease condition and stage of disease the patient presents. With these test results, an individualized treatment regimen can be established.

VII. Consequences of Therapeutic Complement Inhibition

A. Safety and Adverse Effects

1. Safety and Adverse Effects of Established C5 Inhibition. Eculizumab has been the only complement inhibitor approved for clinical treatment until 2020. It
was approved in 2007 for paroxysmal nocturnal hemoglobinuria. These patients have life-long treatment, and treatment has shown a high safety. In a study of 196 patients over 66 months, the drug was safe and well tolerated with no evidence of cumulative toxicity (Hillmen et al., 2013). Four deaths were not related to the treatment and no Neisseria infection was observed. The same was found in a Japanese study of patients with PNH (Kanakura et al., 2013). A systematic review based on 12 databases showed safe treatments without deaths or Neisseria infections (Rathbone et al., 2013). A number of other studies confirm the safety. However, in a large study including 1321 patients followed over 5 years, with patients who were both pediatric and adults, three cases of Neisseria infection was observed and one with a fatal outcome (Rondeau et al., 2019).

Finally, a 10-year observation study comprising 28,518 patient-years confirmed the safety but also the risk of neisserial infection. A few other infections were observed, but whether they were related to treatment was uncertain (Socié et al., 2019). One fatal Neisseria infection was reported with a nongroupable strain has been reported (Nolfi-Donegan et al., 2018). Thus, C5 inhibition with eculizumab is a safe treatment with virtually no adverse effects, but the small risk of getting a Neisseria infection should always be brought in mind.

2. Potential Adverse Effects of New Inhibitors. When new inhibitors come to clinical use, one should certainly be aware of other possible adverse effects. It cannot be excluded that inhibition at the initial recognition phases of all three pathways or inhibition at the level of C3 can lead to increased risk of other infections due to reduced opsonization. Short-time treatment when the patient is under observation should be of less concern and can even be the most effective treatment in certain acute phases.

B. Efficacy of Treatment

1. Fully Complement-Dependent Diseases. As described above under the indications for complement inhibition, the treatment is very efficient and also lifesaving when the disease is mainly complement-dependent, such as with paroxysmal nocturnal hemoglobinuria and aHUS. As mentioned, there are a number of disease candidates in which complement plays a major (although not the whole) role in the pathophysiology. Controlled clinical trials should be performed before new diseases are added to the list. Occasionally, off-label use may be acceptable in patients who are critically ill in which it is reasonable to suggest complement-mediated pathophysiology, in particular when the disease is so rare that it is impossible to perform clinical trials with sufficient power.

2. Diseases with Complex Pathophysiology. Complement activation might be involved as a smaller part of the pathophysiology and in such complex conditions that the challenge will be to use experimental model and clinical trials with a combined strategy to examine whether complement inhibition may partly reduce disease activity.

VIII. Conclusions and Future Perspectives

As stated in the introduction, the aim of this study was to provide some background on the complement system and its physiologic activity with highlight on the mechanisms leading to complement dysfunction, subsequent tissue damage, and disease. We have given priority to these principles rather than include lists of ongoing clinical trials or of all the potential inhibitors that are available as therapeutic candidates. We recommend that the particularly interested reader use “complement” as a search key in https://clinicaltrials.gov.

We hope this review has provided clear information to the reader for considering complement as an interesting and well documented target for treatment of many diseases in the clinic rather than merely being “an elegant model system.” Furthermore, we emphasize the importance of increasing the research activity in this area to provide the basis for opportunities for clinical trials in the near future. It is reasonable to suggest that with an increase in the number of drugs targeting the complement system, the list of conditions to be treated with complement-modulating approaches will increase in parallel.

Authorship Contributions

Wrote or contributed to the writing of the manuscript: Garred, Tenner, Mollnes.

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