Abstract | Monoclonal antibodies (mAbs) are appealing as potential therapeutics and prophylactics for viral infections owing to characteristics such as their high specificity and their ability to enhance immune responses. Furthermore, antibody engineering can be used to strengthen effector function and prolong mAb half-life, and advances in structural biology have enabled the selection and optimization of potent neutralizing mAbs through identification of vulnerable regions in viral proteins, which can also be relevant for vaccine design. The COVID-19 pandemic has stimulated extensive efforts to develop neutralizing mAbs against severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2), with several mAbs now having received authorization for emergency use, providing not just an important component of strategies to combat COVID-19 but also a boost to efforts to harness mAbs in therapeutic and preventive settings for other infectious diseases. Here, we describe advances in antibody discovery and engineering that have led to the development of mAbs for use against infections caused by viruses including SARS-CoV-2, respiratory syncytial virus (RSV), Ebola virus (EBOV), human cytomegalovirus (HCMV) and influenza. We also discuss the rationale for moving from empirical to structure-guided strategies in vaccine development, based on identifying optimal candidate antigens and vulnerable regions within them that can be targeted by antibodies to result in a strong protective immune response.

Since the development of the hybridoma technology enabling the generation of monoclonal antibodies (mAbs) by Köhler and Milstein in the 1970s, mAbs have become a key class of drugs for cancer and immune disorders, with the 100th antibody-based therapeutic recently gaining US Food and Drug Administration (FDA) approval. However, although harnessing antibodies to combat infectious diseases has a history stretching back more than a century to the applications of serum conferring protection against diphtheria toxin, only a small number of mAb drugs are used to treat or prevent infectious diseases. At the time of writing in 2022, six mAbs targeting pathogens have so far been granted full approval by the FDA (Table 1), for indications including prevention of respiratory syncytial virus (RSV) infection, prevention and treatment of anthrax infection, prevention of recurrence of Clostridioides difficile infection and the treatment of Ebola virus (EBOV) infection.

The rapid spread of COVID-19 in 2020 led to intense efforts to develop neutralizing mAbs that target severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) for the treatment and prevention of COVID-19. As a result, more than 20 mAbs entered clinical development. So far, several of these mAbs have received emergency use authorization (EUA) from the FDA (Table 2) and other regulatory agencies worldwide, and more authorizations are anticipated. Although vaccines have been the mainstay of efforts to tackle COVID-19, mAbs can provide an important contribution for vulnerable populations before or after exposure to SARS-CoV-2, such as people who are immunocompromised or people with mild to moderate COVID-19 who are at high risk of developing severe disease. The mAbs under development providing broadly neutralizing activity against coronaviruses could also be instrumental for preparedness against future pandemics. Furthermore, the momentum built and the knowledge gained from the development of mAbs for SARS-CoV-2 may help accelerate the development of mAbs to combat other infectious diseases.

In this Review, we first provide an overview of the technologies for the discovery and engineering of mAbs to target pathogens, with a focus on viruses. We then describe the progress in the development of mAbs against a range of viruses, including SARS-CoV-2, RSV, Ebola, cytomegalovirus (CMV) and influenza. Efforts to harness mAbs to combat bacterial infections have been reviewed elsewhere and are summarized briefly in Box 1. Finally, we also discuss how rapid mAb discovery combined with structural vaccinology can support the development of vaccines and therapeutic mAbs.
Antibody characteristics and engineering

**Antibody structure and function.** Antibodies are natural biomolecules generated by plasma cells or stimulated memory B cells after a pathogen infection or vaccination. Structurally, they are Y-shaped heterodimers composed of two light chains of 25 kDa each and two heavy chains of at least 50 kDa, depending on the immunoglobulin isotype. The heavy and light chains are linked by multiple disulfide bridges and non-covalent interactions (Fig. 1b), with variations in the number of interactions and bridges depending on the immunoglobulin isoatypes.

Antibodies can also be divided into functional components (Fig. 1b). The two fragment antigen-binding domains (Fab) bind to and neutralize pathogens. These are linked to the crystallizable fragment (Fc) domain by a hinge region that gives the Fabs a large mobility such as ADCC and CDC (Fig. 1c), IgG2 and IgG4 are poor CDC activators and IgG3 is a potent CDC activator. Most therapeutic mAbs in clinical use or development against infectious diseases are of the human IgG1 isotype, which has affinity for activating FcyRs but also exhibits binding to the inhibitory FcRnR, thereby limiting protective Fc effector activities.

Antibodies as antivirals. For more than a century, passive immunization with monoclonal or polyclonal antibodies has been used in the treatment and prevention of infectious diseases, particularly in individuals with immunodeficiencies or individuals for whom vaccination is contraindicated.

Antibodies can combat viral infections through several mechanisms. First, antibodies can prevent viral glycoproteins of enveloped viruses or the protein shell of non-enveloped viruses from binding to the target host cells. These viral proteins have two major functions in the viral life cycle: binding to cellular receptors and mediating the fusion of viral and cellular membranes (in the case of enveloped viruses) or penetration into the cytosol (in the case of non-enveloped viruses). For example, the entry of SARS-CoV-2 into host cells is mediated by the interaction between the viral spike (S) glycoprotein and the angiotensin-converting enzyme 2 (ACE2) receptor on the host cell surface. ACE2 is expressed on cells of the respiratory system, gastrointestinal tract and endothelium. The spike–ACE2 interaction can be blocked by antibodies targeting the spike receptor-binding domain (RBD).

**Table 1 | FDA-approved mAbs for infectious disease indications**

| Drug (brand name; company) | Target | Format | Technology | Indication | Year of FDA approval |
|----------------------------|--------|--------|------------|------------|---------------------|
| Palivizumab (Synagis; MedImmune/AbbVie) | RSV | Humanized IgG1 | Hybridoma | Prevention of RSV infection | 1998 |
| Raxibacumab (ABthrax/Anthrin; GlaxoSmithKline/ Human Genome Sciences) | Bacillus anthrasis PA | Human IgG1 | Human scFv phage display library | Anthrax infection | 2012 |
| Bezlotoxumab (Zirplava; Merck & Co.) | Clostridiodes difficile enterotoxin B | Human IgG1 | Transgenic mice | Prevention of C. difficile infection recurrence | 2016 |
| Obiltoxaximab (Anthem; Elusys Therapeutics) | B. anthraxis PA | Chimeric IgG1 | Hybridoma | Prevention of inhalational anthrax | 2016 |
| Ibalizumab* (Trogarzo; TaiMed Biologics) | CD4 receptor (domain 2) | Humanized IgG4 | Mice | Treatment of HIV-1 infection | 2018 |
| Ansvivimab (Ebanga; MedImmune/Ridgeback Biotherapeutics) | Ebola glycoprotein | Human IgG1 | Human | Prevention and treatment of Ebola infection | 2020 |
| Alitivimab, mavitivimab and odesivimab (Immazeb; Regeneron Pharmaceuticals) | Ebola glycoprotein | Human IgG1 | Transgenic mice | Prevention and treatment of Ebola infection | 2020 |

FDA, US Food and Drug Administration; mAb, monoclonal antibody; PA, protective antigen; RSV, respiratory syncytial virus; scFv, single-chain variable fragment. *Antibody with a host target, rather than a pathogen target.

**Fc gamma receptors**
Humans express six Fc receptors (FcγRs) that modulate effector cells upon binding to IgG. FcγRI, FcγRIIA, FcγRIIB, FcγRIIIA and FcγRIIB are activating receptors, whereas FcγRIIB is inhibitory.

**Antibody-dependent cellular cytotoxicity** (ADCC). A mechanism of cell-mediated immune defence whereby an effector cell of the immune system actively lyses a target cell, whose membrane-surface antigens have been bound by specific antibodies.

**Complement-dependent cytotoxicity** (CDC). The complement system is a network of proteins that form an important part of the immune response by enhancing the opsonization of pathogens, cell lysis and inflammation. CDC is a mechanism of complement-mediated immune defence in which an antibody bound to its antigen activates the complement cascade.

**Glycoproteins**
Proteins with oligosaccharide chains (glycans) covalently attached to amino acid side-chains. Virus surface glycoproteins embedded in the membrane often have a role in interactions with host cells, including receptor binding, and are commonly targeted by host antibodies.
which inhibits viral infection, as discussed later in the Review (Fig. 1b).

Antibody effector functions mediated by engagement with complement protein C1q or FcγRs on leukocytes can also be involved in combating viral infections (Fig. 1c). Complement activation by antibodies leads to direct lysis of the virus and/or the infected host cell, and antibodies can also promote or induce phagocytosis, or trigger the release of toxic chemicals, such as cytokines or reactive oxygen species. For instance, it was recently shown that Fc effector functions are required for optimal protection by mAb therapy for SARS-CoV-2; when given after infection, intact mAbs reduced the SARS-CoV-2 burden and lung disease in animals better than loss-of-function Fc variant mAbs by mitigating inflammation and improving respiratory mechanics.

Conversely, in rare cases, suboptimal binding of antibodies to virions can facilitate viral pathogenesis through a process known as antibody-dependent enhancement (ADE), in which recognition of virion–antibody complexes by FcγRs enhances viral entry into host immune cells. ADE was first observed with Dengue virus in the presence of sub-neutralizing antibody concentrations. For example, when the level of maternal antibodies against Dengue virus in newborns wanes, some individuals will experience an interval during which their antibody level will drop below its protective capacity, leading to severe disease following infection. Dengue virus co-circulates as four serotypes, and the increased severity of some secondary infections is thought to be due to enhancement of viral entry by pre-existing antibodies generated following a primary infection with a different serotype that are not able to neutralize the second serotype. Enhanced disease was also observed after vaccination with a formalin-inactivated RSV vaccine in infants. The risk of ADE can be reduced by engineering the antibody Fc domain to reduce binding to FcγRs, as noted below.

The first passive immunization approaches used serum derived from animals actively immunized with an antigen such as diphtheria toxin, but such approaches come with the risk of provoking an immune response against non-human antibodies. These risks can be mitigated by using blood from people who have recovered from an illness as the source of antibodies, known as convalescent plasma therapy (CPT). Although CPT has been historically successful in combating infections, it has proved to be an inconsistent tool, as antibody responses between individuals are highly variable. CPT also has safety risks, including allergic reactions and low risk of infections by other viruses such as HIV, hepatitis B and hepatitis C. Furthermore, widespread use of CPT in the context of a pandemic such as COVID-19 would depend on the availability of a sufficient number of plasma donors and facilities for appropriate processing. Overall, the role of CPT in such cases may be restricted

Table 2 | Selected mAbs marketed or in late-stage clinical studies for COVID-19

| Drug (brand name; company) | Origin | Engineering | Status* | Omicron VOC neutralization | Ref. |
|----------------------------|--------|-------------|---------|--------------------------|------|
| Casirivimab and imdevimab (Ronapreve; Regeneron Pharmaceuticals) | Genetically humanized mice and B cells from a convalescent patient infected with SARS-CoV-2 | Unmodified | EUA granted by FDA for treatment and prevention of COVID-19 in 2020 | –, in vitro study | 118 |
| Bamlanivimab and etesevimab (NA; AbCellera/Eli Lilly) | B cells from convalescent patients infected with SARS-CoV-2 | Unmodified (bamlanivimab); LALA modification in Fc domain to extend half-life (etesevimab) | EUA granted by FDA for treatment and prevention of COVID-19 in 2021 | –, in vitro study | 124 |
| Sotrovimab (Xevudy; Vir Biotechnology/GlaxoSmithKline) | B cells from an individual infected with SARS-CoV | LS modification in Fc domain to extend half-life | EUA granted by FDA for the treatment of mild to moderate COVID-19 in 2021 | ++/−, in vitro and clinical studies | 78 |
| Tixagevimab and cilgavimab (Evusheld; AstraZeneca) | B cells from convalescent patients infected with SARS-CoV-2 | YTE and TM modifications in Fc domain to extend half-life and reduce effector function, respectively | EUA granted by FDA for pre-exposure prophylaxis of COVID-19 in 2021 | ++/−, in vitro study | 129 |
| Bebtelovimab (NA; AbCellera/Eli Lilly) | B cells from convalescent patients infected with SARS-CoV-2 | Unmodified | EUA granted by FDA for treatment of mild to moderate COVID-19 in 2022 | ++++, in vitro study | 131 |
| Regdanvimab (Regkirona; Celltrion) | B cells from convalescent patient infected with SARS-CoV-2 | Modifications to Fc domain to reduce effector function | Approved in Republic of Korea and EU | –, in vitro study | 130 |
| Amubavimab and romlusevimab (NA; Brii Biosciences) | B cells from convalescent patients infected with SARS-CoV-2 | YTE modification in Fc domain to extend half-life | Approved in China; ELIA requested | ++/−, in vitro and clinical studies | 327 |
| Adintrevimab (NA; Adagio Therapeutics) | B cells from convalescent patients infected with SARS-CoV-2 | LALA modification in Fc domain to extend half-life | Phase II/III | ++/−, in vitro study | 328 |

All monoclonal antibodies (mAbs) listed are known to target the receptor-binding domain (RBD) of the spike (S) glycoprotein of severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2). Fc, crystallizable fragment; NA, not available; –, no neutralizing activity; ++/−, partial neutralizing activity; ++++, potent neutralizing activity. Information on status is with regard to the initial US Food and Drug Administration (FDA) emergency use authorizations (EUAs), unless the product was initially further advanced in other regions or countries. *EUA since withdrawn owing to likely ineffectiveness against the Omicron variant of concern (VOC).
Monoclonal antibodies as antibacterial agents

Antimicrobial resistance is one of the top ten global public health threats facing humanity according to the World Health Organization (WHO). Although traditional small-molecule drug discovery still dominates the landscape for antibacterial solutions to antimicrobial resistance, the prophylactic or therapeutic use of monoclonal antibodies (mAbs) for bacterial infection could have an important role. One of the main advantages of these biologics is their low toxicity, as bacterial virulence proteins are targeted instead of proteins required for survival, therefore avoiding the disruption of the microbiome and, potentially, limiting the development of resistance.

So far, three antibacterial mAbs have been approved for the treatment and prophylaxis of bacterial infections. Raaxibucumab (ABThrax/Antihin) and obiltoxaximab (Anthim) are both mAbs targeting the protective antigen (PA) component of the lethal toxin of Bacillus anthracis and are approved to treat inhalation anthrax due to B. anthracis. Bezlotoxumab is a mAb that binds to Clostridoides difficile enterotoxin B (CdtB). This product does not protect against or treat initial C. difficile infection but, rather, is used to reduce the recurrence of infection, which is often seen with C. difficile.

Nevertheless, a naive human single-chain variable fragment (scFv) phage display library was used to develop a potent antitoxin mAb for anthrax, raxibacumab, which was approved in 2012 (BOX 1 and TABLE 1).

A second targeted approach developed subsequently is the direct isolation of antigen-specific memory B cells based on their capacity to bind fluorescent bait antigens, followed by identification of the mAbs they produce. The memory B cells can originate from the plasma of convalescent patients, or from transgenic mice carrying human immunoglobulin loci that produce fully human antibodies in response to immunization with a target antigen. This approach has been particularly successful in the isolation of broadly neutralizing antibodies (bNAbs) targeting the CD4-binding site in the V1/V2 and V3 regions of gp120 and the membrane-proximal external region of gp41 of HIV-1. Furthermore, mAbs against the hepatitis B virus viral S antigen (HBsAg) and the SARS-CoV-2 spike protein have also been obtained, as discussed below.

The major limitation of targeted approaches is that target antigens must be known in advance because the selection process is based on binding affinity to the purified antigen rather than neutralization potency. Target-agnostic approaches present a viable alternative when limited information is available on the pathogen to be neutralized. Various methods to obtain single cell cultures of memory B cells or plasma cells have been described. Memory B cell immortalization using Epstein—Barr virus remains an attractive method because of its limited cost. A limitation of the Epstein—Barr virus approach is the suboptimal immortalization of B cells, which plateaus at approximately 35%. However, the development of single cell cultures without the need for B cell immortalization can overcome this limitation and has been used for the identification of antibodies against pathogens such as group A and group B influenza A viruses and BK/JC polyomaviruses. Recently, a functional organotypic system for antibody generation has been reported. The organoid recapitulates germinal centre features in vitro, such as the production of antigen-specific antibodies with affinity maturation and class-switch recombination from human tonsils, and this is a promising step forward for the field of mAb development.

In all cases, once identified, the mAb candidates must be sequenced for further recombinant expression. Cloning and expression of individual antibodies was traditionally labour-intensive, and throughput was largely limited to a few hundred clones. However, recent technological advances using nanofluidic devices have considerably increased the throughput of this approach. In addition, advances in next-generation sequencing have enabled high-throughput screening and sequencing of paired antibody repertoires. Currently, isolated B memory cells or plasma cells are injected into microfluidic devices (such as the 10x Genomics platform), generating droplets containing a single cell and lysis buffer with microbeads covered by barcoded primers to generate cDNA encoding VH and VL sequences. These approaches allow the discovery of antibodies that are potentially useful as therapeutics, in addition to the
Opsonized is a state of a pathogen in which antibodies or complement factors are bound to its surface.

The possibility of studying the human antibody repertoire at an unprecedented resolution⁶⁷, and could be deployed as emergency response platforms for investigating mAbs from the blood of people who have recovered from emerging viral infections⁶⁸.

Recently, the LIBRA-seq (linking B cell receptor to antigen specificity through sequencing) methodology was developed for high-throughput mapping of paired heavy-chain and light-chain B cell receptor sequences to their cognate antigen specificities⁶⁹. In this approach,

- **Opsonized**
  - A state of a pathogen in which antibodies or complement factors are bound to its surface.

**Fig. 1 | Mechanism of action of monoclonal antibodies during viral infection.**

- **a** | Overview of monoclonal antibody (mAb) structure; heavy chains shown in yellow and red, light chains in blue and green.

- **b** | Binding of severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) to the angiotensin-converting enzyme 2 (ACE2) receptor on host cells is mediated by the viral spike (S) protein, which comprises an S1 subunit (including a receptor-binding domain (RBD) and an amino-terminal domain (NTD)) and an S2 subunit. Priming of coronavirus spike proteins by host cell proteases such as the transmembrane serine protease TMPRSS2 through cleavage at S1/S2 and S2’ sites (see Fig. 3) is essential for viral entry. Therapeutic antibodies and antibodies elicited by vaccination that bind to the RBD or NTD of the spike protein can block viral binding to ACE2, or block fusion between viral and cellular membranes (see Fig. 3). c | Effector functions of antibodies. mAbs can facilitate target cell death via complement fixation and membrane attack complex (MAC) activation, which is known as complement-dependent cytotoxicity (CDC). Antibody-dependent cellular cytotoxicity (ADCC) is a mechanism of cell-mediated immune defence whereby an effector cell (natural killer cell, macrophage, neutrophil or eosinophil) of the immune system actively lysed a target cell, whose membrane has been bound by specific antibodies. Natural killer cells release cytotoxic factors (perforin and proteases known as granzymes) that cause death of the infected cell. Antibody-dependent cellular phagocytosis is the mechanism by which antibody-opsonized target cells activate FcγRs on the surface of macrophages to induce phagocytosis, resulting in internalization and degradation of the target cell through phagosome acidification. Fab, fragment antigen-binding domain; Fc, crystallizable fragment; Fv, variable fragment; VH, variable heavy; VL, variable light.
**Fig. 2 | Antibody discovery approaches.** **a** | Phage bio-panning is based on a library of phages that contain genes coding for variable heavy (VH)/variable light (VL) domains, leading to production of encoded antibodies on phage surfaces. Selection of antibodies produced by phages involves immobilization of the ligand of interest on a solid support (spike (S) protein of severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) shown), followed by applying the phage display library to immobilized ligand to allow binding of specific variants. To eliminate adherent non-binders, multiple rounds of washing are usually performed, and remaining bound phages are eluted and re-amplified. **b** | Recombinant antigens conjugated to a fluorescent marker are incubated with class-switched memory B cells and sorted according to their capacity to bind the antigen of interest (such as the S protein of SARS-CoV-2) by fluorescence-activated flow cytometry, followed by identification of the antibodies they produce. **c** | Target-agnostic approaches using single B cell culture. Single B cells are seeded on a feeder layer in the presence of a cytokine mix and a Toll-like receptor (TLR) activator. Culture supernatant is screened for neutralization activity and clones of interest are retrieved and sequenced. **d** | Workflow for organoid reconstitution from human tonsils to develop an in vitro system that recapitulates key germinal centre features, including production of antigen-specific antibodies, somatic hypermutation and affinity maturation. **e** | Workflow for single cell immune profiling. B cells are encapsulated with barcoded gel beads in a single partition and undergo reverse transcription followed by PCR. Each cDNA is barcoded from its individual cell of origin and processed for next-generation sequencing. Fab, fragment antigen-binding domain; mAb, monoclonal antibody; scFv, single-chain variable fragment. Panel a is adapted from Ref. 329, under a Creative Commons license CC BY 3.0.
B cells are mixed with a panel of DNA-barcoded antigens and both the antigen barcode(s) and B cell receptor sequences are recovered via single cell next-generation sequencing. This enabled the antigen specificity of thousands of B cells from two subjects infected with HIV to be mapped, and the predicted specificities were confirmed for numerous HIV, influenza and SARS-CoV-2-specific antibodies, including known and unknown bNAbs (Fig. 2a). Indeed, large antibody data sets can be analysed computationally to infer antibody sequences and binding modality, and this has emerged as a powerful method for the identification of structurally related antibodies from sequence databases. However, a limitation of the LIBRA-seq methodology, which is inherent to targeted approaches, is the need for a recombinant bait antigen that requires extensive validation.

**Antibody engineering.** Several regions of mAbs can be engineered to improve their therapeutic characteristics.

In addition to the variability induced by isotype usage, several mutations in the Fc domain have been identified to increase or decrease the effectiveness of ADCC and/or CDC. For example, etesevimab, an anti-SARS-CoV-2 mAb developed by Eli Lilly that has received an EUA for COVID-19, has been engineered to lack FcγRI and FcγRII-binding activity with L234A and L235A mutations (the LALA modification) in the Fc domain to reduce safety concerns over the potential to exacerbate disease through ADE mechanisms. Another example is provided by AZD7442, a cocktail of the anti-SARS-CoV-2 mAbs tixagevimab (AZD8895) and cilgavimab (AZD1061) developed by AstraZeneca, which has received an EUA for COVID-19. Both mAbs in the combination have engineered Fc domains including L234A/L235A/P331S substitutions (the TM modification) in the Fc domain to reduce safety concerns over the potential to exacerbate disease through ADE mechanisms. A single bsAb can therefore bind to two different epitopes or different sites on the same protein. A wide range of bsAb formats have been developed, particularly for oncology applications, but there have also been a few bsAbs investigated for infectious diseases. For example, a bsAb targeting both the receptor-binding site (RBS) of the Niemann-Pick C1 (NPC1) protein and a conserved surface-exposed epitope on the EBOV glycoprotein was shown to neutralize all known EBOVs by co-opting viral particles for endosomal delivery and conferred post-exposure protection against multiple EBOVs in mice. Another example, a bsAb for HIV-1, also illustrates the potential of engineering the hinge domain to provide flexibility of orientation and rotation to the Fab region in regard to the Fc fragment. Bispecific anti-Env neutralizing antibodies with an engineered IgG3 hinge domain to increase Fab domain flexibility demonstrated improved neutralization potency and enhanced in vivo protective activity in HIV-1-infected humanized mice.
Extending the multi-specificity concept further, trispecific antibodies engineered to interact with three independent HIV envelope determinants (the CD4-binding site, the membrane-proximal external region and the V1/V2 glycan) conferred complete immunity against a mixture of simian-human immunodeficiency viruses in non-human primates (NHPs), and showed higher potency and breadth than any previously described single broadly neutralizing mAb. One such agent, SAR441236, has entered phase I development (NCT03705169).

Other novel engineered antibody formats that could be applied in antiviral agents include small cameld VHHs (15kDa), known as nanobodies, that retain full antigen specificity, in contrast to mouse and human antibody-binding domains (50kDa). Furthermore, nanobodies possess extended complementarity-determining regions, enabling binding of epitopes that are not normally accessible to conventional antibodies, such as conserved viral domains that are often masked by glycan shields. Clinical trials demonstrated that they are safe and possess low immunogenicity. Interestingly, transgenic mice encoding 18 alpaca, 7 dromedary and 5 Bactrian camel VHH genes were shown to generate potent neutralizing nanobodies against SARS-CoV-2.

Finally, antibody mimetics such as designed ankyrin repeat proteins (DARPin) can provide high affinity and offer multi-specificity. For instance, a multi-DARPin (ensovibe) that binds simultaneously to all three units of the SARS-CoV-2 spike RBD developed by Molecular Partners and Novartis is in clinical trials (NCT04828161).

**Antibodies to combat viral infections**

**Coronaviruses, including SARS-CoV-2.** Coronaviruses are enveloped positive-sense single-stranded RNA viruses belonging to the Coronaviridae family. They can infect a wide variety of mammalian and avian species, causing respiratory and/or intestinal tract diseases. Human coronaviruses are major causes of the common cold and are responsible for up to 30% of mild respiratory tract infections and atypical pneumonia in humans. Four different coronaviruses usually circulate in the human population: HCoV-OC43, HCoV-HKU1, HCoV-NL63 and HCoV-229E.

In the past two decades, three coronaviruses with the potential to cause life-threatening disease in humans have emerged. Severe acute respiratory syndrome coronavirus (SARS-CoV) emerged in 2002 in China and spread, resulting in 8,100 infections and nearly 800 deaths in 37 countries. Ten years later, the Middle East respiratory syndrome coronavirus (MERS-CoV) emerged in the Arabian Peninsula and spread to 21 countries, causing outbreaks in humans and infecting around 2,500 individuals, with a fatality rate of 35%. In 2019, infections by a coronavirus now known as SARS-CoV-2 that can cause fever, severe respiratory illness, pneumonia, diarrhea, dyspnoea and multiple organ failure were identified in China. As of February 2022, more than 414 million cases have been confirmed, leading to at least 5.8 million deaths, according to the World Health Organization (WHO).

All coronaviruses enter the host cells using a trimeric spike (S) transmembrane glycoprotein. The S protein is a type I membrane class 1 fusion protein and is organized into two functional subunits, which remain non-covalently bound in the pre-fusion conformation of the protein. The amino-terminal S1 subunit is responsible for binding to the host cell receptors, whereas the carboxy-terminal S2 subunit is responsible for fusion of the viral and cellular membranes. The S1 subunit is further divided into an N-terminal domain (NTD) and a RBD.

SARS-CoV-2 is phylogenetically closely related to SARS-CoV, sharing approximately 79.6% genomic sequence identity, and similar to SARS-CoV uses the S1 RBD to bind to ACE2 receptors on host cell types such as pneumocytes and enterocytes. After host cell binding, a conformational change in the S2 subunit results in virus fusion and entry into the target cell.

The S glycoprotein has been the primary focus of efforts to develop mAbs to target SARS-CoV-2, as it was already known to be a target of potent neutralizing mAbs against SARS-CoV and MERS-CoV. More than 20 mAbs that target the S glycoprotein, originating either from the B cells of convalescent patients with COVID-19 or immunization of humanized mice, have been tested in clinical trials, and some have received an EUA from the FDA for the treatment of patients with mild to moderate COVID-19 or for pre-exposure prophylaxis. These include sotrovimab, the combination of casirivimab and imdevimab, bamlanivimab (used as a monotherapy or in combination with etesevimab), the combination of cilgavimab and tixagevimab, reglalanivimab and bretelovimab.

However, a major limitation of most mAbs evaluated so far against COVID-19 has been the rapid appearance of SARS-CoV-2 variants of concern (VOCs) that can escape both single mAbs and cocktails of mAbs (TABLE 2). VOCs such as Alpha, Beta, Gamma and Delta have 9–12 mutations in regions of the S glycoprotein, which typically have only a partial impact on the effectiveness of therapeutic mAbs. However, Omicron variants have accumulated more than 35 mutations in the S glycoprotein, of which 15 occur in the RBD, which is not only the site that binds to the host receptor ACE2 but also the key target of therapeutic mAbs, as well as neutralizing antibodies produced by the natural and vaccine-induced immune response. The emergence of Omicron VOCs (BA.1, BA.1.1 and BA.2) has rendered numerous mAbs with EUAs and/or in advanced clinical development partially or almost ineffective. These include the combination of casirivimab and imdevimab, the combination of bamlanivimab and etesevimab, and sotrovimab. These mAbs have all been developed from patients infected with SARS-CoV-2, with the exception of sotrovimab, which was isolated from an individual infected with SARS-CoV. Nevertheless, two recently developed mAbs, bbtelovimab and P2G3, retain full activity against the Omicron VOCs.

Additional, ultrapotent neutralizing antibodies binding the RBD have been identified, and some are...
Currently in clinical development (Table 2). These mAbs are categorized into five groups based on their clustering and binding to the RBD (Fig. 3c). The group 1 mAbs bind the receptor-binding motif (RBM) similarly to ACE2 on the left side of the ridge, binding L455, F456, F486, N487 and Y489; members of this group include REGN10933 (Ref. 134), S2E12 (Ref. 135), COV2–40 (Ref. 136), BD-236 (Ref. 136), C102 and C105 (Ref. 137). The group 2 mAbs also bind the RBM, but are positioned more upright and straddle the centre of the ridge, binding

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**Fig. 3 | The SARS-CoV-2 spike protein and neutralizing antibodies targeting the receptor-binding domain.**

- **a** | Structural representation of the spike (S) protein of severe acute respiratory syndrome coronavirus 2 (SARS-CoV-2) with modelled carboxy-terminal part.
- **b** | SARS-CoV-2 spike protein domain architecture.
- **c** | Structure of the receptor-binding domain (RBD) (grey) bound to angiotensin-converting enzyme 2 (ACE2) (forest green) [PDB:6M0J].
- **d** | Neutralizing antibodies can prevent opening of the S protein to block RBD erection, block the RBD to prevent binding to ACE2, block the amino-terminal domain (NTD) or block the fusion step.
- **e** | Footprints of ACE2 (green) or footprints of different ultrapotent antibody neutralizer classes (blue) shown on the RBD (grey). To generate footprints, the ‘Clashes/Contacts’ tool in UCSF Chimera was used to identify residues on the RBD that contact monoclonal antibodies (mAbs). Default contact criteria with van der Waals overlap of ≥–0.4 Å or more were used. CH, central helix; CD, connector domain; FP, fusion peptide; HR1, heptad repeat 1; HR2, heptad repeat 2; SD1, subdomain 1; SD2, subdomain 2; SS, signal sequence; TM, transmembrane region.
Infusion preparation, named RespiGam, was used prophylactically from the late 1990s to the early 2000s to prevent severe RSV-associated lower respiratory tract disease in young children with bronchopulmonary dysplasia or premature birth\textsuperscript{150,153}. The use of RespiGam was discontinued in 2003 and replaced by prophylaxis with palivizumab, the first neutralizing mAbs developed to treat severe RSV infection in high-risk infants\textsuperscript{148}, which was approved in 1998.

Of the three RSV surface proteins (F, G and SH), F-specific antibodies account for the majority of neutralizing activity in the sera of infected humans\textsuperscript{165,169}, and so the F glycoprotein has been the focus of mAbs development for RSV. F is a trimeric type I fusion glycoprotein responsible for merging the viral membrane with cellular membranes, and similar to many other viral fusion glycoproteins it undergoes major structural rearrangements during the transition from the pre-fusion to the post-fusion state\textsuperscript{166}. Palivizumab is a humanized mAb that binds to antigenic site II of the F glycoprotein. Importantly, site II as well as sites I, III and IV are present in both pre-fusion and post-fusion conformations of the F glycoprotein (Fig. 4a). Therefore, palivizumab does not prevent triggering of conformational changes in F, and presumably blocks entry and membrane fusion by preventing the pre-hairpin to hairpin or the hairpin to post-fusion conformational change\textsuperscript{162}. Motavizumab, an affinity-matured derivative of palivizumab that binds to the F glycoprotein with tenfold greater potency\textsuperscript{163}, was shown to generate a relative decrease of 26% in RSV hospitalization compared with palivizumab. However, results from phase III clinical trials in 2010 showed only a marginal improvement in comparison with palivizumab\textsuperscript{154} and an increase of adverse events\textsuperscript{164}, leading to termination of its development.

Novel mAbs under evaluation for RSV such as nirsevimab (MEDI8897) and suptavumab (REGN2222) target antigenic sites that are present only in the pre-fusion conformation of the F glycoprotein. Nirsevimab, which is derived from a mAb called D25 isolated by AIMM Therapeutics using a target-agnostic approach\textsuperscript{165}, targets the antigenic site Ø\textsuperscript{166} and is more potent than palivizumab in vitro\textsuperscript{167}. It has also been engineered for extended half-life through ‘YTE’ substitutions in the Fc region (see Antibody engineering section), which means that only one dose of nirsevimab may be required to cover a typical 5-month RSV season, rather than the five once-monthly doses that would be required for palivizumab. Indeed, nirsevimab showed an extended half-life offering protection from RSV for a typical 5-month season with a single intramuscular dose in a phase III trial\textsuperscript{166}. However, suptavumab, which targets the antigenic site V, failed to meet the clinical end points in a phase III study due to the appearance of resistant RSV B strains\textsuperscript{168,169}.

**Ebola virus.** Ebolaviruses are single-stranded RNA viruses that belong to the Filoviridae family. The genus *Ebolavirus* contains six species with different designations: *Bundibugyo ebolavirus* (*Bundibugyo virus*), *Reston ebolavirus* (*Reston virus*), *Sudan ebolavirus* (*Sudan virus*), *Taï Forest ebolavirus* (*Taï Forest virus*), *Bomali ebolavirus* (*Bomali virus*) and *Zaire ebolavirus* (*Ebola virus*).
Neutralizing antibodies that bind to glycoproteins from various pathogens.

**a** | Antigenic sites of the fusion protein (F) of respiratory syncytial virus (RSV). Locations of six antigenic sites on pre-fusion structure shown on left [PDB:4MMU], with locations on post-fusion structure shown on right [PDB:6APB]. (ReFS.)

**b** | Crystal structure of Zaire Ebola virus (EBOV) glycoprotein [PDB:5JO3]; GP2 in dark grey, GP1 head in blue, GP1 glycans in yellow. To the right, trimeric gp120 shown with the ZMapp antibody cocktail [PDB:5KEN and PDB:5KEL], with mAb114 [PDB:5FHC] or with three monoclonal antibodies (mAbs) that make up REGN-EB3. e | Glycoproteins of human cytomegalovirus (HCMV) and antibody-binding sites. Trimer gH/gL/gO structure [PDB:7LBE] shown with antibody footprint (orange) for MSL-109 and 13H11 mAb on gH (left) and gH/gL/UL128/UL130/UL131A pentamer [PDB:5VOD] (centre) shown with footprint of antigenic sites present on UL. On the right, structures of gB in pre-fusion [PDB:7KDP] shown with coloured antigenic sites (AD1 in orange, AD2 in red, AD4 in green and AD5 in blue).

**d** | Influenza haemagglutinin (HA) consists of three homotrimers built from two chains, HA1 and HA2, that are linked via a disulphide bond. HA molecules are commonly divided into immunodominant head and immunosubdominant stem. Based on sequence variations of HA, influenza is divided into two phylogenetic groups that are further subdivided into a total of 18 strains. Colour-coded dots represent observed binding of various broadly neutralizing antibodies (bNAbs) against different HAs [PDB:3512]. Footprints of several group 1-specific and group 2-specific antibodies as well as pan-group reactive antibodies on HA stem shown to right. CD, core domain.

**Fig. 4** | Crystal structure of Zaire Ebola virus (EBOV). EBOV causes severe disease with a high case fatality rate of 25–90%176.

The glycoprotein of EBOV is a trimeric class I fusion protein formed by three disulphide-linked GP1-GP2 heterodimers forming a chalice-shaped trimer on the viral surface177. The GP1 subunit binds to the EBOV receptor, NPC1, allowing GP2-mediated fusion of the viral and host cell membranes178. (Fig. 4b). GP1 bears the RBS, glycans and mucin-like domain. GP2 contains an N-terminal peptide, internal fusion loop, stalk and transmembrane domain. Of note, the GP1 subunit contains a core domain which is shielded by a ‘glycan cap’, made by the heavily glycosylated mucin-like domain. The mucin-like domain is dispensable for viral entry, but is a decoy target for host antibody responses179.

**Human cytomegalovirus**. Human cytomegalovirus (HCMV) is an enveloped double-stranded DNA virus with a genome size of more than 235 kb, making it the largest known genome of human herpesviruses. HCMV is a member of the betaherpesvirus family and establishes lifelong latency in healthy individuals.

Primary infection is generally asymptomatic; however, viral reactivation in immunocompromised hosts can be a life-threatening disease and vertical virus transmission during pregnancy is one of the leading causes of congenital birth defects180,181. HCMV utilizes different glycoprotein complexes to allow cellular entry into a large variety of cells. Indeed, two gH/gL-containing complexes regulate viral tropism182,183; more specifically, the gH/gL/gO trimer binds to PDGFRα and is primarily required for infection in fibroblasts184,185, whereas the gH/gL/UL128/UL130/UL131A pentamer binds to neuropilin 2 and is required for viral entry into epithelial, endothelial and myeloid cells186-188. (Fig. 4c). In addition, the gB homotrimer catalyzes membrane fusion between the virus and infected cells189-191.

Several mAbs targeting the gH/gL or gB complexes have been isolated and have shown modest efficacy in an in vitro model of infection. To identify the most potent HCMV neutralizing antibodies, the Lanzavecchia group isolated a large panel of mAbs from memory B cells in naturally infected donors and found that the pentameric complex represented the main target of neutralization against HCMV, eliciting neutralizing antibodies with a potency several orders of magnitude greater than any other HCMV complex192,193. This discovery shed light on the importance of the antigen selection to identify potent neutralizing mAbs and to design efficient vaccines.

The use of HCMV-specific mAbs for the prevention of HCMV infection and disease after allogeneic haemopoietic stem cell transplantation194 or solid organ transplant has been studied extensively195. Among the mAbs that have been evaluated in clinical trials, MSL-109, a human mAb targeting HCMV surface glycoprotein gH196, recombinant purified virus glycoprotein197. REGN3479 (now known as mafitivimab) recognizes the conserved GP2 fusion loop and provides neutralizing activity, whereas REGN3471 (now known as odesitivimab) recognizes the outer glycan cap and has cell-targeting functions. REGN3470 (now known as atoltivimab) binds to the GP1 head and offers both neutralization and cell-targeting activities, including FcγRIIIa and other FcγR-related functions.

During the EVD outbreak that occurred in the Democratic Republic of Congo in 2018, the triple mAb cocktail ZMapp, the monotherapy mAb114 (ansuvimab, also known as VRC 608; developed by Ridgeback Biotherapeutics) and the triple mAb combination REGN-EB3 (mafitivimab, odesitivimab and atoltivimab) were evaluated in an umbrella trial. After an interim analysis, mAb114 monotherapy and REGN-EB3 were both found to be superior to ZMapp with respect to the primary outcome and patient mortality179, and were approved by the FDA for the treatment of EVD in 2020 (ReFS190,191).
was tested as a supplementary treatment for patients with AIDS with HCMV-induced retinitis, but development was halted during phase II/III trials owing to lack of efficacy\(^\text{179}\). RG7667, a mixture of two mAbs binding the gHgL and pentamer complexes\(^\text{179}\), could potentially neutralize HCMV infections of all the cell types tested. However, when evaluated in a phase II trial for recipients of kidney transplants\(^\text{200}\), RG7667 did not meet the primary end point within 12 weeks post-transplant. Next, CSJ148, which consists of two anti-HCMV mAbs, an anti-gB mAb (LJP538) and an anti-pentamer mAb (LJP539)\(^\text{201}\), was evaluated in a phase II trial for prophylaxis of HCMV in patients undergoing haematopoietic stem cell transplantation\(^\text{202}\) and also did not achieve the primary end point. Nevertheless, in NHPs, the presence of durable and potently neutralizing antibodies at the time of primary infection was shown to prevent transmission of systemically replicating maternal rhesus CMV to the developing fetus\(^\text{203}\).

So, despite these significant development efforts, no anti-HCMV mAb has yet been FDA-approved. The only clinically available antibody-based therapy is still CytoGam, a preparation of CMV immunoglobulin used for intravenous injection; however, the neutralization capacity of this preparation is suboptimal\(^\text{204,205}\).

**Influenza.** Influenza, a member of the Orthomyxoviridae family, has four types — influenza A, B, C and D — all of which have a segmented, negative-sense, single-stranded RNA genome. Influenza A and B are responsible for severe infections in humans, whereas influenza C causes only mild symptoms and influenza D is not known to infect humans\(^\text{206}\). The viral genome consists of eight segments that encode at least 12 proteins: haemagglutinin (HA), neuraminidase (NA), PB2, PB1, PB1-F2, PA, PA-X, NP, M1, M2, NS1 and NS2\(^\text{207}\).

The two glycoproteins on the viral surface are HA and NA, with HA being more abundant (~500 molecules per virion) than NA, with a ratio of 4:1 to 5:1\(^\text{208}\). In total, 18 different HA subtypes (H1–H18) have been identified and can be separated into two phylogenetic groups according to their genetic sequences. Group 1 includes H1, H2, H5, H6, H8, H9, H11, H12, H13, H16, H17 and H18, whereas group 2 includes H3, H4, H7, H10, H14 and H15 (Fig. 4d). Eleven different NA subtypes (N1–N11) are known; group 1 includes N1, N4, N5 and N8; group 2 includes N2, N3, N6, N7 and N9; and group 3 includes N10 and N11\(^\text{209}\). Each subtype encompasses several viral strains that represent the seasonal changes of the virus.

The HA head is the major target of the antibody response, providing only limited breadth due to its high sequence diversity and changes in glycosylation sites\(^\text{207}\). Although the structural architecture of HA is conserved overall, the sequence as well as glycosylation patterns differ among different subtypes\(^\text{210}\). Due to its high sequence variability, the elicited antibody response against the immunodominant head is strain-specific and provides only short-lived immunity\(^\text{211}\). The head contains the RBS that is responsible for viral attachment to the host cells through binding to sialic acid receptors. The RBS forms a shallow pocket and consists of four segments: the 190 helix and the 130, 150 and 220 loops\(^\text{211}\). The RBS itself is relatively conserved except for the 220 loop, whereas the remaining head is highly diverse in sequence.

Antibodies generally target five major antigenic sites, Ca1, Ca2, Cb, Sa and Sb for H1, and sites A–E for H3, which are located around the RBS\(^\text{212}\). Antibodies to the RBS site are potent as they block viral attachment or prevent receptor-mediated endocytosis, and therefore neutralize the virus, rendering it unable to infect cells\(^\text{213}\). However, these antibodies are generally strain-specific and thus not capable of providing immunity against drifted strains. Some exceptions have been observed, such as the bNAb C05\(^\text{214}\), S139/1\(^\text{215}\) and F045-092\(^\text{215}\), which target the RBS and are able to neutralize within their group. Furthermore, recently identified bNAb targets hidden epitopes at the HA trimer interfaces near the HA head domain that become accessible during a ‘breathing motion’ of the subunits\(^\text{216}\). FluA-20 has shown broad reactivity to most subtypes by recognizing a conserved epitope at the trimer interface and functions by inhibiting cell to cell spread of the virus\(^\text{217}\).

On the other hand, the HA stem is highly conserved mainly within subtypes and, to some extent, across subtypes, evolving at a much slower rate than the head domain\(^\text{218}\) (Fig. 4d). Several bNAb against this region have recently been characterized in humans after infection or vaccination, making it a highly interesting vaccine design target\(^\text{219}\). These bNAb generally target a hydrophobic pocket around the Trp21 HA2 residue on the HA stem in close proximity to the fusion peptide and block membrane fusion by retaining the HA in its pre-fusion state\(^\text{217}\).

Other mechanisms of action for stem-directed antibodies involve inhibition of proteolytic cleavage, reactivation of viral egress by blocking NA activity through steric hindrance, ADCC and antibody-dependent cellular phagocytosis\(^\text{220}\). These antibodies tend to be less prevalent in humans and often demonstrate little or no neutralizing activity, but are protective in challenge studies\(^\text{217}\). Many of these antibodies are group-specific due to differing glycosylation patterns between group 1 and group 2 influenza HA at position N38, which is only present in the group 2 HA that is the major antigenic site on the HA stem and, thus, can interfere with antibody binding. The F10\(^\text{219}\) and CR6261\(^\text{219}\) mAbs are specific for group 1 strains, whereas CR8020\(^\text{219}\) and CR8043\(^\text{219}\) mAbs are group 2-specific. CR8020 and CR8043 are encoded by VH1-18 and VH1-3 germline regions, whereas group 1 stem-specific neutralizing antibodies have been shown to generally derive from the VH1-69 germline gene\(^\text{220}\). Recognition of both group 1 and group 2 has been shown to commonly involve VH1-18-derived, VH6-1-derived, VH3-23-derived or VH3-30-derived antibodies. Some of the broadest neutralizing antibodies, such as FI6v3\(^\text{219}\), CR9114\(^\text{219}\), 39.29\(^\text{219}\) and MEDI8852\(^\text{219}\), are able to engage HAs from both group 1 and group 2. The exact mechanisms driving differences in immunogenicity of the head versus the stem are not fully understood. However, several explanations have been proposed, including the restricted spatial availability of the stem domain to
B cell receptors because of its close proximity to the viral membrane. MEDI8852 is a human IgG1 mAb isolated from a patient with uncomplicated influenza A infection. MEDI8852 was evaluated in phase IIa trial and is still in development by MedImmune. Vir Biotechnology is developing VIR-2482, an influenza A neutralizing mAb that binds to the conserved region of HA and neutralizes all major strains since the Spanish flu in 1918 (H1N1). It is therefore tempting to speculate that this type of antibody could be used as a universal prophylactic agent, overcoming the limitations of current influenza vaccines, for which the antibody response is dependent on individual seasonal antigens. In addition, because the serum half-life of VIR-2482 has been increased by Fc domain engineering, a single dose can last the entire influenza season of around 5–6 months.

Implications for vaccine development

Antigen–antibody interactions and structural vaccinology for vaccine design. Vaccines have proven to be the most effective prophylactic strategy for infectious diseases. However, traditional vaccine development approaches, which rely on three categories of vaccines (live-attenuated, inactivated and dissociated pathogens), have failed for viruses such as HIV, RSV, influenza, hepatitis C virus, HCMV or EBOV.

For most vaccines, the antibody response is crucial and, thus, the identification of antibodies that can potently neutralize a pathogen is a key factor for accelerating vaccine development. Reverse vaccinology 2.0, also known as antibody-based vaccinology, aims to overcome the limitations of traditional approaches by engineering novel vaccines based on the structural characterization of antigens in complex with their cognate antibodies, with the antigen-specific antibody response acting as a correlate of protection (FIG. 5).

Antigens are generally identified by proteomic analysis of crude homogenates of infected cells and, then, chosen based on immunogenicity and their ability to stimulate an immune response. However, this approach is time-consuming, and structural integrity or immunogenicity is not guaranteed upon expression of a newly identified protein. Experimentally driven isolation of neutralizing mAbs and identification of their target is still the best approach to identify vaccine candidates. One of the main advantages of antibody-driven vaccinology is isolation of potent neutralizing mAbs. These mAbs will be useful for passive immunization of immunocompromised patients, and/or as therapeutic agents during the acute phase of infection. An additional advantage of the antibody-driven approach is that mAbs can be instrumental in identifying the optimal vaccine antigen for which mAb binding can block virus transmission.

This identification step can be done by immunoprecipitation coupled to mass spectrometry. In addition, once identified, both the antigen and the mAb can be used for structural vaccinology. The latter approach aims at elucidating the atomic structures of the viral antigens with a neutralizing Fab (FIG. 5). Structural vaccinology is a valuable source of information to engineer antibodies for stabilization purposes. The combination of antibody-driven and structure-based antigen design strategies is particularly efficient in developing rapid responses to emerging infectious disease threats.

Facilitated by the improvements in high-throughput B cell technologies, the structural insights into human mAbs have been instrumental in directing the immune response to conserved antigenic sites. Complementary to this approach, structure-based design has been employed to completely remove domains containing sites targeted by non-neutralizing antibodies or to identify possible positions for the introduction of glycosylation sites to mask such epitopes.

Respiratory syncytial virus. RSV provides a case study for the reverse vaccinology 2.0 concept. For RSV, neutralizing antibodies mainly target the head domain of the fusion (F) protein. To focus the immune response against antigenic sites in the F head, Boyington et al. designed truncated F immunogens containing only the head region. These immunogens were successful in eliciting neutralization titres comparable with full-length pre-fusion F protein. These proteins are especially vulnerable in their pre-fusion state, and thus have been the target of structure-guided stabilization efforts to lock antigens in their pre-fusion conformation in order to promote a neutralizing antibody response.
that prevents membrane fusion\textsuperscript{(180).} This structural information can be used to identify stabilizing mutations such as disulfide bonds or proline mutations to rigidify the protein backbone and mutations to fill cavities to impede transition from pre-fusion to post-fusion states. Additionally, structure-based design has been applied to fuse domains for multimerization or to remove potential unstable regions\textsuperscript{(142).}

A prime example for the successful stabilization of a pre-fusion antigen is the F protein, where the elicitation of a potent neutralizing antibody response relies on targeting the pre-fusion state\textsuperscript{(167).} Multiple structure-based design strategies were applied to stabilize the F protein in its pre-fusion conformation. McLellan et al. designed intra-protomer disulfide bonds coupled with additional mutations to fill cavities in the native antigen to achieve a stabilized pre-fusion antigen (Ds-Cav1)\textsuperscript{(166).} A different design approach by Krarup et al. focused on introducing proline residues to prevent structural rearrangements of the F protein from occurring during the adoption of the post-fusion conformation\textsuperscript{(245).} Immunization studies in animals confirmed the improved neutralization potential of pre-fusion F constructs over post-fusion antigens, further supported by a subsequent study by Joyce et al. which highlighted that stability improvements of the DS-Cav1 immunogen in the pre-fusion state increased the neutralizing antibody response fourfold as compared with Ds-Cav1 (REF\textsuperscript{(246)})

A comparatively young field in structure-based vaccine design is epitope scaffolding, which relies on the transplantation of viral epitopes onto unrelated carrier proteins, so-called scaffolds, to focus the immune response against conserved, functional sites that are known to be targeted by neutralizing antibodies\textsuperscript{(246).} This strategy has first been applied to the design of novel immunogens for HIV\textsuperscript{(246–251),} although one limitation in antigen transplant is the stabilization of distant discontinuous epitopes\textsuperscript{(252).} For RSV, computational design of immunogens was first achieved by McLellan et al., who designed immunogens for antigenic site II of the F protein by grafting of the epitope of motavizumab onto an unrelated protein scaffold\textsuperscript{(253).} However, immunization of mice with this epitope scaffold did not elicit neutralizing antibodies, although it did elicit sera with F-binding activity\textsuperscript{(254).} Subsequent design efforts focusing on the same antigenic site resulted in a novel epitope scaffold that engaged site-specific antibodies with high affinity and boosted subdominant antibodies with enhanced neutralization\textsuperscript{(254,255).} A recent study by Sesterhenn et al. demonstrated that a cocktail strategy of three computationally designed immunogens displaying RSV F sites I, II and IV elicited a neutralizing antibody response and allowed for the focusing of the antibody response against specific antigenic sites upon immunization of mice and NHPs\textsuperscript{(256).} Using a strategy termed motif-centric design, the authors computationally designed de novo topologies around the extracted F antigenic sites to improve epitope stabilization and accurate display of the antigenic site\textsuperscript{(257).}

Although epitope scaffolding approaches have facilitated the design of novel immunogens that display neutralization epitopes, these designed immunogens have mostly been restricted to small, continuous epitopes\textsuperscript{(246,248,254)} with few exceptions\textsuperscript{(256,257).} In contrast, observed epitopes often consist of multiple segments\textsuperscript{(258,259)} that are challenging for structure-based design. As an example, Marcandalli et al. demonstrated that the use of structure-based design of a self-assembling protein nanoparticle presenting a pre-fusion-stabilized DS-Cav1 in a repetitive array on the nanoparticle exterior induced neutralizing antibody responses up to tenfold higher than trimeric DS-Cav1 alone\textsuperscript{(260).} The same nanoparticle was used to derive promising vaccines against SARS-CoV-2 (REF\textsuperscript{(261)}, influenza\textsuperscript{(262)} and HCMV\textsuperscript{(263)}.)

\textit{Ebola virus.} Vaccines from several platforms have shown protection against EVD in human and NHPs, and two that are not based on the reverse vaccinology concept have been licensed: Ervebo (rVSV-ZEBOV) and a two-dose combination of Zabdeno/Mvabea\textsuperscript{(271–279).} Both vaccines are based on viral vectors, and recent analysis of human B cell responses from vaccinated humans demonstrated low levels of somatic hypermutations in mAbs induced by the vaccine, even if they were protective\textsuperscript{(270).} Rational design efforts for protein-based subunit vaccines are ongoing\textsuperscript{(271).}

Ervebo is a live, attenuated recombinant vesicular stomatitis virus-based vector expressing the envelope glycoprotein gene of Zaire EBOV instead of the VSV-G gene (FIG. 4b). NHP studies demonstrated rVSV-ZEBOV efficacy in stringent conditions, where a single inoculation at \(1 \times 10^7 \text{pfu} \) was shown to protect against illness, viraemia and death after challenge with a high dose of EBOV (1,000 pfu, generally thought to represent 100–1,000 times the lethal dose (median lethal dose) in experimental animal studies)\textsuperscript{(266–270).} Zabdeno/Mvabea each contain a monovalent replication-incompetent virus, Zabdeno, an adenoviral vector of serotype 26 that encodes the EBOV-GP Mayinga variant (Ad26.ZEBOV), is used for the priming injection, and Mvabea, a modified vaccinia virus Ankara-Bavarian Nordic Filo-vector encoding the same glycoprotein (MVA-BN-Filo)\textsuperscript{(271)}, is used for the booster.

Other vaccines in development include cAd3-EBO Z, an attenuated version of a chimpanzee adenovirus (cAd3) encoding the EBOV-GP glycoprotein\textsuperscript{(272).} Finally, a recombinant nanoparticle vaccine with the EBOV-GP Makona strain was found to induce a potent immune response\textsuperscript{(273).}

\textit{Human cytomegalovirus.} The development of HCMV vaccines began in the early 1970s, and two attenuated virus strains were isolated for laboratory work: AD169 and Towne\textsuperscript{(274,275).} The AD169 attenuated strain was quickly abandoned whereas the Towne attenuated strain progressed to extensive testing in recipients of solid organ transplants and healthy volunteers\textsuperscript{(276).} Recipients of kidney transplants were shown to be highly protected against serious CMV disease and graft rejection. However, protection against viral infection was not statistically significant. The next development in the quest for CMV vaccines was the identification of a surface protein of CMV called glycoprotein B or gB (FIG. 5c). When combined with the MF59 oil-in-water adjuvant,
The vaccine was safe and partially effective\textsuperscript{272}, but the levels of neutralizing antibodies were weak in humans after three injections\textsuperscript{277-279}. The subunit gB protein was also combined with the AS01 adjuvante to stimulate Toll-like receptor 4 (TLR4), which elicited higher and more prolonged levels of anti-gB antibodies in humans. Unfortunately, the adjuvanted vaccine was never tested for efficacy.

The gB antigen is a class III trimeric fusion protein and is used in the post-fusion conformation\textsuperscript{198}. Therefore, it was suggested that gB in the pre-fusion conformation could generate a higher neutralizing response, but recent results have, surprisingly, indicated that this might not be the case\textsuperscript{290}. The pentameric complex of proteins present on the surface of CMV consists of glycoprotein H (gH), glycoprotein L (gL) and the products of genes UL128, 130 and 131A. The complex has been shown to generate far higher titres of neutralizing antibodies than gB\textsuperscript{198,281} (Fig. 4c). This discovery has since driven much of the HCMV vaccine field and vaccine trials are ongoing. However, if neutralizing antibodies are necessary to prevent infection and spread of HCMV, a strong T cell response is also needed to suppress reactivation of the virus in patients who are seropositive. Forthcoming clinical trials using either a recombinant protein subunit (NCT05089630) or mRNA (NCT05105048) will indicate whether the pentamer can generate an expanded influenza-specific humoral immune response when compared with trivalent inactivated influenza vaccine\textsuperscript{294}. Together, these results demonstrate that immunogens lacking the immunodominant head domain can elicit a group-specific, hetero-subtypic immune response\textsuperscript{293,294}. Recently, a ferritin nanoparticle-based vaccine incorporating the ectodomain of HA from an H2N2 pandemic strain was demonstrated to be safe and immunogenic in a phase I clinical trial, supporting its potential application in pandemic preparedness and universal influenza vaccine development\textsuperscript{295,296}.

A different approach to focus the immune response against conserved antigenic sites that is structurally less demanding relies on the masking of non-neutralizing antigenic sites in the HA head through hyper-glycosylation and removal of glycans from the HA stem\textsuperscript{297,298}. Eggink et al. applied this strategy to hyper-glycosylated immunodominant epitopes in the HA head, resulting in the silencing of immunodominant sites. Immunization of mice confirmed a shift of the antibody response towards the immunorecessive stem domain and improved protection of mice after viral challenge\textsuperscript{299}. Beyond focusing the immune response against the HA stem, this strategy can be also used to improve the immune response against subdominant epitopes in the HA head\textsuperscript{117}.

**Influenza.** One of the major obstacles for effective influenza vaccines is the immunodominance of the highly diverse HA head that is responsible for viral attachment. Structure-based design methods have been focused on the conserved but immunorecessive HA stem region (Fig. 4d).

To shift the antibody response towards the stem domain, chimeric HA molecules have been engineered, consisting of a common HA stem paired with different HA heads\textsuperscript{291-293}. Repeated immunizations of mice demonstrated that a stem-specific antibody response can be mounted, resulting in heterologous and hetero-subtypic immunity\textsuperscript{284}. In ferrets, these constructs were shown to reduce viral loads after influenza virus challenge\textsuperscript{296,297}. Chimeric HA has been tested in a phase I clinical trial and was found to be safe, and able to induce a broad, strong, durable and functional immune response\textsuperscript{298}.

Although Kanekiyo et al. demonstrated that the isolation of RBDs from different HA strains and multimerization on ferritin nanoparticles resulted in the elicitation of a B cell response against conserved epitopes\textsuperscript{199}, the most promising design strategies are based on isolated HA stem-only antigens created by removing the HA head\textsuperscript{290,291}. Two such design strategies are headless HAs from Yassine et al.\textsuperscript{220} and the mini-HAs from Impagliazzo et al.\textsuperscript{215}.

The headless HA immunogens were developed by removing the HA head from an H1 strain, followed by multiple rounds of structure-based design that yielded stabilized stem immunogens\textsuperscript{272}. Lethal influenza challenge with a hetero-subtypic H5N1 strain showed complete protection in mice and partial protection in ferrets; however, no cross-group reactivity with group 2 viruses was observed\textsuperscript{292}. Applying the same structure-based design strategy to H3 and H7 HA resulted in two group 2 headless immunogens that were able to elicit protective, homo-subtypic antibodies in mice\textsuperscript{292}. Immunization of NHPs with headless H3 generated neutralizing antibodies against divergent H3N2 strains and selected H1N8 as well as H7N9 strains\textsuperscript{293}.

The mini-HA molecules developed by Impagliazzo et al.\textsuperscript{220} follow a similar design approach. A combination of rational and library-based design approaches was applied to generate stabilized HA stem molecules that lack the immunodominant head and the transmembrane region\textsuperscript{293}. The designs were based on an H1 subtype and were shown to be protective in lethal heterologous and hetero-subtypic challenge in mouse models\textsuperscript{231}. In pre-exposed NHPs, the mini-HAs elicited an expanded influenza-specific humoral immune response when compared with trivalent inactivated influenza vaccine\textsuperscript{294}. Together, these results demonstrate that immunogens lacking the immunodominant head domain can elicit a group-specific, hetero-subtypic immune response\textsuperscript{293,294}. Recently, a ferritin nanoparticle-based vaccine incorporating the ectodomain of HA from an H2N2 pandemic strain was demonstrated to be safe and immunogenic in a phase I clinical trial, supporting its potential application in pandemic preparedness and universal influenza vaccine development\textsuperscript{295,296}.

**COVID-19.** Acquisition of structural data for the SARS-CoV-2 S protein played an important role in the development of multiple COVID-19 vaccines, including the extraordinarily successful mRNA vaccines. Another critical aspect in the development of effective vaccines was the intensive experience gained from studies of the S protein of other coronaviruses such as HCoV-HKU1 [REF. 300] and MERS-CoV\textsuperscript{301,302} led by the McLellan and Veesler laboratories. This allowed the design of stabilizing mutations in the versions of the S protein encoded in the mRNA vaccines (BNT162b2 mRNA from Pfizer-BioNTech and mRNA-1273 from Moderna) and some vectored vaccines (Ad26.COV2.S from Janssen)\textsuperscript{303}, such as deletion of the polybasic cleavage site, inclusion of stabilizing mutations\textsuperscript{304} and inclusion of trimerization domains\textsuperscript{305,306}.
HIV. Research on the development of an HIV vaccine has strongly benefited from the isolation of human bNAbS, reviewed in depth elsewhere97–100. The use of bNAb-instructed stabilization of HIV Env trimers is the basis for many promising immunogens111. In addition, the most promising strategies use stabilized trimers with the aims of activating particular bNAb-producing cell precursors and guiding their affinity maturation to generate mature bNAbS — a strategy called germline targeting112. Recently, 97% seroconversion was reported113 in healthy subjects vaccinated with the eOD-GT8 immunogen114,115, demonstrating the strong potential of the germline-targeting strategy. Molecular analysis of the B cells induced during the clinical trial (NCT03547245) will provide a road map to accelerate progress towards an HIV vaccine. A phase I clinical trial in which an mRNA vaccine encoding the eOD-GT8 immunogen is used for the priming injection has recently been initiated (NCT05001373).

Closing perspectives

The emergence of SARS-CoV-2 and the devastating COVID-19 pandemic have emphasized the severity of the threat of emerging infectious diseases, especially those of zoonotic origin. In response, there has been unprecedented success in the discovery and development, manufacturing and regulatory evaluation of several anti-COVID-19 vaccines and mAb therapeutics in a very short time, achieved through exceptional mobilization of public and private resources. These advances have been based on the enormous scientific progress made in immunology and vaccinology over the last few decades, and also the crucial previous knowledge on the biology of coronaviruses, such as the S protein as a target for neutralizing antibodies.

Efforts to harness antibodies to combat COVID-19 have also benefited from technological advances and expertise gained particularly in the area of human bNAbS from the HIV field. It is expected that the generation of anti-infective mAbs that are urgently needed will benefit from the recent clinical successes and case studies highlighted in this article and elsewhere116. However, mAbs still have limitations. The potential for mutations in the viral targets of antibodies to allow viruses to escape neutralization has recently been highlighted with SARS-CoV-2 Omicron VOCs, and also by the development of resistant variants when selective pressure is applied in the setting of drug treatment, which has been observed in immunocompromised patients treated with bamlanivimab and etesevimab117, sotrovimab118 or REGN-COV2 (REF119).

With the recent advances in machine-learning algorithms, and the dramatic increases in the repertoire of available antibodies, further research should provide insight into the structural properties required for bNAbS92. Moreover, the progress made with gene-editing technology opens the possibility of engineering functionalities into human cells. Indeed, B cells engineered to carry broadly neutralizing B cell receptors are likely to represent a milestone to address pathogens with no vaccines or inefficient vaccines111. Vaccination by edited B cells able to differentiate into memory B cells, plasmablasts and long-lived plasma cells could be a valuable option for HIV and influenza prevention. Finally, given that a key current limitation of mAbs is still their limited distribution in tissues, novel formats or ways to specifically direct mAbs to the targeted tissues may render mAbs even more powerful tools in fighting viral infections.

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