The breakdown of the glomerular filtration barrier (GFB) manifests as a protein leak from the plasma into the urine (proteinuria), is associated with loss of normal kidney function and is a powerful risk factor for disease progression to organ failure. Clinically, patients may present with nephrotic syndrome (a high-grade form of proteinuria with oedema), isolated proteinuria or proteinuria with hypertension. Abnormal kidney function may also be detected upon blood testing. Childhood nephrotic syndrome is managed by clinical assessment and is determined to be sensitive to steroid therapy or steroid-resistant nephrotic syndrome (SRNS). In adults, GFB disease is frequently named according to the pathological lesions detected histologically following a kidney biopsy. The classical adult pathological descriptors of GFB disease are minimal change disease and focal segmental glomerulosclerosis (FSGS). However, common and rare diseases that present with protein leak into the urine, including hypertensive nephroangiosclerosis, diabetic kidney disease, lupus nephritis, pre-eclampsia and Alport syndrome, frequently have GFB disease as a major component upon histological inspection (Table 1). GFB breakdown results in glomerular cell activation, local scar tissue formation, occlusion of glomerular capillaries, impaired blood flow to the nephron and, ultimately, glomerular obsolescence. In addition, plasma proteins that leak into the glomerular filtrate directly injure the tubular epithelium, resulting in loss of tubular function and interstitial scarring. Therefore, the burden of GFB disease on human health is enormous. Occasionally, patients have reduced kidney function without damage to the GFB. In these infrequent cases, patients do not have proteinuria, and inspection of the kidney indicates that damage to the tubules and their supportive capillaries underlies the kidney disease.

Existing therapies for kidney disease have largely treated complications, symptoms or late manifestations of disease, and have had limited disease-modifying effects. Encouragingly, in recent trials, sodium–glucose co-transporter 2 (SGLT2) inhibitors have improved hyperglycaemia control as well as cardiovascular and renal outcomes in patients with advanced type 2 diabetes mellitus. However, despite their recognized pleiotropic effects on glomerular cells, SGLT2 inhibitors act predominantly on the kidney tubule and have little direct role in GFB function, and thus have only minor effects on proteinuria.

Recent large epidemiological studies have demonstrated an important predictive link between proteinuria and disease progression to organ failure in both...
common and rare diseases\textsuperscript{12}. These observations have prompted regulatory organizations to recommend proteinuria as an approvable primary end point in settings where protein levels in the urine are high. This simpler primary end point will likely enable the evaluation of investigational therapeutics in patients and accelerate the drug discovery process. These welcome changes coincide with new insights into the mechanisms by which kidney spil protein into the urine, following 20 years of research on the GFB. In this Review, we provide an overview of the GFB, introduce novel candidate therapies that specifically target the key aspects of GFB function and discuss challenges and potential opportunities for future research and development (Table 2).

### Overview of the GFB

The GFB (Fig. 1) was first described more than 50 years ago, and is a unique structure comprising highly fenestrated and specialized endothelial cells, an unusual basement membrane and a high density of arborized interdigitating cells known as podocytes\textsuperscript{11}. The GFB is the terminal unit through which the kidney filters water, salts and low molecular weight organic molecules from the blood. Podocytes form the architectural backbone and provide the mechanical stability of the GFB\textsuperscript{12,13}. Because of the mechanically demanding environment, cytoskeletal integrity of podocytes is paramount for their function\textsuperscript{14}. The specialized projections that interdigitate to form the slit diaphragm are key elements in the GFB and contribute to its overall physical stability\textsuperscript{14}. The central molecular components of the slit diaphragm were first identified in 1998 and 2000, when the genes that cause childhood nephrotic syndrome, \textit{NPHS1} (which encodes nephrin) and \textit{NPHS2} (which encodes podocin), were discovered and found to be exclusively expressed at the slit diaphragm\textsuperscript{14,17}. These proteins anchor the cytoskeleton to the plasma membrane. Nephrin (also known as \textit{NEPHS1}), along with similar proteins such as \textit{FAT1} and \textit{NEPH1/2}, form a multilayered bipartite scaffold between each of the interdigitating podocyte projections (tertiary foot processes)\textsuperscript{16}. The proteins form flexible spring-like protein bridges that prevent macromolecules from passing through. Since the discovery of these key proteins, an array of molecular components that maintain the podocyte tertiary foot processes through dynamic regulation of the actin cytoskeleton (such as those encoded by \textit{ACTN4}, \textit{INF2}, \textit{ARHGAP24}, \textit{MYH9} and \textit{TRPC6}) have been associated with the development of FSGS and nephrotic syndrome\textsuperscript{19–23}. Additional proteins that maintain slit diaphragm proteins (for example, \textit{CD2}-associated protein (\textit{CD2AP})) as well as proteins that tether the podocyte to the basement membrane (such as \textit{a3B2} integrin) all play vital roles in GFB maintenance.

One of the earliest structural features of GFB dysfunction is the retraction of podocyte tertiary foot processes, which reduces the podocyte surface area and disrupts the slit diaphragm\textsuperscript{14}. Persistent podocyte dysfunction leads to deposition of abnormal basement membrane matrix, which makes the tissue sclerotic and, ultimately, results in podocyte depletion\textsuperscript{17}. Recent ultrastructural analyses indicate that alterations in the number and the morphology of foot processes and/or shortening of the slit diaphragm in experimental FSGS leads to capillary dilation due to reduced compressive forces that counteract filtration pressure, which contributes to GFB breakdown and, ultimately, results in protein leak\textsuperscript{26}.

Podocytes regulate glomerular endothelial cell (GEC) growth, survival, differentiation and permeability by releasing the essential paracrine cytokine, vascular endothelial growth factor A (VEGFA), which binds to its cognate receptor, VEGFR2, on GECs\textsuperscript{27,28}. GECs are highly specialized cells with fenestrae and a charged luminal endothelial surface layer also known as the ‘glycocalyx’, which is a negatively charged network of proteoglycans, glycoproteins and glycolipids\textsuperscript{29–31} that helps maintain a charge-selective barrier\textsuperscript{32–34}. GEC dysfunction can also initiate and contribute to GFB breakdown\textsuperscript{12,13,34}.  

### Table 1 | Kidney diseases in which the GFB is implicated in pathogenesis

| Disease                     | Clinical manifestations | Potential molecular targets                                                                 |
|-----------------------------|-------------------------|----------------------------------------------------------------------------------------------|
| Minimal change disease      | Nephrotic syndrome      | APOL1, TRPC6, TRPC5, suPAR, ET\textsubscript{A}, ET\textsubscript{R}, CoQ\textsubscript{10} biosynthesis |
| Focal segmental glomerulosclerosis | Nephrotic syndrome, proteinuria, hypertension, CKD | APOL1, TRPC6, TRPC5, suPAR, ET\textsubscript{R}, CoQ\textsubscript{10} biosynthesis |
| Alport syndrome             | Nephrotic syndrome, proteinuria, haematuria, hypertension, CKD | miR-21, NRF2 |
| Diabetic kidney disease (GFB partially implicated) | Nephrotic syndrome, proteinuria, hypertension, CKD | TRPC5, TRPC6, ET\textsubscript{R}, miR-21 |
| Hypertensive nephroangiosclerosis (GFB partially implicated) | Proteinuria, hypertension, CKD | APOL1, TRPC6, ET\textsubscript{R} |
| Pre-eclampsia               | Nephrotic syndrome, hypertension | sFLT1 |
| Lupus nephritis             | Nephrotic syndrome, proteinuria, hypertension, CKD | APOL1, ET\textsubscript{R} |
| Other immune complex disease | Nephrotic syndrome, proteinuria, hypertension, CKD | ET\textsubscript{R}, complement |

APOL1, apolipoprotein L1; CKD, chronic kidney disease; CoQ\textsubscript{10}, coenzyme Q10; GFB, glomerular filtration barrier; ET\textsubscript{R}, endothelin type A receptor; miR-21, microRNA-21; NRF2, nuclear factor erythroid 2-related factor 2; sFLT1, soluble fms-like tyrosine kinase 1 (also known as soluble vascular endothelial growth factor receptor 1); suPAR, soluble urokinase plasminogen activator surface receptor; TRPC, transient receptor potential channel.
Some human kidney diseases resulting in GFB breakdown, such as pre-eclampsia and haemolytic uremic syndrome, are caused by factors directly targeting the glomerular endothelium. Human genetic association studies, combined with models of podocyte injury in gene-targeted animals, suggest that endothelial-specific nitric oxide synthetase (eNOS) plays a role in disease susceptibility and severity. Furthermore, abnormalities of the components of the glomerular basement membrane (GBM), such as mutations and variants in collagen type IV basement membrane proteins that are found in common and rare diseases, revealed the essential role of GBM constituents for the integrity of the GBF. The emerging picture is that podocytes, GECs and the GBM all contribute to the overall structure and function of this complex barrier.

Genetic studies of patients with familial and sporadic forms of rare kidney diseases have provided variants and mutations in many disease-causing proteins that are highly expressed by podocytes, and, less frequently, highly expressed by GECs. Emerging whole-exome sequencing and genome-wide studies indicate that these same genes are abnormally expressed or harbour mutations in a significant proportion of patients presenting with common forms of kidney disease. Nevertheless, GFB breakdown for many is caused predominantly by environmental factors including xenobiotics, infections, local immune protein activation or immune cell activation, local metabolic disturbance from a diabetic environment or elevated glomerular pressure.

As our understanding of the GFB has evolved during the past 20 years, cell biology, comparative medicine and genetic experiments from humans have pointed to podocytes as critical to maintaining GFB function, and shown that podocyte injury or death causes scarring lesions in the glomerulus. A number of genes that are mutated or contain variants that confer increased disease risk are expressed by podocytes. Several of these have emerged as potential therapeutic targets. Circulating proteins that contribute to GFB breakdown following kidney transplantation may also be targeted for therapeutic benefit. Such proteins and factors can stimulate an injury response in podocytes, although the mechanisms by which they cause injury are not uniformly understood. Circulating complement proteins, immune complexes or autoantibodies are known to cause glomerular cell injury, in part through receptor-mediated interactions. Other circulating proteins and factors may similarly cause cell injury through receptor–ligand interactions.

The majority of pathological mutations in genes identified to cause isolated FSGS or SRNS are reported to regulate the assembly, turnover and degradation of the actin cytoskeleton or regulate its attachment at the cell surface, and other mutations cluster in the proteins...
that regulate the biosynthesis of mitochondrial constituents (Table 2). The following sections cover candidate therapies in development that directly target the GFB and have advanced to clinical trials or are at the preclinical stage of development. Many of these candidate therapies impinge directly on functions of mitochondria or the actin–myosin contractile machinery, which are central to GFB function.

Fig. 1 | Introduction to the GFB. Sequentially magnified schematic showing the nephron, the functional unit of the kidney, followed by the glomerulus, the primary interface between the microvascular circulation and the nephron. Each glomerular capillary loop forms a glomerular filtration barrier (GFB) comprising the glycocalyx and fenestrated glomerular endothelial cells (GECs), which block negatively charged molecules and plasma proteins; the glomerular basement membrane (GBM), which provides a physical scaffold that supports the GECs and podocytes, but contributes little directly to GFB function; and podocytes, which form primary, secondary and tertiary (foot) processes. Foot processes attach to the GBM by matrix tethering receptors including dystroglycans and integrins that, in turn, regulate the actin–myosin contractile apparatus to maintain foot process projections. Primary and secondary processes have an elaborate actin–myosin contractile apparatus and cross-link proteins running inside foot processes. The slit diaphragm, an ultrastructural molecular barrier, connects interdigitating foot processes, and contains NEPH1/2, P-cadherin, protocadherin FAT1, nephrin and roundabout homologue 2 (ROBO2). This molecular bridge forms part of the GFB and connects with the actin–myosin contractile apparatus via CD2-associated protein (CD2AP) and podocin. External short transient receptor potential channels (TRPCs) open in response to diverse stimuli, including mechanical forces, to release Ca2+ and regulate cellular responses. The function and localization of mitochondria in both podocytes and GECs, particularly in tertiary processes, is important in the maintenance of the GFB. Adapted from REF.70.
**Targeting mitochondrial function**

Mitochondria are critical to cellular metabolism, homeostasis and stress responses, and their dysfunction is linked to diseases of the GFB. Mitochondria have roles in the pathological generation of excessive reactive oxygen species (ROS), the regulation of intracellular calcium levels, cell death pathways and cytoskeletal dynamics. In this section, we discuss genetic variants in mitochondrial proteins that disrupt mitochondrial electron transport or biogenesis and contribute to GFB disruption. Although the precise contribution of mitochondria to oxidative phosphorylation in podocytes and GECs is not fully elucidated, mitochondria produce ROS, control cell survival by maintaining the mitochondrial membrane potential and regulate additional cell processes including metabolism, protein translation, iron homeostasis and DNA repair.

Abnormalities of podocyte function have been linked to mitochondrial disorders. Podocytes use large amounts of ATP to maintain the high surface area of tertiary foot processes via the actin cytoskeleton, and mutations in mitochondrial DNA and proteins could cause breakdown of the GFB because this ATP is lost; experimental evidence supports this hypothesis. In humans, xenobiotics, including tenofovir, that inhibit mitochondrial DNA replication can cause nephrotoxicity with glomerular scarring and proteinuria, underscoring the important role for mitochondrial homeostasis in maintaining the GFB.

Recent experimental data support additional roles for mitochondria beyond ATP production, including ROS production, cell death, integration of signalling pathways and responses to cell stress. High-content imaging studies demonstrate a relationship between mitochondrial dysfunction and changes in mitochondrial motility, fission and fusion events, localization and shape. Mitochondrial dysfunction in GECs, rather than podocytes, could also contribute to GFB loss. Mitochondrial oxidative stress lesions in GECs have been reported in patients and in experimental models of FSGS, as well as in segmental glomerular scars in patients with diabetic kidney disease. Mitochondrial stress in GECs is associated with loss of fenestrations and the glycocalyx.

Although much remains to be learned about the precise roles of mitochondria in GFB health and disease, mitochondrial function could be boosted as a broad-based approach to treat GFB disease. Several agents that may promote mitochondrial health are currently under clinical evaluation. These include a small-molecule activator of the transcription factor nuclear factor erythroid 2-related factor 2 (NRF2) from Reata Pharmaceuticals, which improves kidney function in a number of glomerular diseases, although it did not reduce protein leak at the GFB (ClinicalTrials.gov identifier NCT03019185). NRF2 activates both antioxidative and mitochondrial biogenesis programmes in cells. Lademirsen, developed by Sanofi-Genzyme, is an antisense oligonucleotide (ASO) that inhibits microRNA-21 (miR-21), has been reported to enhance mitochondrial function in podocytes and is under evaluation in patients with Alport syndrome (NCT02855268). ZyVersa is planning to evaluate VAR-200 (2-hydroxypropyl-β-cyclodextrin) in patients with FSGS. VAR-200 may upregulate cholesterol efflux in podocytes independently of the phospholipid-transporting ATPase ABCA1, a common target for cholesterol efflux-mediated strategies, thereby reducing mitochondrial toxicity from intracellular lipid accumulation. None of the approaches is selective for the GFB, and the safest and most specific way to target mitochondrial health in podocytes and GECs in the setting of human disease is currently unclear. Nevertheless, a positive outcome in these ongoing clinical trials will serve as new impetus to pursue mitochondrial function improvement for GFB disease.

The rest of this section on mitochondrial dysfunction focuses on targets and approaches to intervene in two key pathways: the apolipoprotein L1 (APOL1) pathway and the coenzyme Q10 (CoQ10) biosynthetic pathway. High-risk variants in APOL1 as well as the mutations in the CoQ10 biosynthetic pathway contribute to loss of mitochondrial function, pointing to a common pathogenic mechanism leading to GFB breakdown.

**APOL1 inhibition or silencing**

The APOL1 gene region was originally identified as a risk factor in patients of recent African ancestry who were predisposed to FSGS in the setting of HIV infection. Although the genetic association was initially attributed to a neighbouring gene, MYH9, subsequent genome-wide association studies in patients with idiopathic FSGS identified the same locus but, with reasonable certainty, identified coding variants in APOL1 itself that were strongly associated with FSGS in patients of recent African ancestry. These common variants in the coding region of APOL1 are known as G1 (p.S342G:I384M) and G2 (p.N388_Y389del), and are now established genetically defined risk factors for the development of kidney disease, with relative risk exceeding 20-fold when both alleles contain a variant (that is, APOL1 G1/G1, G1/G2 and G2/G2 genotypes all confer high risk). Nevertheless, many healthy individuals with the high-risk genotypes will not develop kidney disease, indicating that additional factors are required to trigger disease. Follow-up studies have associated the G1 and G2 variants with increased risk of many forms of kidney disease including hypertensive kidney disease, lupus nephritis and sickle cell disease. The G1 and G2 variants are only found in individuals who have current or recent African ancestry, including African Caribbean, African American and Latino individuals. The prevalence of the high-risk genotype in healthy individuals of these ethnicities is 20–40%. These high-risk variants are strongly enriched among patients with kidney disease in these ethnic groups.

APOL1 is only found in humans, gorillas and baboons. In other primates, APOL1 protein is not produced but APOL1 can be found as a pseudogene or gene fragment, indicating evolutionary gene loss. Such an evolutionary pattern is consistent with potential toxicity from APOL1 (Ref. 79). Indeed, in assays performed in a range of systems including human cells and Drosophila melanogaster, the normal form of APOL1 (G0) is toxic to cells. The G1 and G2 forms have an even greater and more extensive toxic gain-of-function phenotype.
This toxicity is partly related to the level of expression, which can be induced by inflammation or infection. Exogenous interferons and infections including HIV and COVID-19 can cause FSGS in patients with the high-risk APOL1 genotype28,90, adding further evidence that G1 and G2 are toxic in the kidney.

APOL1 is produced primarily in the liver and circulates in the high-density lipoprotein (HDL) complex, where its primary function is to kill trypanosomes that enter the bloodstream. One serious trypanosome infection, sleeping sickness, is caused by Trypanosoma brucei rhodesiense and T. brucei gambiense. Sleeping sickness arose in and is endemic to parts of central Africa. T. brucei can evade the protective effects of the normal form of APOL1. It is thought that the APOL1 G1 and G2 variants arose in Africa to specifically kill these resistant trypanosomes. APOL1 oligomerizes to form multisubunit cationic pores in trypanosome lysosomes and mitochondria, resulting in osmotic swelling and mitochondrial depolarization, which kill the organism. Lysosomal pores contribute to osmotic swelling, as does insertion of pores into the outer membrane, but pore insertion in the trypanosome mitochondrial membrane results in mitochondrial depolarization and loss of ATP production, which causes parasite death in a manner analogous to apoptosis81–83.

In addition to circulating in HDL complexes, splice variants of APOL1 that have intracellular membrane localization domains and are not released extracellularly are expressed in the vascular wall throughout the kidney, including in GECs and podocytes. This expression is induced during inflammation when interferon levels are high, particularly in response to viral infection. As with circulatory APOL1, intracellular APOL1 can be toxic to intracellular trypanosomes and may also disrupt viral packaging by inserting into membranes of intracellular vesicles84.

Notably, the intracellular form of APOL1 is also toxic to human kidney cells85–87 and podocytes88,89. The G0 (wild type), G1 and G2 variants form oligomeric structures that can function as cation pores, producing currents that can be detected at the plasma membrane in cells expressing APOL1. However, the G1 and G2 high-risk variants selectively form oligomeric complexes in mitochondria that open the mitochondrial permeability transition pore and cause mitochondrial depolarization in human kidney cells, including podocytes. The G1 and G2 variants also traffic selectively to membranes in the endoplasmic reticulum and mitochondria in podocytes, and may directly introduce APOL1 pores into mitochondrial membranes. Therefore, although the toxic mechanism in podocytes is not fully elucidated, intracellular G1 and G2 variants selectively form multimeric complexes with pore properties in membranes that lead to loss of mitochondrial function, suggesting that mitochondria may be the primary site of toxicity from the G1 and G2 variants72,90 (Fig. 2a). Interestingly, in podocytes, the G1 and G2 variants cause loss of mitochondrial function and cell toxicity only when expressed at high levels, suggesting that a toxic threshold needs to be met.72,90

The tendency of APOL1 to form cation pores has been reproduced in many different laboratories in cell-based and cell-free systems. Pore insertion in kidney cells is the most likely mechanism through which APOL1 injures podocytes in humans, partially because this is the innate immune mechanism through which APOL1 prevents trypanosome infections81. Nevertheless, alternative mechanisms of APOL1 variant-induced podocyte injury have been proposed, including depletion of intracellular K+ and the subsequent induction of stress-activated protein kinase91, abnormal endosomal trafficking and intracellular compartment acidification92, activation of endoplasmic reticulum stress through double-stranded RNA and protein kinase R activation, which inhibits protein translation93,94, and activation of intracellular inflammasomes95. These mechanisms may be activated as a consequence of pore formation, or could contribute to podocyte injury through independent mechanisms.

The role of APOL1 in toxicity in GECs and vascular wall cells beyond the glomerulus is poorly understood. Transgenic expression of human APOL1 variants in podocytes of mice at high levels is sufficient to cause breakdown of the GFB95,96. Nevertheless, APOL1 is induced broadly in the vascular wall, not just the GFB, and APOL1 high-risk variants contribute to the development and progression of hypertensive kidney disease (nephrosclerosis), which is characterized by arteriolar vascular wall pathology as well as disease of the GFB. It is therefore possible that APOL1 contributes to vascular wall toxicity in both arterioles and peritubular capillaries of the kidney, where the pathological features of hypertensive nephrosclerosis are also prominent97.

Two therapies that inhibit the toxic functions of APOL1 are in clinical trials. Vertex Pharmaceuticals has VX-147, a small-molecule oral inhibitor of APOL1, in a phase II trial in adults with FSGS and a G1/G1, G1/G2 or G2/G2 genotype (NCT04340362). The primary end point in this trial is the percentage change in proteinuria or G2/G2 genotype (NCT04340362). The primary end point in this trial is the percentage change in proteinuria. The role of APOL1 in toxicity in GECs and vascular wall cells beyond the glomerulus is poorly understood. Transgenic expression of human APOL1 variants in podocytes of mice at high levels is sufficient to cause breakdown of the GFB95,96. Nevertheless, APOL1 is induced broadly in the vascular wall, not just the GFB, and APOL1 high-risk variants contribute to the development and progression of hypertensive kidney disease (nephrosclerosis), which is characterized by arteriolar vascular wall pathology as well as disease of the GFB. It is therefore possible that APOL1 contributes to vascular wall toxicity in both arterioles and peritubular capillaries of the kidney, where the pathological features of hypertensive nephrosclerosis are also prominent97.

Both the small-molecule APOL1 inhibitor and the ASO targeting APOL1 are potential first in class agents for a disease with high unmet need. Major challenges for these clinical studies, however, include identifying patients who have the G1 or G2 high-risk variants, as patients with FSGS are not routinely genotyped. A second challenge for the ASO will be to demonstrate target engagement in podocytes [BOX 1]. Because both of these candidate medicines block the function of APOL1, they may also block the ability of APOL1 to kill trypanosomes, although the risk of trypanosome infection is low in the developed world. Of note, an individual has been found who is homozygous null for APOL1 [REF97], suggesting that blocking APOL1 will be safe in humans. This individual, who had normal kidney function, presented in India at age 45 years with an infection with Trypanosoma evansi, normally a pathogen of cattle. He was treated with suramin and made a full recovery.

APOL1 may also play a broader role in diseases of the GFB. The risk of developing other kidney diseases including hypertensive nephrosclerosis, lupus nephritis, pre-eclampsia, kidney transplant nephropathy and sickle
Fig. 2 | Therapeutic approaches targeting the GFB in clinical or late preclinical development. Glomerular filtration barrier (GFB), toxic mechanisms that affect podocytes and strategies to address podocyte dysfunction currently in development are shown. a | Apolipoprotein L1 (APOL1) inhibition. APOL1 genetic variants G1 and G2 accumulate in intracellular membranes and structures, particularly mitochondria (inset), where they form oligomers that can function as cation pores and leak cations, such as K⁺, across membranes. Oligomers also bind to mitochondrial inner membrane proteins, activating the mitochondrial permeability transition pore (MPTP), depolarizing mitochondria and causing mitochondrial dysfunction. VX-147 inhibits these toxic functions of APOL1. b | Coenzyme Q10 (CoQ₁₀) restoration. Mutations in genes encoding components of the CoQ₁₀ biosynthetic pathway reduce levels of CoQ₁₀, an electron acceptor of the inner membrane electron transport chain (inset), leading to mitochondrial dysfunction. Pharmacological delivery of CoQ₁₀ or 2,4-dihydroxybenzoic acid (2,4-diHB) bypasses the defects and restores CoQ₁₀ levels in the mitochondrial inner membrane. c | Short transient receptor potential channel (TRPC) inhibition. Mutant forms of TRPC6, overactivated TRPC6 or overactivated TRPC5 increase levels of intracellular Ca²⁺, which acts via calmodulin and calcineurin to activate the transcription factor nuclear factor of activated T cells (NFAT). Intracellular calcium activates RhoA (mainly via TRPC6) and RAC1 (mainly via TRPC5 and calcineurin). These signalling events remodel actin to form stress fibres and disrupt the normal actin–myosin contractile apparatus, resulting in foot process effacement, proteinuria and cell dysfunction. GFB-887 and AM-1473 inhibit TRPC5 and TRPC6, respectively. d | Soluble urokinase plasminogen activator surface receptor (suPAR) inhibition. Elevated levels of circulating suPAR, particularly a fragment containing the D2 and D3 domains, can bind to and activate αvβ3 integrin at focal adhesions on podocytes. This interaction activates focal adhesion kinase (FAK), which remodels actin to form stress fibres and disrupts the normal actin–myosin contractile apparatus, resulting in foot process effacement, proteinuria and cell dysfunction. Antibodies blocking suPAR from binding to αvβ3 integrin are one potential therapeutic approach when suPAR is the disease driver. Cytc, cytochrome c; Dyn, dynamin; GBM, glomerular basement membrane; GEC, glomerular endothelial cell.
Box 1 | Antisense oligonucleotides as candidate therapies in the GFB

Two novel candidate therapies in clinical development to treat diseases of the glomerular filtration barrier (GFB) include antisense oligonucleotides (ASOs) against apolipoprotein L1 (APOL1) mRNA and the non-coding microRNA-21 (miR21). ASOs have a single-stranded DNA backbone, freely enter cells by pinocytosis and form double-stranded RNA–DNA hybrids that induce RNA cleavage by endogenous RNase H, thereby degrading the target. Single-stranded DNA is unstable, can activate innate immune responses and is not able to pass cell membranes. Advances in DNA and RNA chemistry have resulted in substantive chemical modifications to the deoxyribose backbone that serve to stabilize the molecule, protect it from endogenous exonucleases and endonucleases, reduce its charge, render the molecules cell penetrant, reduce innate immune activation and increase affinity for target RNA, increasing activation of RNase H. Such modifications include replacing some phosphodiester bonds with phosphorothioate bonds. Other modifications increase the rigidity between bases, including 2'-O-methyl, and increase affinity of the oligo for the target RNA. These modifications may contribute to translational suppression by a steric hindrance mechanism.

Despite these advances, a major challenge for systemically delivered ASOs is still to get the molecule to the nucleus of the target cells. Subcutaneous or intravenous delivery of ASOs results in high levels of biodistribution to the liver and kidney (mainly to hepatocytes and proximal tubule cells), with lower levels to a range of organs and tissues. Studies of ASO distribution to podocytes and glomerular endothelial cells (GECs) of the GFB have shown limited uptake compared with tubular structures. However, in rodent models, diseased GFBs have an increased uptake of ASOs, and studies from both Ionis and Sanofi have demonstrated high degrees of silencing in the glomerulus in animal models of APOL1 disease or Alport syndrome. Nevertheless, in other organ settings, such as muscle, ASO uptake and target engagement in rodent models did not translate to similar target engagement in human clinical studies. Therefore, a clear understanding of dose and glomerular cell target engagement in large animals, preferably in humans, is required before embarking on interventional studies for glomerular disease. Finally, ASOs have demonstrated mild tubular toxicity at high doses in certain settings, which has limited their advancement to clinical evaluation. Such toxicity findings are idiosyncratic and depend on charge, backbone modifications and sequence, and therefore toxicity can be minimized through careful preclinical evaluation. A recent comprehensive study from Ionis suggests that toxicity to kidney tubules is a minor risk.

cell nephropathy is elevated in patients with APOL1 high-risk genotypes. Success in proof of concept trials could lead to subsequent evaluation in these additional indications, and definition of a group of clinical presentations, all of which include GFB breakdown, as APOL1-mediated kidney diseases.

**Coenzyme Q10 biosynthetic pathway**

SRNS arises in childhood and is frequently identified pathologically as minimal change disease or FSGS on kidney biopsy. Whole-exome sequencing studies of children with isolated SRNS, as well as syndromes that include SRNS, have identified >50 genes that are thought to cause GFB breakdown; the vast majority have an autosomal recessive inheritance pattern. Several of these cluster in the proteins that regulate the biosynthesis of the lipophilic membrane constituent CoQ10, which is a benzoquinone with an isoprenoid side chain. The implicated genes include COQ2, COQ4, COQ6, COQ7, COQ9, PDS51, PDS52, ADCK3 and ADCK4 (refs 105,106). CoQ10 is found principally in the mitochondrial inner membrane, as well as in the Golgi and plasma membranes. In mitochondria, CoQ10 plays a pivotal role in mitochondrial oxidative phosphorylation, as it shuttles electrons between cyclooxygenase complexes I–II and II–III of the electron transport chain. Moreover, it has a critical antioxidant role in mitochondria because of its redox potential. The genes that are mutated in individuals with SRNS contribute to CoQ10 biosynthesis and all the mutations are associated with low levels of CoQ10, dysfunction of the electron transport chain complexes and mitochondrial dysfunction.

To understand the importance of CoQ10 in GFB disease, several studies have inserted human mutations into mice or specifically disrupted individual COQ genes somatically in mouse podocytes. Loss of function of these CoQ10 biosynthetic genes in podocytes is sufficient to recapitulate the hallmarks of human disease: nephrotic syndrome and FSGS. Importantly, in these studies, podocytes with COQ gene mutations have abnormal mitochondrial morphology, dysfunction of the electron transport chain and loss of inner mitochondrial membrane potential. Podocyte cytoskeletal proteins are also reduced, in keeping with a role for mitochondria in maintaining foot processes.

Several clinical case series have now tested a therapeutic role for oral CoQ10 or its synthetic analogue idebenone, and demonstrated that SRNS and FSGS can be stabilized or reduced in children if the disease is treated fairly early in its course (refs 107,108) (Fig. 2b). CoQ10 and idebenone have been used at high doses (30 mg·kg−1·day−1) and have limited oral bioavailability owing to the lipophilic nature of CoQ10, which limits its transport to the mitochondrial inner membrane. The study of the CoQ10 biosynthetic pathway has identified an intermediate, 2,4-dihydroxybenzoic acid (2,4-dHBH), that could be more therapeutically useful than CoQ10. Hydrophilic properties give 2,4-dHBH superior bioavailability to exogenously supplied CoQ10 and exogenous 2,4-dHBH can also bypass many of the loss-of-function mutations and normalize CoQ10 levels (ref. 109). 2,4-dHBH is found naturally in certain foods including coffee and avocados, and has a good safety profile. If given in drinking water to mice with COQ6 or ADCK4 mutations, 2,4-dHBH reversed the progression of disease; the compound also normalized mitochondrial function in cells from a patient with a homozygous COQ7 mutation (ref. 110). This substrate will likely be tested more comprehensively in human studies in the near future, offering the possibility of halting or reversing disease in children with mutations in COQ genes. Additional substrates, including vanillic acid, may also bypass the CoQ10 biosynthetic pathway and elevate levels of this key membrane quinone. One recent study of CoQ10 deficiency in mice identified that the podocytes of these mice had elevated levels of polyunsaturated fatty acids, which can activate stress-activated kinase pathways. Treatment with GDC-0879, a small-molecule inhibitor of the BRAF kinase pathway, rescued podocyte injury and kidney function in those models. These observations suggest that mitochondrial dysfunction in podocytes, as a result of CoQ10 deficiency, may confer toxicity through excessive production of fatty acid metabolites as well as dysfunction of the electron transport chain (ref. 111).

**Outlook on mitochondrial therapies**

The two promising areas of therapeutic intervention in patients highlighted here support an emerging picture that GFB breakdown can result directly from
mitochondrial dysfunction. The selective role of these genetically defined targets in kidney disease suggests that, similar to neurons and myofibres, podocyte function is highly dependent on correctly functioning mitochondria. The strong validation of APOL1 as a target for kidney disease and the high prevalence of APOL1 variants in patients with kidney disease suggest that APOL1 inhibition has the potential to have broad effects on kidney health in humans. Recent studies point to additional mitochondrial functions in podocytes and GECs, such as the control of fatty acid metabolism\(^{111}\), which could uncover new mitochondrial targets. Novel biomarkers are required to study target engagement to guide drug dosing in clinical trials.

### Targeting actin and myosin

The majority of pathological mutations in genes identified to cause isolated FSGS or SRNS are reported to regulate the assembly, turnover and degradation of the actin cytoskeleton or regulate its attachment at the cell surface\(^{114,46}\). The formation and maintenance of podocyte tertiary processes requires complex actin filament assembly and disassembly, and integration at focal adhesions and slit diaphragm protein complexes. Expression of mutated forms of genes including ACTNA, INF2, AHRGAP24 and AHRGDIA in animal models directly interferes with podocyte tertiary process formation\(^{123-114}\). For example, the AHRGDIA gene product is a dissociation inhibitor of the GTPase family (Rho-GDI\(\alpha\)), which locks Rho\(A\) and another Rho family member, RAC\(1\), in their inactive states. Rho\(A\) functions dynamically in the assembly of actin filaments by activating a group of actin polymerizing proteins called formins. Mutant forms of Rho-GDI\(\alpha\) fail to interact with Rho\(A\); podocytes expressing these proteins adopt a hypermigratory phenotype and can no longer maintain the GFB. Another protein that has been found to be mutated, inverted form 2 (INF2), is a member of the actin polymerizing family of formin proteins and plays important roles in cell spreading. Mutations in INF2 act dominantly and fail to promote normal cell spreading\(^{115}\).

Mutations in numerous proteins involved in actin–myosin dynamics are found in individuals with FSGS or SRNS, and these dynamics are vital to maintaining the GFB. Collectively, these findings suggest that normalizing dysfunctional actin–myosin dynamics should have therapeutic utility. Drugging these proteins directly, however, is challenging because of the ubiquitous nature of this bioactive cytoskeleton. Nevertheless, the presence of isolated GFB disease caused by loss-of-function mutations in individual genes of the actin–myosin machinery suggests that facets of its function that are specific to podocytes and GECs could be identified. To that end, studies of podocyte-specific actin–myosin machinery components have identified that a complex containing actin-related protein 2 (ARP2) and ARP3 is central to tertiary process formation through the formation of branched actin networks. In particular, ARP3 plays an important role in maintaining normal podocyte architecture. In the absence of ARP3, the actin–myosin machinery is overactivated, leading to stress fibres and, ultimately, to cell detachment\(^{116}\). It is currently unclear whether ARP2 or ARP3 function could be therapeutically restored in disease settings.

Another potentially druggable target is dynamin. Dynamin, encoded by DNM1, DNM2 and DNM3, is another GTPase that has well-recognized roles in clathrin-mediated endocytosis. This process forms multimeric ring structures that pinch off sections of cell membranes to form endosomes\(^{117}\). Dynamin has also been reported to directly interact with filamentous actin and promote actin assembly, particularly in the formation of filopodia, membrane ruffles and cell spreading\(^{118}\). The dynamin-mediated actin assembly seems to be independent of Rho-GTPases and formins. Several studies suggest that dynamin oligomerization caused either by a gain-of-function point mutation in DNM1 or by the use of a divalent small molecule, Bis-T-23, that stimulates actin oligomerization can overcome both genetic and environmental defects in actin assembly in podocytes, restore foot process formation and restore the GFB\(^{119}\). Bis-T-23 was originally identified as an inhibitor of dynamin-mediated endocytosis and has poor drug-like properties\(^{120}\), and therefore its precise mode of action in restoring the GFB is unclear. Moreover, mutations in dynamin genes have not been associated with diseases of the GFB in humans. Nevertheless, if small molecules that selectively promote dynamin oligomerization can be developed and overcome potential safety liabilities, they may have broad utility in restoring GFB function.

Therapies targeting other aspects of the actin–myosin machinery are further along in development. In the following sections, we discuss therapies in development that target channels and receptors that directly regulate the function of the actin–myosin cytoskeleton in the GFB. In particular, molecules targeting short transient receptor potential channels (TRPCs), roundabout homologue (ROBO) proteins, integrins and endothelin receptors to treat GFB disease are currently in clinical or late preclinical evaluation. The molecules presented are a selection of druggable targets for therapeutic intervention with varying degrees of human target validation.

### Podocyte-directed therapies

**TRPC6 and TRPC5.** Approximately 1–2% of patients with FSGS have mutations in TRPC6, which has an autosomal dominant inheritance pattern. Similar to other TRPCs, TRPC6 is a non-selective cation channel that permits Ca\(^{2+}\) entry into the cell upon activation by stretch, oxidative stress or local factors including angiotensin release. Influx of Ca\(^{2+}\) into podocytes acts as a second messenger in microdomains of the cell, as it regulates actin–myosin assembly and activates transcription factors such as nuclear factor of activated T cells (NFAT)\(^{121}\). TRPC6 localizes to foot processes, and its expression is increased in a range of kidney diseases. Although many patients have mutations that increase channel activity, others have mutations that do not increase channel activity\(^{122}\). Nevertheless, a common feature of disease-associated mutations in TRPC6 is that they increase activation of calcineurin, a calcium-dependent phosphatase that regulates the activity of ERK1/2 and NFAT family member NFATc1 (Ref\(^{123}\)). Mutant forms of TRPC6 activate NFATc1 through a
dominant mechanism that rapidly leads to podocyte effacement and proteinuria. TRPC6 mutations also cause defects in the actin cytoskeleton by acting through the Rho-GTPases, Rho and RAC, and hence contribute to abnormalities of the actin–myosin machinery that maintains foot processes and adhesion to the basement membrane (Fig. 2c). TRPC6 mutations in podocytes contribute to cell detachment and cell death during cell stress. TRPC6 transgenic mice expressing extra copies of wild-type or mutant forms of TRPC6 develop albuminuria. Although the number of patients who have TRPC6 mutations is small, there are several studies in animal models suggesting that wild-type TRPC6 is an important intermediate in the kidney disease cascade downstream of angiotensin receptor type 1 activation, raising the possibility that TRPC6 inhibition may be useful in kidney diseases with high angiotensin levels, such as hypertensive kidney disease.

Ammgen have generated highly potent and selective inhibitors of TRPC6 channel activity, including AM-1473. These inhibitors used novel cryogenic electron microscopy structures of the channel and patient-derived pathological mutations to identify a novel binding site that can be used to selectively block TRPC6 function (Fig. 2c). As yet, no clinical studies in patients have been announced. Selecting the optimal patient population to evaluate such inhibitors may be challenging; genotyping studies of cohorts of patients with FSGS indicate that TRPC6 mutations are rare. An alternative patient population in which TRPC6 inhibition may be assessed is those with hypertensive kidney disease, which has high circulating angiotensin levels. In preclinical studies, overactivity of wild-type TRPC6 has been reported to drive disease pathology. TRPC6 activity appears to require calcineurin and NFAT, a pathway inhibited by the calcineurin inhibitors cyclosporin and tacrolimus. As calcineurin inhibitors are already in use for FSGS, novel TRPC6 inhibitors may not be additive with existing therapies, and will therefore be required to show superior characteristics or a cleaner safety profile. It should be noted that TRPC6 channels play a role in vascular smooth muscle tone and cardiac contractility in pathological settings, so it will be interesting to understand whether TRPC6 inhibition has negative inotropic or hypotensive effects.

A second TRPC family receptor, TRPC5, has also been reported to play a role in calcium and calcineurin signalling in podocytes. TRPC6 has been reported to form a molecular complex with RhoA and inhibit cell migration, whereas TRPC5 has been reported to form a molecular complex with RAC1 and promote cell migration. Unlike TRPC6, no human TRPC5 mutations have been reported to be associated with kidney disease, and levels of expression of TRPC5 do not seem to be regulated in response to disease. Nevertheless, mutations in genes encoding regulatory proteins result in RAC1 overactivation and cause human disease, and TRPC5 loss-of-function in mice protects podocytes and the GFB in settings where podocytes are injured. Selective small-molecule inhibitors of TRPC5 calcium flux have been developed, and demonstrated efficacy in a transgenic rat model of FSGS as well as in angiotensin-driven hypertensive kidney disease in rats. Administration of a small-molecule TRPC5 inhibitor during established GFB breakdown reduced the rate of progression of kidney disease and preserved GFB integrity.

Goldfinch Bio has developed a small molecule, GFB-887, that selectively inhibits TRPC5. The company has initiated a phase II basket study trial (NCT04387448) to evaluate this compound in patients with SRNS, FSGS or diabetic nephropathy (Fig. 2c). The primary end point is the percentage change in the ratio of urine protein to creatinine at 12 weeks. The questions concerning patient selection and demonstration of superiority over calcineurin inhibitors posed above for TRPC6 inhibition also apply to studies of TRPC5 inhibition. It also remains unclear whether TRPC5 inhibition alone will be effective if TRPC6 currents and calcineurin activation remain intact in podocytes. Although TRPC5–RAC1 and TRPC6–RhoA reportedly have opposite effects on podocyte migration, forced activation of either RhoA or RAC1 in podocytes can cause GFB breakdown. It therefore remains unclear whether TRPC5 inhibition will be different from TRPC6 inhibition in the diseased GFB of patients. One potential solution Goldfinch Bio has undertaken is to identify patients with elevated RAC1 signalling as an exploratory biomarker for evaluation of patients receiving GFB-887. A urinary biomarker may identify patients who will be responsive and guide further evaluation of the study drug.

**Soluble urokinase plasminogen activator surface receptor inhibition.** Urokinase plasminogen activator surface receptor (uPAR) is a glycosylphosphatidylinositol-anchored receptor that regulates plasminogen activation. Activated plasminogen is converted into plasmin and acts in clot lysis. However, uPAR is multifaceted as it can be cleaved to soluble forms, known as suPAR. suPAR isoforms have been identified as candidate circulating factors that cause disruption of the GFB in patients whose severe proteinuria recurs after kidney transplantation.

The integrin αvβ3 has long been implicated in cell motility and vascular permeability, but has been most widely studied in thrombosis, arthritis, osteoporosis, inflammation, angiogenesis and viral entry. Activation of
αvβ3 by one of its many ligands, including fibronectin or fibrinogen, changes the cellular cytoskeleton by activating focal adhesions. Focal adhesions are macromolecular complexes through which force can be transmitted from the extracellular matrix to the cell. Such activation may also depend on local shear stress, such as that created by flow. αvβ3 is widely expressed by various cell types in the body. It is detected at low levels on podocytes as well as other vascular cells, including endothelium. As well as ligands such as fibronectin and fibrinogen, αvβ3 may become activated by binding suPAR.

The plasma suPAR concentration has been identified as a robust biomarker of FSGS and other kidney diseases. In addition, its levels predict the development and progression of kidney and cardiovascular diseases in very large patient populations, highlighting the utility of suPAR as a biomarker that predicts outcomes in a number of diseases. suPAR levels, however, do not predict which patients will develop rFSGS after transplantation, and neither do they identify pediatric patient populations with FSGS or SRNS. Blood levels of suPAR do not correlate well with the severity of the phenotype, suggesting that suPAR requires additional factors to play a toxic role. Consistent with this hypothesis, several investigators have not reproduced the toxic gain of function that has been reported in...
Foot process (FP) fusion reduces the number of slit diaphragms, causing protein leakage (FSGS). In both settings there is breakdown of the glomerular filtration barrier (GFB). As observed in pre-eclampsia, endothelial injury may also result in the release of GEC-derived pro-apoptotic factors (right) causing focal segmental glomerulosclerosis (FSGS). A silencing small interfering RNA (siRNA) directed at the placenta compromises the GFB. A glucocorticoid receptor antagonist, the synthetic steroid dexamethasone, reduces circulating levels of sFLT1 in the setting of pre-eclampsia. GBM, glomerular basement membrane; MRL-CNM myosin, non-muscle myosin regulatory light chain.

Fig. 3 | Therapeutic approaches targeting podocyte–endothelial cell crosstalk in clinical or late preclinical development. a Glomerular capillary loops showing injury to glomerular endothelial cells (GECs) from toxins, cellular stress or depletion of podocyte-derived trophic factors. Such injury can result in endotheliosis (left), as observed in pre-eclampsia. Endothelial injury may also result in the release of GEC-derived pro-apoptotic factors (right) causing focal segmental glomerulosclerosis (FSGS). In both settings there is breakdown of the glomerular filtration barrier (GFB). b Roundabout homologue 2 (ROBO2) inhibition. Endothelial cell-derived SLIT2 activates ROBO2 in podocytes to negatively regulate actin polymerization and muscle myosin activity, acting as a counterbalance to nephrin signalling. ROBO2 signals via the adaptor protein slit-robo rho GTPase activating protein 1 (SRGAP1) and protein intermediates NCK and CDC42. A ROBO2–crystallizable fragment (Fc) fusion protein (PF-06730512) blocks ROBO2 activation, enhancing nephrin function in disease states. c Endothelin type A (ETA) receptor inhibition. In response to podocyte stress or injury, ETα receptor is upregulated on GECs and the ligand endothelin (ET1) is released from podocytes. ETα receptor activation in adjacent GECs causes mitochondrial depolarization, resulting in damage from reactive oxygen species (ROS) and a reduction in nitric oxide synthase (NOS). These changes cause loss of GEC fenestrae and degradation of the glycocalyx. Such GECs release pro-apoptotic factors, which in turn cause podocyte stress fibre formation and, ultimately, cell depletion. d Soluble fms-like tyrosine kinase 1 (sFLT1) inhibition. In pre-eclampsia, high levels of sFLT1, an endogenous inhibitor of vascular endothelial growth factor (VEGF) signalling, are released from the placenta into the circulation. Differentiation and maintenance of specialized functions of GECs relies on high levels of VEGF, released from podocytes, engaging VEGF receptor 2 (VEGFR2) on GECs. High concentrations of sFLT1 ‘mop up’ podocyte VEGF and block this tonic signal, causing GEC swelling and dysfunction. Although podocyte foot processes remain intact, the glycocalyx and fenestrae of GECs are disrupted, which is sufficient to compromise the GFB. A silencing small interfering RNA (siRNA) directed at the placenta reduces circulating levels of sFLT1 in the setting of pre-eclampsia. GFB, glomerular basement membrane; MRL-CNM myosin, non-muscle myosin regulatory light chain.

animal models. Overall, through binding to and activating αvβ3, suPAR is likely an important factor in GFB breakdown in some patients.

Several approaches to targeting suPAR are being considered (Fig. 2d). Walden Biosciences is developing neutralizing antibodies that prevent suPAR fragments from interacting with αvβ3 integrin but has not yet announced clinical studies. The molecular hypothesis predicts that inhibition of αvβ3 integrin with antibodies or small-molecule inhibitors could also offer therapeutic benefit. Of note, Janssen/Vascular Therapeutics’ VPI-2690B is an antibody that blocks αvβ3 signalling and has been evaluated for protection of the GFB in diabetic nephropathy. The study completed in 2017 but the results have not been disclosed. Several other small-molecule, peptide and antibody inhibitors of αv integrins (which may include αvβ3, αvβ5 or αvβ1), including MK-0429, cilengitide (EMD121974) and Vitaxin (LM609), have been generated and evaluated for a number of indications, particularly vascular growth in cancers. All have failed to show significant efficacy and are no longer being developed. Although none of them has been tested in rFSGS, these completed studies indicate that αvβ3 integrin inhibition is unlikely to be successful. It will be important to understand whether suPAR has differential binding and activation of αvβ3 integrin or whether other receptors are responsible for the pathological effects of suPAR on the GFB.

The challenges for demonstrating efficacy in humans are significant. Which patients with FSGS have suPAR-driven disease? Should the many patients with FSGS with underlying mutations that cause podocyte dysfunction be excluded? Despite the controversies surrounding the precise toxic role of suPAR, therapies targeting this cytokine merit safe evaluation in patients.

Targeting podocyte–GEC crosstalk

As in many other systems, interdependence exists between cells in the GFB, which cooperate to maintain the structure and function of the barrier. Research has begun to uncover the broad reach of intercellular communications between podocytes and GECs. For example, podocytes and GECs secrete factors and extracellular vesicles that deliver detrimental signals or block trophic signals, and thereby disrupt the barrier and promote disease (Fig. 3a). Intercellular signalling molecules that mediate survival or dysfunction of podocytes and GECs have also emerged as promising therapeutic targets for GFB diseases. Some of these molecules have progressed to clinical evaluation, underscoring the importance of intercellular signalling to health and pathology.

ROBO2–SLIT2 inhibition. ROBOs are receptors found on cell surfaces. ROBOs and their protein ligands, SLITs, were first described as repulsive factors in axon guidance. However, ROBOs and SLITs are more widely expressed and regulate repulsion and guidance elsewhere, including in the vasculature. Loss-of-function mutations in ROBO2 and its adaptor protein, slit-robo rho GTPase activating protein 1 (SRGAP1), are associated with congenital abnormalities of kidney and urinary tract syndrome, which is characterized by vesicoureteral reflux. SLIT2 binds to ROBO2 to direct repulsive signalling through SRGAP1 and limit branching of the ureteric bud, and ROBO2 deficiency results in ectopic or supernumerary collecting ducts. In the glomerulus, SLIT2 and SLIT3 are secreted proteins predominantly derived from GECs and mesangial cells, whereas ROBO2 is exclusively expressed by podocytes. Therefore, GECs and mesangial cells play an active role in regulating tertiary processes and slit diaphragm maintenance through cellular crosstalk (Fig. 3b).

ROBO2 and SRGAP1 are also highly expressed in developing podocytes and expressed at foot processes of mature podocytes, where ROBO2 signalling inhibits nephrin-mediated polymerization of actin, which is required to form tertiary processes. ROBO2, through interaction with SRGAP1, downregulates the GTPase activity of a Rho GTPase, CDC42, which in turn interacts actin polymerization, thereby directly inhibiting foot process extension and function. Loss-of-function of ROBO2 in mouse podocytes results in wider tertiary processes, but the GFB is preserved until adulthood, whereupon very mild dysfunction of the GFB can be detected. However, when ROBO2 was mutated in mice with concurrent nephrin mutations, mice were protected from proteinuria and disease. Rare mutations in the slit diaphragm protein nephrin cause severe childhood-onset SRNS and FSGS. Nephrin has a structural role in the slit diaphragm and also signals via NCK and CDC42 to promote actin polymerization to maintain foot processes. In human diseases of the GFB,
nephrin levels and function are consistently diminished. Inhibition of ROBO2 in such settings would be predicted to counteract disruption of the GFB as a result of diminished nephrin function.

ROBO2 and SRGAP1 directly inhibit foot process elongation by a second mechanism. ROBO2 and SRGAP1 form a complex with non-muscle myosin II proteins, which comprise two heavy chains and two light chains. There are three isoforms of heavy chains encoded by three genes, MYH9, MYH10 and MYH14 [Ref.156]. The non-muscle myosins work with actin to maintain podocyte foot processes, the slit diaphragm and the GFB. Patients with autosomal dominant mutations in MYH9 can present with glomerular disease, including haematuria, proteinuria, nephrotic syndrome and foot process effacement associated with macrothrombocytopenia and hearing loss151.

There is variable penetration of the phenotype, which can occur in childhood but often presents in adults. The disease is thought to be caused by loss-of-function and haploinsufficiency of MYH9. MYH9−/− mice do not develop kidney disease but are more susceptible to injury-induced kidney disease152. In the setting of salt loading, MYH9-deficient mice develop proteinuria; concurrently ROBO2 deficiency protects mice from proteinuria. Absence of ROBO2 in podocytes in mice results in partial protection from a range of toxic and environmental injuries to the GFB including proteamine sulfate and nephrotic serum153. These findings point to regulation of podocyte actin–myosin contractile apparatus playing a significant role in regulating the GFB in health and disease.

Pfizer has developed a protein that is a fusion between ROBO2 and the crystallizable fragment (Fc) antibody domain. This protein, PF-06730512, blocks the interaction of ROBO2 with SLIT2 when administered intravenously. Pfizer is currently evaluating the activity of PF-06730512 in phase II clinical studies in adults with FSGS (NCT03448692) (Fig. 3b). The primary end point is the percentage change in the ratio of urinary protein to creatinine at week 13. Fc fusion proteins have been used successfully to block receptor signalling in other human diseases, but generally have to be delivered intravenously. Challenges to the ROBO2–Fc fusion clinical trial include the risk of redundancy, as both ROBO1 and ROBO4 are also expressed in the kidney, and the need to understand which patients have ROBO2-dependent protein leak, as the level of expression of ROBO2 may be an important factor in its pathogenic role. In addition, because ROBO2 deficiency in mouse podocytes itself results in adult-onset mild proteinuria, it will be important to understand the therapeutic window, dose–response curve and pharmacodynamics of the fusion protein in humans to avoid excessively blocking the pathway.

**Endothelin receptor A inhibition.** Endothelin 1 (ET1; also known as Edn1) is a potent vasoconstrictor that acts by binding to endothelin type A (ETA) receptor and ETB receptor, which are expressed in a wide variety of cell types154. Increased urinary ET1 excretion has been observed in patients with diabetic kidney disease or FSGS155–157. Variation in the regulation or expression of ET1 or ETB receptor in humans is associated with vascular diseases including coronary artery disease, hypertension, renal artery stenosis, arterial dissection and migraine, but no genetic association with disease of the GFB has been demonstrated158,159.

ET1 has been investigated by infusion in healthy humans160 and rodents161 in the setting of diabetes162 and pre-eclampsia163, and in rodent models of FSGS157. Whereas ET1 activation of ETα receptor results in vasoconstriction and systemic blood pressure. Infusion of ET1 directly increased glomerular permeability to albumin and renal inflammation via ETα receptor activation, independent of changes in arterial pressure164.

ET1-induced contraction of vessels occurs through RhoA activity and phosphatidylinositol 3-kinase (PI3K) activity, which is induced downstream of ETα receptor and increases the level of myosin light chain phosphorylation163. Activation of RAC1 by ET1 contributes to the hypertrophic response of cardiac myocytes by modulating mitogen-activated protein kinase (MAPK) signalling cascades164. ET1 can also mediate disruption of the F-actin cytoskeleton and promote dysfunction of the slit diaphragm through activation of the Rho and PI3K pathways via ETβ receptor in a mouse immortalized podocyte line165,166. These preclinical studies point to an important role for ET1 in proteinuric chronic kidney diseases and highlight the potential for ETα receptor inhibition in maintenance of the GFB167,168.

In the glomerulus, ET1 and its receptors are expressed by mesangial cells, GECs and podocytes and have a specific local role in maintaining the GFB. ET1 has been reported to signal via both ETα and ETβ receptors in podocytes and drive the development of glomerulosclerosis157. In hypertensive rats, selective ETα receptor blockade prevented renal fibrosis and glomerulosclerosis via inhibition of RhoA and Rho-associated protein kinase (ROCK) activity170,172. An important role for ET1–ETα receptor signalling in the GFB was highlighted by three findings: ET1 receptor expression was only detected in GECs of human biopsies and mouse kidneys in samples that had signs of diabetic kidney disease42 or FSGS157,172; ETα receptor-expressing GECs exhibit mitochondrial oxidative damage; and pre-proendothelin (ET1 precursor protein) is produced locally in high quantities by adjacent podocytes43,174. In these studies, ET1 released from podocytes induced ETα receptor-dependent signalling in GECs. This signalling caused endothelial dysfunction characterized initially by loss of GEC fenestrations and loss of the glyocalyx, followed by podocyte injury and loss of the GFB149,150,152,154. Selective ETα receptor antagonism prevented loss of the glyocalyx in mice with type 2 diabetic kidney disease and ameliorated proteinuria173,174. The studies found an unexpected interdependence between GECs and podocytes of the GFB in which ET1–ETα receptor directly contributes to loss of the GFB, and support the notion that selective pharmacological inhibition of ETα receptor can directly preserve the GFB.

Endothelin receptor antagonists have been evaluated in patients with proteinuric glomerular diseases.
including diabetic kidney disease and FSGS, have demonstrated encouraging anti-proteinuric effects and may stabilize kidney function177,178. The SONAR phase III clinical trial, which tested the selective ET<sub>a</sub> receptor antagonist atrasentan versus placebo in people with type 2 diabetic kidney disease, showed a 35% reduction in the relative risk of composite renal events177. The trial was prematurely terminated owing to fluid retention-related adverse events; however, adverse events related to fluid retention may be less of an issue in patients with little or no cardiovascular disease. Atrasentan is entering new phase III studies in rare diseases of the GFB (NCT04573920). In FSGS, the phase II DUET trial found that atrasentan, a combined ET<sub>a</sub> receptor inhibitor and angiotensin receptor blocker (ARB), reduced proteinuria (urinary protein to creatinine ratio) by ~50% compared with ARB treatment alone in people with primary FSGS180. In the 84-week open-label extension, the investigational agent was well tolerated, stabilized kidney function and reduced proteinuria to near normal levels in 40% of those studied. Following the success of the proof of concept study, a phase III DUPLEX trial has recruited 300 patients with FSGS with a primary end point of change in the estimated glomerular filtration rate, which measures the change in glomerular function over time (NCT03541174). An interim analysis of the proteinuria effect will be available in 2021. ET<sub>a</sub> receptor inhibitors are being pursued as a treatment for other glomerular and renal vascular diseases, including IgA nephropathy (phase III PROTECT; NCT03762850) and in patients with kidney disease caused by systemic sclerosis (phase II ZEBRA; NCT02047708).

**Outlook on actin–myosin therapies**

The expanding list of mutations in genes involved in actin–myosin assembly and disassembly provides important insight into the underlying functions of podocytes in the GFB. Together with evidence from drug toxicities that cause kidney disease by challenging the actin cytoskeleton architecture in podocytes181, or from other acquired perturbations in cytoskeleton regulation in podocytes, targeted therapies that stabilize actin-associated proteins in podocytes hold the potential to maintain the structural integrity of podocytes and treat GFB diseases.

**Targeting systemic factors: sFLT1**

Disruption to the GFB can be precipitated by systemic physiological changes. For example, pre-eclampsia, a serious condition that occurs in 2–8% of all pregnancies, is responsible for thousands of deaths annually, particularly in low and middle-income countries, where an infrastructure of surveillance and management is not readily available. Proteinuria is one of the diagnostic criteria of pre-eclampsia; others include hypertension and oedema. Approximately 25% of patients with pre-eclampsia have high-grade proteinuria and nephrotic syndrome. Biopsies of pre-eclamptic kidneys typically show endotheliosis and thrombotic microangiopathy with vacuolated podocytes. GECs are swollen, lose fenestrations, and frequently show sub-endothelial fibrinoid deposits182. Podocytes, however, usually are present in normal numbers and have preserved their foot processes. This observation demonstrates that it is possible to disrupt the GFB without disrupting podocytes [FIG. 3a].

Proteomic analysis of patients’ plasma samples led to the identification of high circulating levels of a soluble decoy receptor, soluble Fms-like tyrosine kinase 1 (sFLT1; also known as soluble vascular endothelial growth factor receptor 1 (sVEGFR1)), in many patients with pre-eclampsia183,184. The abnormally high levels of sFLT1 arise from the placenta weeks before clinical disease185,186. The placenta is a rapidly growing highly vascularized organ. It is thought that sFLT1 is an important negative regulator of placental vascular growth and that excessive sFLT1 production results in inadequate new vessel formation, leading to fetal growth retardation. Excessive sFLT1 spills into the maternal circulation and can exert anti-angiogenic activity by binding circulating VEGF, thereby blocking signalling from VEGF receptor 2 (VEGFR2) [REF. 185]. In the kidney, sFLT1 inhibits the activation and signalling of VEGFR2 on GECs by VEGF from adjacent podocytes. VEGF2 signalling is critical for GEC maturation, migration into the glomerular tuft, maintenance of fenestration and normal GEC function185,186. GEC function in pre-eclampsia can also be affected by reduced levels of pro-angiogenic placental growth factor or increased levels of soluble endoglin, which blocks endoglin–TGFβ signalling.

Disturbance of these signalling pathways from placent-derived factors may also have deleterious effects on the GFB181. In support of a critical role for sFLT1 in pre-eclampsia, anti-VEGF antibody therapy, which is used to treat some cancers, can cause GECs to lose fenestrations and function, leading to loss of GFB integrity as an on-target side effect186,187. These clinical findings underscore the impact of VEGF sequestration on GEC function, and also highlight the role of cell–cell crosstalk between podocytes and GECs in maintaining the highly differentiated endothelial state that is required for an effective GFB [FIG. 3d].

The mechanisms of the pathological overflow of sFLT1 from the placenta in pre-eclampsia are unknown. However, the ensuing endothelial dysfunction characterized by impaired vasodilation results from a decrease in nitric oxide synthase (NOS) activity and nitric oxide (NO) bioavailability (normally stimulated by VEGF) in the endothelium and increased oxidative stress, which combine to further stimulate systemic hypertension185,189. Low levels of endothelial NO therefore likely exacerbate glomerular disease and contribute to proteinuria190–194. Degradation of the GEC glycocalyx has also been linked to reduced VEGF signalling by sFLT1 in pre-eclampsia, and may contribute to GFB permeability195–197 [FIG. 3d].

GEC dysfunction is the main feature of glomerular involvement in pre-eclampsia. Nevertheless, the reduced expression of podocyte slit diaphragm proteins nephrin and synaptopodin as well as the detection of detached podocytes in the urine are consistent with podocyte injury198. Although homeostatic local sFLT1 production has been shown to stabilize podocyte function, excess sFLT1 can also prevent autocrine VEGF from binding to receptors on podocytes, thereby directly affecting the podocyte actin cytoskeleton189,190.
Pre-eclampsia is well managed in the USA with antihypertensive medications, and results in few deaths. Nevertheless, serious complications including liver injury, acute kidney injury, bleeding and seizures are frequent. Clinical studies involving sFLT1 elimination by dextran sulfate apheresis in humans have shown promise in patients with severe disease; apheresis reduces proteinuria and prolongs pregnancy209, but does not effectively remove sFLT1. Of note, the therapeutic goal is to reduce sFLT1 levels by 30–40%, as a higher degree of reduction can result in reduced uterine blood flow202,203.

An academic team from The University of Massachusetts RNA Therapeutics Institute has developed a novel therapeutic approach that blocks the production of sFLT1 by the placenta in rodent and primate models using small interfering RNA (siRNA)-mediated silencing of sFLT1 mRNA. Cholesterol conjugation of siRNAs resulted in stable and sufficient accumulation in rodent and baboon placenta and selectively lowered circulating sFLT1 protein levels by 50%206. This, in turn, alleviated hypertension and proteinuria of pre-eclampsia that was induced in the preclinical models, without adverse consequences to the developing fetus. Therefore, an siRNA-based therapeutic for pre-eclampsia could be highly effective. This therapeutic approach may be taken forward by Tiger Therapeutics.

### Enabling technologies

In addition to the advances in target identification through family studies, genomics, systems biology and proteomics highlighted in the preceding sections, several other breakthroughs have occurred in drug discovery that will enable the advancement of candidate molecules targeting the GFB. With advances in cell isolation and cell culture methods, detection of in situ binding and assays using fluorescent read-outs, it is possible to perform secondary screens of candidate molecules on human patient-derived podocyte monocultures. Such assays may be more relevant than cell lines that have little or no relevance to GFB disease. Still, these monocultures bear only a superficial relationship to podocytes in vivo, and do not recapitulate the complex cellular interplay between GECs and the GBM205,206. Protocols for establishing mature podocytes from human induced pluripotent stem cells have been developed207, and are now being integrated into more complex model systems such as 3D cultures and organoids. These emerging technologies can enable low- to medium-throughput evaluation of candidate therapies in model microfluidic systems that contain cells derived from patients. For example, a glomerulus on a chip that hosts GECs and podocytes separated by a porous and flexible poly(dimethylsiloxane) (PDMS) membrane, together with fluids and strain to better mimic the structure and function of the glomerular capillary wall, can measure the function of the permeability barrier that normally prevents proteinuria208. Other developments include a glomerulus on a chip in which endothelial cells and podocytes interact to generate a layer of basal lamina that resembles the GBM, whereas GECs develop a glycocalyx in this system209. Exposure of chips to the serum from patients with membranous nephropathy, or use of podocytes derived from a patient with Alport syndrome, showed protein leakage, supporting the chip’s potential for use in personalized medicine210. These advances allow the evaluation of podocyte function in maintaining the GFB and the assessment of effects of candidates on restoring the GFB in a highly controlled setting210,211. Modern homologous recombination and transgenic technologies can be used to place human genes and their regulatory sequences into rodents and test the effects of candidate therapies in vivo against the human target211. This has proved important for establishing target engagement and dose in vivo, with the caveat that allometry (modelling dose and pharmacokinetic properties from one species to another) remains a fine art.

In addition to developing better model systems, several additional important advances made over the past decade could enable therapies targeting the GFB to be developed. Structure-based drug design212, enabled by the refinement and industrialization of cryogenic electron microscopy213, has helped to identify compounds with sub-nanomolar effective inhibitory concentrations. In silico predictive tools have helped tailor arrayed small-molecule screens to focus on classes of compounds, rather than all compounds214, and advances have been made in pooled small-molecule screens using DNA-barcoded libraries of small molecules215.

The use of in silico tools to predict toxicity, metabolism, production of toxic metabolites or drug–drug interactions has improved the early de-prioritization of molecules with potential toxicity215. Screening promising small molecules using standardized assays for potential cardiac toxicity, hepatotoxicity and genotoxicity has become standard practice in the early stages of small-molecule improvement from initial screening hits, but screening against renal toxicity remains challenging. However, human kidney tubule 2D and, particularly, 3D cultures have shown promise in detecting toxicity in panels of known molecules with nephrotoxicity216.

Finally, new therapeutic modalities, including nucleic acid therapies such as ASOs (BOX 1), are proving robust in targeting the GFB. Current algorithms enable the design of ASO molecules that not only engage the target mRNA or microRNA for protracted periods but also appear to adequately engage the target mRNA in the nucleus of podocytes and GECs. Other therapeutic approaches are also being identified that can precisely correct the disease-causing variants and mutations in the GFB. These DNA editors, including CRISPR–Cas9, base editors and prime editors, have the potential to precisely correct the underlying cause of kidney diseases, if they can be delivered to cells of the GFB successfully218,219.

### Outlook and future directions

New insights into genetics, defined environmental insults and the structure–function of the GFB have resulted in new candidate therapeutic targets for kidney disease that selectively target the GFB, and offer the possibility of tailored therapies that correct defects within or between cells of the GFB. Modern structure-based small-molecule drug design, advances in nucleic acid therapies and protein therapies have enabled successful drug candidates to emerge that are now in clinical evaluation. The use
of precision medicine will be required to make these candidate therapies successful in clinical practice.

One of the themes in this Review has been the selection of the right patients to evaluate investigational drugs. An increasing number of trials fail to meet their primary end points, and then a post hoc analysis identifies a subgroup in whom the drug was effective. Development of clinically available biomarkers to identify kidney injury and monitor therapy effectiveness will help guide clinical decisions in the future. For APOL1 inhibitors, their efficacy will likely be seen only in patients who have a diagnosis of FSGS on kidney biopsy and carry high-risk variants on both copies of APOL1. Similarly, an siRNA against sFLT1 will probably only work in patients with pre-eclampsia who also have elevated levels of sFLT1 in the circulation. Kidney disease has long been defined by pathological descriptors on biopsy, but modern genetics and other techniques can increasingly identify patients with a potential genetic or immunological cause, and therapies targeting the GFB are becoming more tailored to those causes.

Although certain rare kidney diseases that alter the GFB are caused by inherited mutations, most adult-onset disease has previously been thought to be acquired. Such conclusions are now being challenged by the finding that common forms of kidney disease, such as hypertensive nephrosclerosis and diabetic kidney disease, occur in patients with heterogeneous loss-of-function mutations or high-risk variants that are identical to those that cause rare, early-onset disease. For example, mutations in uromodulin (encoded by UMOD) and collagen type IV trimers (encoded by COLA43, COLA44 and COLA45) cause autosomal dominant tubulointerstitial kidney disease and Alport syndrome, respectively, but the same mutations or common variants are also detected in patients presenting with FSGS, hypertensive kidney disease and diabetic kidney disease. The continued whole-exome sequencing of patients with common forms of kidney disease is likely to refine current targets and identify novel genetically defined therapeutic targets. It is likely that more routine genotyping of patients with kidney disease will determine their eligibility for the next generation of GFB targeted therapies.

Nevertheless, improved and validated assays of human disease are required to investigate promising therapeutic targets, enable drug discovery and provide the confidence needed to advance these therapies to the clinic. Current animal models are poor predictors of human disease. Patient-derived 3D cell systems could bridge this translational gap.

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