Noncoding RNA therapeutics — challenges and potential solutions

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Abstract | Therapeutic targeting of noncoding RNAs (ncRNAs), such as microRNAs (miRNAs) and long noncoding RNAs (lncRNAs), represents an attractive approach for the treatment of cancers, as well as many other diseases. Over the past decade, substantial effort has been made towards the clinical application of RNA-based therapeutics, employing mostly antisense oligonucleotides and small interfering RNAs, with several gaining FDA approval. However, trial results have so far been ambivalent, with some studies reporting potent effects whereas others demonstrated limited efficacy or toxicity. Alternative entities such as antimiRNAs are undergoing clinical testing, and lncRNA-based therapeutics are gaining interest. In this Perspective, we discuss key challenges facing ncRNA therapeutics — including issues associated with specificity, delivery and tolerability — and focus on promising emerging approaches that aim to boost their success.

Noncoding RNAs (ncRNAs) are generated from the larger part of the genome that does not encode proteins but produces noncoding transcripts that regulate gene expression and protein function. The two major classes of ncRNA are the well-studied short microRNAs (miRNAs) and the more recently identified long ncRNAs (lncRNAs) (Box 1). Deregulation of both types of transcript has been linked to every major cancer investigated to date and affects all major cancer hallmarks. In addition, they have been linked to complex biological processes such as immune cell development and function, immune disorders, neural development and neurological diseases. Therapeutic targeting of such naturally occurring ncRNAs thus represents a very promising approach for the treatment of various diseases.

Various RNA-based therapies, including antisense oligonucleotides (ASOs), small interfering RNAs (siRNAs), short hairpin RNAs (shRNAs), ASO anti-microRNAs (anti-miRs), miRNA mimics, miRNA sponges, therapeutic circular RNAs (circRNAs) and CRISPR–Cas9-based gene editing have been developed and several excellent reviews describe these agents (Box 2). Currently, 11 RNA-based therapeutics are approved by the FDA and/or the European Medicines Agency (EMA) (Table 1), aiming at gene modifications in liver, muscle or the central nervous system. All these therapeutics are either siRNAs or ASOs that cause specific gene downregulation, or ASOs that target pre-mRNA splicing (that is, inducing exon skipping or inclusion). In addition, a plethora of RNA therapeutics are in phase II or III clinical development, including newer entities such as miRNA mimics and antimiRs (Table 2), but no lncRNA-based therapeutics have entered the clinic.

The use of miRNA-based therapeutics has dual advantages. First, miRNAs are naturally occurring molecules in human cells, unlike man-made chemotherapy compounds or ASOs, and therefore have all the mechanisms in place for their processing and downstream target selection (Fig. 1). Second, miRNAs act by targeting multiple genes within one pathway, thus causing a broader yet specific response. The miR-15–miR-16 cluster, downregulating multiple anti-apoptotic factors including BCL-2 and MCL1, represents an excellent example of a miRNA acting at multiple levels to affect the same cancer hallmark. The use or targeting of naturally occurring miRNAs could therefore represent a promising alternative to existing RNA-based therapies and may potentially boost therapeutic effects compared with synthetic siRNAs or ASOs that influence only a single target gene.

The diverse functional repertoire of lncRNAs reveals various opportunities for their therapeutic targeting, the means of which need to be adjusted to the mode of action of the lncRNA. LncRNA targeting may include transcriptional inhibition, post-transcriptional inhibition, steric hindrance of secondary structure formation or protein interactions, introduction of synthetic (for example, circular) lncRNAs, and modulation of lncRNA genomic loci or expression patterns via CRISPR–Cas9 and CRISPR–Cas13, respectively (reviewed in ref. ). An interesting development is the exploration of natural antisense transcripts (NATs): lncRNAs that are transcribed in the antisense direction to coding genes, and negatively regulate them in cis. ASOs that target NATs, termed ‘antagoNATs’ have shown very promising preclinical results for gene reactivation in the central nervous system. AntagoNATs successfully upregulated brain-derived neurotrophic factor (BDNF), a protein highly involved in memory formation, as well as the healthy allele of SCN1A, the haploinsufficiency of which causes the brain disorder Dravet syndrome. Notably, BDNF-AS-targeting antagoNATs were successfully delivered across the blood–brain barrier using a minimally invasive nasaldepop (MIND) in a mouse model. MIND is aimed at directing drug delivery to the olfactory submucosal space and achieved approximately 40% efficacy compared with riskier invasive delivery techniques. Such promising developments suggest that the entrance of lncRNA-based therapeutics into clinical testing is imminent.

The translation of all RNA-based therapeutics into the clinic has been hampered by issues associated with specificity, delivery and tolerability. Specificity issues include undesired on-target effects due to uptake in cells other than the cells of interest, or off-target effects caused by either sequence similarities or overdosing to levels much higher than expected endogenously. Delivery issues are related to three major points: the instability of ‘naked’, chemically unmodified RNA structures; their inefficient intracellular delivery, which requires the exploitation of endosomal
escape mechanisms; and the lack of delivery vehicles suitable for targeting the organ and cell type of interest. In line with this, clinical trials have been terminated most often owing to a lack of efficacy (TABLE 3), one example being the nuclease-resistant ASO Genasense (G3139) that targets BCL2 miRNA, which was discontinued owing to limited therapeutic effect; this stands in sharp contrast to the very successful use of venetoclax, a small-molecule inhibitor of the BCL-2 protein. Tolerability issues are caused by the recognition of RNA structures by pathogen-associated molecular pattern (PAMP) receptors, such as Toll-like receptors (TLRs), leading to adverse immune effects. A prime example is the miR-34 mimic MRX34, which caused significant adverse reactions in five patients, including a case of cytokine release syndrome, in a multicentre phase I clinical trial in patients with advanced malignancies. Conversely, miR-16 restitution therapy in patients with mesothelioma (MesomiR-1) was second, the processing of pri-miRNAs by the nuclear ribonuclease Drosha and Dicer5 into a 70 nt long precursor miRNA (pre-miRNA) with a stem–loop structure; third, the nuclear export of pre-miRNAs by exportin 5 and Ran-GTPase; and fourth, the cleavage of pre-miRNAs by RNase III enzyme Dicer to yield a mature, double-stranded miRNA. Gene regulation is mediated through the unwinding of the miRNA by RNA helicase and the incorporation of the miRNAguide strand into the RNA-induced silencing complex (RISC)276–279. Post-transcriptional gene silencing is mediated via nucleotide complementarity to mainly the 3′ UTR, or less prevalently the 5′ UTR or coding region, of target miRNAs278,279. The miRNA ‘seed sequence’ is situated from nucleotides 2 to 7 at the 5′-end of the miRNA sequence. Seed sequence binding with perfect complementarity results in the deadenylation and degradation of the targeted mRNA, whereas binding with imperfect complementarity, which is more common, results in translational inhibition, both of which are facilitated by RISC.

Long noncoding RNAs (lncRNAs) are larger transcripts (>200 nt in size) that are synthesized similarly to miRNAs, but not translated into protein55. LncRNAs contain two types of functional element, the interactor elements involved in direct physical interaction with other nucleic acids, with proteins or lipids, and the structural elements, leading to the occurrence of secondary and/or tertiary 3D RNA structures, which direct their functional interactions281. It is this capacity to interact with DNA and RNA as well as proteins via base pairing in linear form or chemical interactions in secondary structures, that allows lncRNAs to function in more variable ways than miRNAs. Gene-regulatory roles have been identified for many lncRNAs, for example, by influencing transcription factor binding or epigenetic marks. Interactions with miRNAs may influence their stability or rate of translation. Similarly, lncRNA–protein interactions may influence the stability, activity or localization of the protein282,283. Furthermore, circular RNAs, which have a similar length range to lncRNAs, have become known for their potent roles as miRNA sponges.

In this Perspective, we discuss potential approaches to address the drawbacks that have been experienced regarding the clinical translation of RNA-based therapeutics so far, including the hurdles of immune responses, low specificity and aspecific delivery, with a particular focus on miRNA- and lncRNA-based therapeutics. By giving an overview of new and promising preclinical and clinical advancements, we hope to further facilitate the development of RNA-based life-saving interventions for the treatment of any type of disease in which RNAs are pathogenic, including RNAs produced from the human genome and RNAs from xeno-genomes such as RNA viruses (for example, SARS-CoV-2) (Box 3). The hurdle of immunogenicity

As a viral defence mechanism, our immune system recognizes both single-stranded (ss) and double-stranded (ds) RNA via diverse extra- and intracellular PAMP receptors. Extracellular recognition is mediated by TLRs 3, 7 and 8 in endosomes, and intracellular recognition is mediated by receptors in the cytoplasm such as EIF2AK2, RIGI, IFIH1 and NOD1/2. Immune activation by RNA interference (RNAi) was first described in vitro, with seven individual siRNAs causing a IFN-α-mediated response through cytoplasmic recognition of the 21 nt dsRNA molecules by the tyrosine kinase EIF2AK2 (REF.27). Although delivery of naked siRNA was first shown not to trigger an interferon response in animals, intraocular injections of anti-vascular endothelial growth factor receptor 1 (VEGFR1) siRNAs bevasiranib (NCT00499590) and AGN 211745 (NCT00395057) were later shown to owe their anti-angiogenic effects to direct TLR3 stimulation instead of to the desired silencing effect, resulting in termination of clinical development (TABLE 3). The predominant pathway recognizing RNA therapeutics is TLR signalling, which is mediated through myeloid differentiation factor 88 (MyD88) and activates various pathways, resulting in nuclear factor-κB (NF-κB) activation and the production of pro-inflammatory cytokines (IL-6, IL-8, IL-12 and TNF), or a type I interferon response, which ultimately leads to the activation of diverse downstream immune responses. TLR7 and TLR8 stimulation by miRNA was shown to be dependent on the presence of GU-rich sequences (for example, 5′-UGUGU-3′ or 5′-GUCCUCUCAA-3′) and caused a IFNα-mediated inflammatory response, activation of NF-κB and production of inflammatory cytokines (for example, IL-6 and TNF) in dendritic cells and macrophages. This effect observed in vitro after transfection and in vivo after delivery using liposomes was triggered by both ssRNA and dsRNA and occurred in the endosomal compartment.

It was furthermore shown that ssRNA is more prone to cause immune stimulation than dsRNA; consequently, all siRNAs currently in clinical use or development are double-stranded (TABLE 1, 2).

Endogenous miRNAs may also act as TLR agonists, as shown for multiple tumour-secreted miRNAs. Lung tumour cell-derived, exosome-secreted miR-21 and miR-29a could reach the endosomal compartment of surrounding macrophages to activate TLR8 (and TLR7 in mice), causing NF-κB activation and production of inflammatory cytokines (for example, IL-6 and TNF). Of note, the miRNA-mediated activation of TLR8 in tumour-associated macrophages has been shown to be involved in the acquisition of resistance to chemotherapy. The let-7 miRNA family caused TLR7 stimulation in macrophages, microglia and neurons, which was also dependent on the presence of GU-rich sequences (5′-UGUGU-3′ and variants thereof). Moreover, not only cellular miRNAs, but also viral miRNAs directly activate TLRs and their downstream signalling, showing that this is a more widespread response to miRNAs than initially reported. Single-stranded small
GU-rich RNAs putatively transcribed from the SARS-CoV genome also induced inflammatory cytokine secretion (TNF, IL-6, IL-12) via TLR7 and TLR8 stimulation in vitro44. Second-generation chemical modifications now commonly applied to RNA-based therapeutics were developed to reduce the immunostimulatory potential of synthetic RNA therapeutics. 2′-ribose modifications on siRNAs (for example, 2′-F, 2′-O-Me and 2′-H), especially when applied to uridines within GU-rich sequences, can abrogate TLR stimulation38,45,46. The presence of 2 nt 3′-overhangs, as present in endogenous miRNAs, assists evasion from RNA recognition protein RIG1, which only responds to blunt-ended short RNAs42. However, 5′-triphosphate-modified siRNAs caused RIG1 recognition44, showing that chemical modifications may also be immunogenic. Despite significant advancements, there are still reports of adverse immune responses in clinical trials employing such modified reagents. For example, systemic injection of a liposomically-delivered siRNA targeting apolipoprotein B (TKM-ApoB, PRO-040201) for treatment of hypercholesterolaemia was terminated after phase I owing to the occurrence of flu-like symptoms in a patient receiving the highest dose (see Related links). In addition, while the 23 nt ds miR-34a mimic MRX34 had shown promising preclinical results regarding efficacy and safety, with liposome-delivered MRX34 demonstrating efficient antitumour effects in preclinical studies49,50, the first-in-human MRX34 clinical trial was terminated owing to immune-related adverse effects in five patients, which included enterocolitis, hypoxia, systemic inflammatory response syndrome, cytokine release syndrome, hepatic failure and respiratory failure41. This potent immune effect was surprising, as there had been no reports of immunogenicity of miR-34a in preclinical studies as measured by IL-1β, IL-6 and TNF secretion in mice41. Notably, a study using the same delivery vehicle to deliver ssDNA molecules (PNT2258) also showed no evidence of immune stimulation41. In non-human primates, MRX34 most strongly

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**Box 2 | Types of RNA-targeting therapeutics**

RNA-targeting therapeutics are used to induce miRNA-like functions, restore or deplete the levels of a microRNA (miRNA), or to inhibit the interaction of a miRNA with its targets. As RNA therapeutics are naturally unstable and unable to cross cell membranes owing to their negative charge, various chemical modifications are applied to improve their pharmacokinetics and pharmacodynamics (reviewed in REFS 291,292). First-generation modifications replace phosphodiester with phosphothioate (PT) backbone linkages to improve stability. Fomivirsen, the first RNA-based therapeutic to be approved for clinical use in 1998 was a first-generation antisense oligonucleotide (ASO) targeting the cytomegalovirus (CMV) IE-2 mRNA for treatment of CMV retinitis (TABLE 1). Second-generation modifications replace the 2′-O-alkyl group of the sugar moieties with, for example, 2′-O-Me, 2′-MOE or 2′-F to improve bioavailability, enhance efficacy and reduce toxicity and immunostimulation. As 2′-sugar modifications inhibit RNase H activity, second-generation ASOs are chimeric molecules (‘gapmers’) with a central stretch of DNA monomers (to support RNase H cleavage) flanked by 2′-modified nucleotides. Third-generation chemical modifications apply changes to the furanose ring to create, for example, locked nucleic acids (LNAs), peptide nucleic acids (PNAs) and phosphorothioate morpholino oligomers (PMOs). All RNA therapeutics currently approved for clinical use carry second- or third-generation chemical modifications (TABLE 1). ASOs are single-stranded DNA molecules with full complementarity to one select target mRNA and may act by blocking protein translation (via steric hindrance), causing mRNA degradation (via RNase H cleavage) or changing pre-mRNA splicing (via interference with cis-splicing elements causing exon inclusion or exclusion)293,294. Small interfering RNAs (siRNAs) may be single or double stranded and exploit the endogenous miRNA pathway and mediate silencing of one, fully complementary mRNA via their loading into the RNA-induced silencing complex (RISC)295. Short hairpin RNAs (shRNAs) exploit the miRNA maturation pathway, being cleaved by Dicer into a double-stranded mature product before RISC loading. ShRNAs are traditionally introduced into cells using viral vector systems such as adenovirus-associated viruses, retroviruses or lentviruses. Bisfunctional shRNAs have higher knock-down efficacy as they simultaneously produce transcripts with perfect and imperfect complementarity to target degradation as well as translational inhibition296. Two liposomally delivered bisfunctional shRNA constructs are currently being tested in phase I clinical trials: pbi-shRNA EWS/FLI1 (REF. 297), which targets the mRNA creating the EWS–FLI1 fusion protein, a driver of Ewing sarcoma (NCT02736565), and pbi-STMN1 (REF. 297), which targets stathmin 1 mRNA in advanced solid cancers (NCT01505153). MiRNA mimics exploit the main advantage of endogenous miRNAs being able to target multiple mRNAs at once. miRNA mimics have the same sequence as an endogenous miRNA while the passenger strand carries a few mismatches to prevent RISC loading and potential action as an anti-microRNA (antimiR)298. Two miRNA mimics have been tested in clinical trials for cancer treatment: the miR-34 mimic MRX34 (REFS 299,300) (TABLE 3) and the miR-16 mimic Mesomir-1 (REF. 301). AntimiRs are essentially ASOs designed to be fully or partially complementary to an endogenous miRNA to prevent the interaction with its target genes. AntimiRs may also be referred to as ‘antagomiRs’ if they are conjugated to cholesterol to improve intracellular delivery302. Two miR-122 antimiRs have been clinically tested as novel hepatitis C virus (HCV) therapeutics, RG-101 (N-acetylgalactosamine-conjugated ASO; TABLE 3) and miravirsen (SPC3649; β-oxyo-LNA)303 (TABLE 2, BOX 3). Moreover, anti-miR-92a (MRG-110) was tested for its capability to induce angiogenesis and improve wound healing (NCT03603431), and anti-miR-21 (RG-012) was tested for its ability to prevent kidney fibrosis in patients with Alport syndrome (NCT03737386). MiRNA sponges are artificial transcripts that contain multiple miRNA binding sites to trap and sequester it295,296. miRNA sponges may target one specific or multiple different miRNAs297,298, for instance, to simultaneously inhibit miR-21, miR-155 and miR-221/miR-222 in tumour cells299, or they may target a whole miRNA seed family, for instance, to sequester miR-181a, miR-181b and miR-181c300. Although this strategy has shown great utility as an experimental tool301, miRNA sponges have not yet been applied in the clinic. MiRNA-masking ASOs represent an inverted approach that masks the binding site of a miRNA within the target gene302 and offers a gene-specific and safe therapeutic strategy in cases where seed-family members have dual effects. Tiny 8–10 nt LNAs may also be used to mask seed sequences303. The 16 nt oligonucleotide-mediated masking of miR-16 binding sites in TYRP1, the mRNA of which acts as a miRNA sponge via three non-canonical miR-16 binding sites in its 3′ UTR, resulted in the restoration of the tumour-suppressive function of miR-16 in melanoma cells304. Despite these promising developments, miRNA-masking ASOs have not yet been used in the clinic. LncRNA therapeutics have only become the focus of investigations in the past decade and no lncRNA-targeting therapeutics have entered clinical development so far. LncRNAs are being actively explored as biomarkers, supporting their prevalent link with diseases (for example, preeclampsia, NCT03903393; lung cancer, NCT03806619; acute ischaemic stroke, NCT04175691). In the future, lncRNAs are expected to broaden the amount of RNA interference (RNAi) and CRISPR targets, and specific lncRNA types such as circular RNAs or natural antisense transcripts represent entirely new therapeutic approaches.
located to liver, bone marrow and spleen. Pharmacodynamic analysis in phase I trial patients showed downregulation of miR-34a target genes in white blood cells and increased levels of miR-34a in tumour tissue. Despite this, it remains unclear whether the three patients who responded to MRX34 therapy (4% response rate) experienced gene silencing-mediated effects or immune-mediated antitumour activity.

Of note, the prominent immunotherapy target programmed cell death 1 ligand 1 (PDL1) is a known target of miR-34a and could be involved in the responses observed. As the natural sequence of miR-34a contains GU-rich sequences (5'-UGGCAUGUCUAAUGGUGUGU-3') and the precise chemical formulation of MRX34 is elusive, TLR stimulation by this therapeutic cannot be excluded and requires further investigation. On the other hand, MesomiR-1 (miR-16 mimic) administration showed a favourable safety profile in a phase I clinical trial, with minor immune reactions attributed to the carrier. Third-generation modifications such as phosphoramide morpholino oligomers (PMOs) can further reduce immunogenicity by neutralizing the charge of small RNA therapeutics, thereby prohibiting their interaction with proteins, including TLRs. This technology is used in eteplirsen, a 30 nt ASO approved for treatment of Duchenne muscular dystrophy, which induces the skipping of exon 51 of dystrophin mRNA.

In summary, the discussion above indicates that although there have been some advancements addressing the issue of
| Therapeutic | Type | Modification and delivery | Route of administration | Target organ | Disease | Target gene and pathway | Phase Identifier |
|-------------|------|---------------------------|-------------------------|--------------|---------|------------------------|------------------|
| rAAV5-miHTT (AMT-130) | Pri-miR-451 backbone | Adeno-associated viral vector (AAV5) | Intrastriatal | Brain | Huntington disease | Huntingtin (HTT) mRNA | I/II NCT04120493 |
| WVE-120102 | ASO (allele-selective) | Stereopure ASO | Intrathecal | Brain | Huntington disease | U-variant of SNP rs362331 (SNP2) in HTT mRNA | I/II NCT02225846, NCT04617860 |
| RG-125 (AZD4076) | Anti-miR-103/107 | GalNAc-conjugated antagomiR | Subcutaneous | Liver | Type II diabetes, nonalcoholic fatty liver disease | miR-103/107 | I/II NCT02612662, NCT02826525 |
| Remlarsen (MRG-201) | miR-29 mimic | Cholesterol conjugated | Intradermal | Skin | Keloid (pathological fibrosis) | miR-29 targetome | II NCT02603224, NCT03601052 |
| siG12D-LODER siRNA | Biodegradable polymeric matrix (PLGA) | Intratumoral | Tumour | Advanced pancreatic cancer | G12D-mutated KRAS mRNA | II NCT01188785, NCT01676259 |
| Prexigebersen (BP1001-A) | ASO | Liposomal | Intravenous | Blood and/or immune cells | Acute myeloid leukaemia, chronic myeloid leukaemia | GRB2 mRNA | II NCT01159028, NCT04196257, NCT02781883 |
| Olpasiran (AMG 890, ARO-LPA) | siRNA | GalNAc conjugated | Subcutaneous | Liver | Cardiovascular disease | Apolipoprotein A (LPA) mRNA | II NCT03626662, NCT04270760 |
| Vupanorsen (AKCEA-ANGPTL3-LRx) | ASO | GalNAc conjugated | Subcutaneous | Liver | Dyslipidaemias, hyperlipidaemias, hyperlipoprotein-aeimias | Angiopoietin-like 3 (ANGPTL3) mRNA | II NCT04459767, NCT03371355, NCT04516291 |
| Miravirsen (SPC3649) | Anti-miR-122 | PS-β-d-oxy-LNA gapmer ODN | Subcutaneous | Liver | Hepatitis C virus infection | miR-122 | II NCT01646489, NCT01727934, NCT01872936, NCT01200420 |
| Donidalorsen (IONIS-PKK-LRx, ISIS 721744) | ASO | GalNAc-conjugated PS-2′-MOE ODN | Subcutaneous | Liver | Hereditary angio-oedema, COVID-19 | Prekallikrein (PKK) mRNA | II NCT02263507, NCT04030598, NCT04307381, NCT04549922 |
| BMT 101 (cp-asirRNA) | Cell-penetrating asymmetrical siRNA | Carrier-free | Intradermal | Skin | Hypertrophic scar | Connective tissue growth factor (CTGF) mRNA | II NCT03133130, NCT04012099 |
| Danvatisen (IONIS-STAT3-2.5Rx, AZD9150) | ASO | GalNAc conjugated | Intravenous | Tumour | Metastatic NSCLC, resectable early-stage NSCLC, pancreatic cancer, mismatch repair-deficient colorectal cancer | STAT3 mRNA | II NCT03819465, NCT03794544, NCT02983578 |
| Bamosiran (SYL040012) | siRNA | Carrier-free | Topical | Eye | Ocular hypertension, glaucoma | β-Adrenergic receptor 2 (ADRβ2) mRNA | II NCT00990743, NCT01227291, NCT01739244, NCT02250612 |
| Cemdisiran (ALN-CC5) | siRNA | GalNAc conjugated | Subcutaneous | Blood | Paroxysmal nocturnal haemoglobinuria, IgA nephropathy, Berger disease, glomerulonephritis | Complement 5 mRNA | II NCT04601844, NCT02352493, NCT03841448, NCT03999840 |
| Apatortsen (OGX-427) | ASO | 2′-O-MOE-PTO gapmer | Intravenous | Tumour | Squamous cell lung cancer, non-squamous NSCLC, urological neoplasms, metastatic bladder cancer, urinary tract neoplasms, castration-resistant prostate cancer | HSP27 mRNA | II NCT01120470, NCT01454089, NCT01829113, NCT02423590 |
immunogenicity of RNA therapeutics, the hurdle is not yet overcome and there is still much to explore. Potential solutions to help address this issue will be discussed below.

**Potential solutions**

*Expanding immune-related adverse reaction screening with TLR interaction methods.* Not all miRNAs and miRNA therapeutics similarly induce immunogenicity, and the apparent difficulty of predicting such responses prompts the use of broader screening methods in preclinical studies.

As immune responses may differ between animal models and humans, such screening methods should employ various cell types of human origin. Primary cells should be preferred as cell lines may have impaired response pathways. The use of co-culture and organoid systems as well as patient-derived xenograft models could furthermore grant a better assessment of systemic responses. In the clinical assessment of kinase inhibitors, screenings for cytokine production or kinase activity are already standard practice. Such practice should also be adapted for RNA therapeutics and include in vivo immune screening, the assessment of large panels of immune cell markers and testing of direct TLR stimulation. A comprehensive database can be developed for the systematic collection of examples of miRNAs that target TLRs (out of more than 3,000 human miRNAs already known), or of exact immune adverse reactions and gravity of symptoms for each tested RNA-based therapeutic. This will support large data analyses and allow selection of the RNA therapeutics with the smallest potential immunogenicity before the initiation of clinical studies.

### Table 2 (cont.)  RNA therapeutics in phase II or III clinical development

| Therapeutic | Type | Modification and delivery | Route of administration | Target organ | Disease | Target gene and pathway | Phase | Identifier |
|-------------|------|---------------------------|-------------------------|--------------|---------|------------------------|-------|------------|
| Sepofarsen (QR-110) | ASO | Chemically modified | Intravitreal | Eye | Leber congenital amaurosis type 10 (LCA10), blindness, LCA, vision disorders, sensation disorders, neurological manifestations, eye diseases, hereditary or congenital eye diseases | c.2991+1655A>G-mutated CEP290, pre-mRNA splicing | II/III | NCT03140969, NCT03913143, NCT03913130 |
| Alicaforsen (ISIS 2302) | ASO | Phosphorothioate-modified | Oral | Intestine | Crohn's disease | ICAM1 mRNA | III | NCT03473626, NCT00063830, NCT00063414, NCT00048113, NCT02525523 |
| Tivanisiran (SYL1001) | siRNA | Carrier-free | Topical | Eye | Dry eye disease | Transient receptor potential cation channel subfamily V member 1 (TRPV1) | III | NCT01438281, NCT01776658, NCT02455999, NCT03108664 |
| AKCEA-TTR-LRx | ASO | GalNAc conjugated | Subcutaneous | Liver | Hereditary transthyretin-mediated amyloid polyneuropathy | Transthyretin (TTR) mRNA | III | NCT04302064; NCT03728634; NCT04136184; NCT04136171 |
| Tominersen (RO7234292, HHT ASO, IONIS-HTRRx, ISIS-443139, ISIS-HTRRx, RG 6042) | ASO (allele-nonselective) | PS-2′-MOE gapmer | Intrathecal | Brain | Huntington disease | HTT mRNA | III | NCT02519036, NCT04000594, NCT03342053, NCT03761849, NCT03842969 |
| Pelacarsen (AKCEA-APO(a)-LRx, TQJ230) | siRNA | GalNAc conjugated | Subcutaneous | Liver | Hyperlipoproteinaemia | Apolipoprotein A mRNA | III | NCT03070782, NCT03070782, NCT04023552 |
| Nedosiran (DCR-PHXC) | siRNA | GalNAc conjugated | Subcutaneous | Liver | Primary hyperoxaluria type 1 and primary hyperoxaluria type 2, kidney diseases, urological diseases | Lactate dehydrogenase A enzyme (LDHA) mRNA | III | NCT03392896, NCT04555486, NCT04580420, NCT03847909, NCT04042402 |

ASO, antisense oligonucleotide; GalNAc, N-acetylgalactosamine; LNA, locked nucleic acid; LODER, local drug eluter; NSCLC, non-small cell lung cancer; siRNA, small interfering RNA; SNP, single nucleotide polymorphism.
Using ‘tiny’ antisense RNAs. As efficient activation of TLRs requires a length of at least 21 nucleotides for ssRNA\textsuperscript{[92,93]}, the design of smaller RNA therapeutics could circumvent the issue. Locked nucleic acid (LNA)-modified antimiRs with a short sequence of 7–8 nucleotides, termed ‘tiny’ LNAs, target the 5′-seed region of miRNAs and can enable antagonism and inhibition of an entire miRNA family sharing that seed sequence. A shorter sequence obviously increases the potential for off-target effects, although targeting of two miR-122 family members using tiny LNAs showed few off-target effects in vitro two miR-122 family members using tiny LNAs showed few off-target effects in vitro with no measurable effects on miRNAs that contained the complementary sequence\textsuperscript{[99]}. Tiny LNAs targeting the miR-17/miR-18/miR-19 (REF.\textsuperscript{[64]}), miR-122 and let-7 (REF.\textsuperscript{[63]}) families successfully downregulated multiple family members simultaneously. Intravenous injection of tiny LNAs in murine cancer models showed antitumour effects after treatment with anti-miR-21 in breast cancer\textsuperscript{[62]} and anti-miR-17 and anti-miR-19 in medulloblastoma\textsuperscript{[66]}. In addition, miR-34-targeting tiny LNA injection in a murine cardiac stress model showed promising results, improving cardiac parameters, an effect that could not be achieved by inhibition of a single miR-34 family member\textsuperscript{[65]}. Despite these successes in preclinical studies, tiny LNAs have not yet been clinically assessed. The successful use of shorter RNA therapeutics will depend on the detailed analyses of the correlation between the lengths of the RNA therapeutic, the in vivo efficacy and off-target binding to other RNAs or DNA by perfect or imperfect complementarity to exclude any unexpected and/or undesired effects. LNA-antimiRs that have been in clinical development (miravirsen, MRG-110 and cobomarsen (terminated, see Table 3)) are small sequences with a length of 14–16 nt, which may assist in escaping TLR recognition, but are not considered ‘tiny’ LNAs.

Applying metronomnomic miRNA therapy. By analogy with metronomic chemotherapy\textsuperscript{[71]}, we define metronomic RNA therapy as regular and frequent administration of limited drug doses over a prolonged period, to achieve a low, but active, dose range without inducing excessive toxicity or immunogenicity. The addition of metronomic cyclophosphamide to the combination trastuzumab plus pertuzumab showed markedly longer progression-free survival for older patients with metastatic HER2-positive breast cancer\textsuperscript{[72,73]}. Continuous administration of RNA therapeutics over months, instead of daily administration in cycles of weeks interrupted by periods of no administration, could achieve similar results.

To safely perform this, extensive pharmacodynamic and pharmacokinetic studies are necessary to grant a precise understanding of delivery systems as well as the circulation half-life of the chemically diverse subsets of RNA therapeutics. siG12D-LODER (local drug elutor) is a biodegradable polymeric matrix implanted into the pancreas to deliver a siRNA targeting G12D-mutated KRAS mRNA over a period of 12 weeks\textsuperscript{[69]}. A currently ongoing phase II clinical trial (NCT01676259) will show whether siG12D-LODER together with chemotherapy improves the survival of patients with unresectable advanced pancreatic cancer. For the clinical application of metronomic RNA therapy, such locally implanted delivery devices that release their therapeutics over a long period of time could be appealing if drug release from the device is stable. Furthermore, such low-dose RNA therapy may grant ‘second chances’ for therapeutics that showed immune stimulation such as the miRNA mimic MRX34. Indeed, the miR-16 mimic MesomiR-1 was reported to be dosed approximately 1,000-fold lower compared with MRX34 with no adverse immune effects observed in its phase I clinical trial\textsuperscript{[61]}. Metronomic RNA therapy is furthermore well suited as a combinatorial therapy, for instance, the combination of chemotherapy and miRNA metronomic therapeutics for elderly and/or frail patients.

Combinatorial RNA therapeutics. Combination therapies can also be used to reduce the required dose of RNA therapeutics and thus their immunogenicity, for example, if used in synergy with chemotherapy, radiotherapy, small-molecule-based or immunotherapies\textsuperscript{[72,73]}. Moreover, combinatorial RNA therapeutics could induce a dual effect, as ncRNAs are involved in resistance to commonly used therapeutics\textsuperscript{[74]}. A large-scale screening revealed that individual siRNAs, for instance, those that target genes involved in mitotic spindle assembly, could increase the sensitivity of lung cancer cell lines to paclitaxel, allowing a 1,000-fold reduction in the required dosage of the drug\textsuperscript{[75]}. Natural miRNAs can play a similar role, as shown for miR-155 in lung cancer, which caused significant resistance to chemotherapy via a negative feedback mechanism with TP53. The application of anti-miR-155 therapy resensitized lung cancer cells to chemotherapy both in vitro and in vivo\textsuperscript{[76]}. Similarly, miR-34a mimics sensitized lung cancer cells to erlotinib, an EGFR-specific tyrosine kinase inhibitor\textsuperscript{[77]} and radiation\textsuperscript{[78]}. In addition, many lncRNAs have also been linked to drug resistance in cancer\textsuperscript{[79]}, thus broadly enlarging the pool of ncRNA targets that may be exploited in the future. Furthermore, combinatorial therapeutics are being investigated for non-cancerous diseases. A recent phase II clinical trial reported the use of a single injection of the anti-miR-122 therapeutic RG-101 in combination with viral protein inhibitor GSK2878175, administered orally for 12 weeks as a single-visit curative regimen for chronic hepatitis C (EudraCT 2015-004702-42). This study reported high cure rates with an oral GSK2878175 administration scheme of longer than 9 weeks\textsuperscript{[80]}; however, the development of RG-101 was halted owing to adverse effects observed in a different clinical trial (discussed below)\textsuperscript{[81]}. Long circulating NPs with a pH-regulated drug release mechanism have been developed to deliver siRNA–mitoxantrone drug complexes directly into the tumour tissue. The delivery of siRNA targeting PLK1 (Polo-like kinase, a mitotic cell cycle regulator) together with a miR-200c mimic (miR-200c acts as a tumour suppressor by blocking epithelial-to-mesenchymal transition) caused tumour regression in a murine breast cancer model\textsuperscript{[82]}. Furthermore, the combination of miR-520d-3p and anti-Eph2A siRNA therapeutics in liposomal NPs was tested in preclinical models. EphA2 is an ovarian cancer oncoprotein, the downregulation of which suppresses tumour growth and tumour cell migration and invasion\textsuperscript{[83]}. The synergistic effect of anti-Eph2A siRNA and miR-520d-3p therapy was likely caused by the concordant targeting of related receptor EphB2 by miR-520-3p\textsuperscript{[84]}. Another interesting approach is the combination of multiple RNA therapeutics to target not only multiple genes but also multiple cell types. Proof of this concept has been provided by use of siRNAs that target vascular...
endothelial growth factor (VEGF) and placental growth factor (PIGF) delivered by multi-functionalized NPs to both breast cancer cells and tumour-associated macrophages. This approach caused a synergistic effect, addressing the role of VEGF and PIGF in promoting tumour proliferation and immunosuppression by creating a pro-oncogenic nature in the tumour microenvironment.

Another interesting approach for combinatorial RNA therapy is the use of certain molecules and compounds for gene therapy. This includes the use of galactosaminic acid (GalNAc) for targeting specific cell types, cell-penetrating peptides for nuclear entry, and metal nanoparticles for precise delivery. Other methods include Oligonucleotide conjugation with antibodies, lipids, polymers, viral vectors, bacterial minicells, and bacteriophages, as well as exosomes for cargo delivery.
MicroRNA biogenesis pathway and ways to interfere therapeutically. MicroRNA (miRNA) biogenesis is a multistep process (see blue boxes) consisting of transcription of a pri-miRNA by RNA polymerase II or III, its nuclear processing into a pre-miRNA by Drosha and Dicer, nuclear export of the pre-miRNA by exportin 5, cytoplasmic processing by Dicer and TRBP into a mature miRNA duplex and its helicase-mediated unwinding. The passenger strand is degraded, and the mature miRNA strand is integrated into the RNA-induced silencing complex (RISC) to mediate either translational repression or miRNA degradation depending on the extent of complementarity to the miRNA target. Translational repression is mediated through effects on translation initiation, elongation and termination as well as co-translational degradation. miRNA degradation is mediated through mechanisms resulting first in mRNA deadenylation (step 1), followed by de-capping (step 2) and concluded by co-translational degradation (step 3). Ways to interfere with the endogenous miRNA pathway (see red boxes) include inhibition of biogenesis at the nuclear or cytoplasmic level, miRNA replacement therapy and functional inhibition of the mature miRNA or the interaction with its target mRNA. As oligonucleotides are not readily taken up into cells, commonly used delivery methods are shown and include conjugation to various chemical or biological entities as well as delivery within lipid particles, polymers and viral or bacterial vector systems.

Small-molecule inhibitors of miRNAs. Small molecules have important advantages over the use of oligonucleotides, that is, good solubility, bioavailability, and metabolic stability. They may act on miRNAs at the transcriptional or post-transcriptional level or may influence miRNA processing. The main methods to screen for small-molecule inhibitors of miRNAs (SMIRs) are either computer-assisted design through sequence screening or high-throughput screening of (existing) compounds for interaction with miRNAs or miRNA precursors or effects on miRNA pathways. Computational models to predict the miRNA–SMIR association may employ miRNA structure-based models (that mainly work on the miRNA precursor 3D structure to predict the appropriate small-molecule interactors), and gene expression profile-based models (developed on the basis of the expression profile of genes targeted by small molecules and/or miRNAs or on the basis of previously known miRNA–SMIR interactions), which are sometimes combined with the structural similarity of small molecules and miRNAs. Moreover, databases that record the relationships between small molecules and miRNAs, such as SM2mir, Informa 2.0, mTD, NRDTD and ChemiRs offer a reasonably comprehensive repository of data regarding the influence of small molecules on miRNA expression. High-throughput screening of large compound libraries use either the interactions of specific miRNAs with their biogenesis machinery or effects on miRNA expression levels as efficacy measures. For example, >100,000 compounds were assessed for interactors with let-7 binding motifs in RNA binding protein LIN28, which identified TPEN and Li71 as inhibitors of LIN28-mediated let-7 biogenesis. Screening for miRNA–Dicer interactions revealed that 4 of 14 tested aminoglycosides (streptomycin, neomycin, tobramycin and amikacin) specifically inhibited pre-miR-27a processing, and 3 polypeptide derivatives from a 640-compound library interfered with pre-miR-372 processing. Of the latter, the most active inhibitor, a spermine–amidine conjugate, showed anti-proliferative activity in gastric cancer cells that overexpressed miR-372. Last, testing of more than 40,000 compounds for their ability to inhibit oncogenic miR-29a led to the discovery of five inhibitors of pre-miR-29a.

Functionally, SMIRs may interfere with the transcription or the nuclear or cytoplasmic processing of a specific miRNA. The first identified SMIRs interfered with miRNA transcription and include miR-21-inhibiting azobenzene-2 (REF 104) and miR-122-reducing ’mIR-122 inhibitor 2. The latter efficiently reduced the expression of liver-specific miR-122, resulting in a reduction in hepatitis C virus replication. Aza-flavanones specifically inhibited transcription of miR-4644, an inhibitor of cell death, and induced cell death in human carcinoma cells. Other SMIRs act by inhibiting the biogenesis of specific miRNAs, blocking either nuclear pri-miRNA processing by Drosha or cytoplasmic pre-miRNA processing by Dicer. Targapremir-96 is a small dimeric molecule that binds to the internal loop of miR-96 to interfere with Drosha binding. Targapremir-96 treatment in breast cancer cells triggered apoptosis without affecting healthy cells and this effect was reversed by overexpression of pri-miR-96 (REF 105). Owing to its prevalence as an oncomiR in many types of cancer, multiple compounds were identified that interfere with Dicer processing of pre-miR-21, including the aminoglycoside streptomycin (a tuberculosis therapeutic), a peptide possessing a binding pocket for pre-miR-21 (REF 106) and the compound AC1MMYR2, which showed potent antitumour effects in murine glioma and breast cancer models. BzDANP binds to pre-miRNAs possessing a C bulge near the Dicer processing site, that is, miR-29a and miR-136. The binding of BzDANP to miR-136 caused formation of a ternary complex together with Dicer, slowing the kinetics of pre-miR-136 processing and consequently lowering mature miR-136 levels. Targapremir-210 binds close to the Dicer processing site on pre-miR-210. MiR-210 is a major regulator of the hypoxic response (via a positive feedback loop with hypoxia-inducible factor 1α) and thereby influences breast cancer metastasis, and accordingly, treatment of triple-negative breast cancer cells with targapremir-210 triggered apoptosis under hypoxic conditions and inhibited tumorigenesis in a murine xenograft model. Other examples include the multi-tyrosine kinase inhibitor linifanib, which inhibits cytoplasmic miR-10b maturation through an unknown mechanism, and an aminoglycoside–nucleobase conjugate that inhibited Dicer processing of multiple oncogenic miRNAs including miR-372, miR-373, miR-17 and miR-21, but possibly also other miRNAs. A more tailorable approach can be developed using a miRNA binding molecule linked to an effector domain. One such approach links a short 2′-5′ poly(A) oligonucleotide to a miRNA binding molecule, causing selective recruitment of RNase L to degrade the RNA structure. This principle has been shown to work in vitro, where selective degradation of oncogenic pri-miR-96 induced apoptosis in triple-negative breast cancer cells, and in vivo, where selective degradation of pre-miR-21 reduced the occurrence of metastases in a murine breast cancer model. A similar approach consists of linkage of a miRNA binding molecule to a weak Dicer inhibitor, a method termed ‘proximity-enabled Dicer inhibition.’
| Therapeutic     | Type     | Modification and delivery | Route of administration | Target organ | Disease                                      | Target gene and/or pathway | Reason for termination                                      | Ref. |
|----------------|----------|---------------------------|--------------------------|--------------|----------------------------------------------|-----------------------------|------------------------------------------------------------|------|
| Aganirsen      | ASO      | 1st gen; PT Topical       | Eye                      | Ischaemic central retinal vein occlusion, neovascular glaucoma | Insulin receptor substrate 1 (IRS1) mRNA | Formulation stability issues                                |      |
| Cobomarsen     | AntimiR  | 3rd gen; LNA Subcutaneous or intravenous | Blood or lymphoid organs | Various lymphomas | miR-155                                      | Non-safety or efficacy-related company decision              |      |
| PRO-040201     | siRNA    | Liposomal (stable nucleic acid lipid particle) | Intravenous | Liver | Hypercholesterolaemia | Apolipoprotein B (APOB) mRNA | Potential for immune stimulation (flu-like symptoms) |      |
| AGN 211745     | siRNA    | Chemical composition unclear; carrier-free | Intravitreal | Eye | Age-related macular degeneration, choroidal neovascularization | Vascular endothelial growth factor receptor 1 (VEGFR1) mRNA | Lack of clinical efficacy, TLR3 stimulation (sequence-independent TLR3-mediated therapeutic effect) | 31   |
| RG-101         | AntimiR  | GalNAc conjugated Subcutaneous | Liver | Hepatitis C infection | miR-122                                    | High levels of bilirubin in the blood                       |      |
| MRX34          | miRNA mimic | Liposomal Intravenous or intratumour | Tumour | Primary liver cancer, advanced or metastatic cancer with or without liver involvement, haematological malignancies | miR-34a targetome | Immune-related adverse events | 24   |
| Oblimersen sodium (G3139, Genasense) | ASO | 1st gen; PT Subcutaneous | Tumour | Various malignancies | BCL2 mRNA | Lack of clinical efficacy, insufficient delivery, primary end points not met | 21   |
| Suvodirsen     | ASO      | 1st gen; PT, stereopure Intravenous | Muscle | Duchenne muscular dystrophy | Dystrophin (DMD) pre-mRNA splicing (exon 51 skipping) | Lack of clinical efficacy, failure to increase dystrophin levels |      |
| DCR-MYC        | siRNA    | Liposomal Intravenous | Tumour | Advanced solid tumours, multiple myeloma, lymphoma | MYC mRNA | Lack of clinical efficacy despite MYC reduction |      |
| DCR-PH1        | siRNA    | Liposomal Intravenous | Liver | Primary hyperoxaluria type 1 (PH1) | Lactate dehydrogenase A (LDHA) mRNA | Development shifted to GalNAc-conjugated variant (DCR-PHXC) |      |
| Custirsenen    | ASO      | 2nd gen; 2’-MOE gapmer Intravenous | Tumour | Prostate cancer, breast cancer | Clusterin (CLU) mRNA | Lack of clinical efficacy, primary end points in phase III trials not met |      |
| Bevasiranib    | siRNA    | 1st gen; PT Intravitreal | Eye | Age-related macular degeneration, diabetic macular oedema | Vascular endothelial growth factor (VEGF) mRNA | Lack of clinical efficacy, TLR3 stimulation (sequence-independent TLR3-mediated therapeutic effect) | 31   |
| AEG351556      | ASO      | Mixed backbone oligonucleotides Intravenous | Tumour | Various malignancies | X-linked inhibitor of apoptosis (XIAP) mRNA | Lack of clinical efficacy, increased incidence of chemotherapy-induced peripheral neuropathy | 255  |
| ISIS 329993    | ASO      | 2nd gen; 2’-MOE Subcutaneous or intraperitoneal | Heart or joints | Paroxysmal atrial fibrillation, rheumatoid arthritis | C-reactive protein (CRP) mRNA | Lack of clinical efficacy despite CRP mRNA reduction | 269  |
This system was used to successfully inhibit miR-21 biogenesis in vitro, and the use of a photocleavable linker enabled inactivation of the reaction by light\textsuperscript{20,21}. Other small-molecule inhibitors can be envisioned that target the hairpin structure, which can be very similar between two or more oncogenic miRNAs such as miR-10b and miR-21, the concomitant downregulation of which reduces the proliferation of glioma cells\textsuperscript{122}. Such an approach could synergize the killing of malignant cells and in addition reduce potential resistance, as the two miRNAs induce cell death through independent targets. Although there are so far no reports of small molecules targeting miRNA hairpin structures, the hairpin structure of a promoter G-quadruplex structure in hTERT has recently been targeted successfully using the small molecule RG1603 (REF\textsuperscript{123}), implying the feasibility of this approach. Targeting of the mature miRNA or its seed sequence could represent another tactic, whereby the latter would enable inhibition of an entire miRNA family, akin to what was achieved using tiny LNA antisense technology.

Although SMIRs have been reported as efficient miRNA inhibitors, several challenges remain regarding specificity, optimal potency, delivery and lack of toxicity that still hinder their clinical application, and there are currently no clinical trials involving SMIRs.

**Targeting ncRNAs with interaction element blockers and structural element lockers.** A new type of RNA therapy can be envisaged based on the targeting of interactor elements (IEs), that is, docking sites on ncRNAs for multiple types of molecule (DNA, RNA, proteins and lipids) and/or targeting structural elements (SEs), which directly affect the conformation of a ncRNA and indirectly the functional interactions with interactor molecules (FIG. 2b). Consequently, such new therapeutics will influence signalling pathways and disease phenotypes. Small molecules are obvious candidates for such developments owing to their stability, lower toxicity and pharmacokinetic and dynamic properties. IE blockers (IEBs) can also be envisioned as antisense oligonucleotides able to hide the IE from the interaction with its effector, while an oligonucleotide SE locker (SEL) could disrupt the SE sequence to affect secondary and tertiary structures of ncRNAs.

The small molecule NP-C86 has been shown to inhibit the interaction of the GAS5 IncRNA with UPF1, a protein involved in nonsense-mediated decay and degradation of GAS5. Blocking this interaction resulted in increased stability of GAS5, which in turn increased its interaction with the insulin receptor and caused an upregulation of glucose intake in adipocytes\textsuperscript{124}. NP-C86 is thus an IEB and may find future applications in the treatment of patients with diabetes. MALAT1 is strongly associated with cancer metastasis and contains a triple helix structure at its 3′-end that protects the IncRNA from exonuclease degradation and assists its nuclear accumulation. Multiple studies have attempted to identify SEs that target this stabilizing triple structure. Fluorescence-based screening of a diphenylfurran (DPF)-scaffold-based library identified a DPF-based small molecule that specifically targets the triple helix structural element in MALAT1 (REF\textsuperscript{125}). However, it remains to be elucidated whether the binding of this molecule results in efficient MALAT1 destabilization. Similarly, another high-throughput screening identified two small molecules that target the MALAT1 triple helix, but not other similar structures present for instance in NEAT1. The compounds significantly reduced levels of MALAT1 as well as downstream genes and attenuated the branching of mammary gland organoids\textsuperscript{126}. However, the development of MALAT1-targeting small molecules should be addressed with caution as a dual function has been described for this IncRNA\textsuperscript{127}. Accordingly, these molecules have not yet entered clinical testing.

**The hurdle of specificity.** The quality of an RNA therapeutic is determined by the strength of its on-target specificity as well as the absence of off-target and undesired on-target effects. Many developments in RNAi therapeutic design have already improved on-target specificity and decreased unwanted off-target effects. Second-generation chemical modifications have significantly improved the potency of antisense therapeutics\textsuperscript{128}, and third-generation LNA modifications show the highest potency\textsuperscript{29}. Studies on siRNA specificity have shown that off-target genes with three or four base mismatches and additional G-U mismatches can be efficiently downregulated through translational repression\textsuperscript{130}, similar to the effect of miRNA binding with mismatches (FIG. 1). Sequence-mediated off-targeting was mainly caused by complementarity to the first eight nucleotides at the 5′-end of the siRNA, again mirroring the endogenous function of miRNA seed sequences\textsuperscript{131,132}. Placement of a 2′-O-methyl ribosyl substitution at position 2 of the guide strand could abrogate this seed effect\textsuperscript{129}. Other off-target effects are caused by loading of

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**Table 3 (cont.)** | RNA therapeutics for which clinical development was halted

| Therapeutic | Type | Modification and delivery | Route of administration | Target organ | Disease | Target gene and/or pathway | Reason for termination | Ref. |
|-------------|------|--------------------------|-------------------------|-------------|--------|---------------------------|----------------------|------|
| PF-4523655 (PF-655) | siRNA | 2nd gen; liposomal | Intravitreal | Eye | Age-related macular degeneration, diabetic macular oedema | DNA damage-inducible transcript 4 (DDIT4) mRNA | Lack of clinical efficacy, no improvement on current standard of care | 270 |
| ISIS 104838 | ASO | 2nd gen; 2′-MOE gapmer | Oral | Joints | Rheumatoid arthritis | TNF mRNA | Cost and competition-related company decision | 271 |
| ISIS 5132 (CGP 69846 A) | ASO | 1st gen; PT | Intravenous | Tumour | Breast cancer, ovarian cancer | Raf mRNA | Lack of clinical efficacy | 272 |
| Aprinocarsen (ISIS 3521, LY900003) | ASO | 1st gen; PT | Intravenous | Tumour | Non-small cell lung cancer | Protein kinase Ca mRNA | Lack of clinical efficacy | 273 |

AntimiR, anti-microRNA; ASO, antisense oligonucleotide; GalNac, N-acetylgalactosamine; gen, generation; LNA, locked nucleic acid; PT, phosphothiorate; siRNA, small interfering RNA; TLR3, Toll-like receptor 3. aSee Related links.
Positive-strand RNA viruses resemble endogenous mRNAs and can consequently be directly bound and regulated by microRNAs (miRNAs). This can influence viral replication positively or negatively via various mechanisms such as RNA stabilization, and thus differs from miRNA interaction with endogenous mRNAs, which generally results in downregulation. In addition, viral infection may sequester endogenous miRNAs or change their expression, which can similarly lead to pro- or antiviral effects (reviewed in references 305,306). As a result, both miRNA replacement and miRNA inhibitory therapies can be applicable for the treatment of viral infection.

Several examples of pro- and antiviral miRNA effects are shown in the figure. Pro-viral effects include those mediated by liver-specific miR-122 [ref. 319], which binds to three sites in the 3′ and 5′ UTRs of hepatitis C virus (HCV) RNA319–321, and miR-17 binding to the 3′ UTR of bovine viral diarrhoea virus (BVDV) RNA322, both causing stabilization and increased translation. Although the interaction of miR-142-3p with Eastern equine encephalitis virus (EEEV) inhibits viral translation, the haematopoiesis-specific expression of miR-142-3p turns this into a pro-viral effect as it assists viral escape from immune recognition331. Antiviral effects include inhibition of porcine epidemic diarrhoea virus (PEDV) translation and enterovirus 71 (Ev71) replication by binding of miR-221-5p332 and let-7a333, respectively, and the pro-inflammatory effect of miR-9 sequestration by human coronavirus OC43, resulting in derepression of nuclear factor-κB (NF-κB)334.

Fomivirsen was the first FDA-approved antisense oligonucleotide (ASO) targeting the IE-2 mRNA of cytomegalovirus (CMV)335, which was used to treat CMV-induced rhinitis before the emergence of antiretroviral therapies336. Two miR-122 antiimiRs (miravirsen and RG-101) successfully reduced hepatitis C virus (HCV) RNA levels in clinical trials, resulting in transiently undetectable levels of HCV RNA in approximately 50% of test subjects for up to 14 (miravirsen) or 76 (RG-101) weeks after dosing305,319,320. Of note is that stabilizing mutations could render HCV independent of miR-122-mediated stabilization. Disruption of the miR-122 binding motifs on HCV RNA by point mutations caused viral resistance to anti-miR-122 therapy in vitro312,313. Although serial passage in the presence of miravirsen showed a HCV 5′ UTR C3U substitution, but no mutations in miR-122 binding sites S1 to S3314. A pressing question at hand is whether RNA interference (RNAi) or miRNA-based therapeutics can be used to tackle the currently surging serious acute respiratory syndrome coronavirus 2 (SARS-CoV-2) infections causing coronavirus disease 2019 (COVID-19; reviewed in reference 312).

Alternative approaches to treat cytokine release syndrome in severe COVID-19 with stem cell-derived, miRNA-loaded exosomes have also been suggested324,325. A plethora of in silico studies indicate that numerous human miRNAs can bind to SARS-CoV-2 RNA within its UTRs, all structural and several nonstructural protein-coding regions326,327. However, the functional implications of this remain largely unstudied, although a general sequestration of endogenous miRNAs by SARS-CoV-2 RNA has been suggested328. Of note, comparison of strength of miRNA binding to different coronavirus UTRs showed weaker binding in the novel SARS-CoV-2 compared with older, less-pathogenic variants, suggesting evolutionary evasion from inhibition by endogenous miRNAs327.

Furthermore, the virus itself is thought to encode multiple miRNAs328,334–336, the endogenous miRNA interactors of which have so far only been explored in prediction models. Moreover, endogenous miRNAs may target proteins required for viral entry such as the ACE2 receptor (miR-200c337) or the neural-entry cofactor neuropilin-1 (miR-24)338. Last, SARS-CoV-2 strongly influences the noncoding transcriptome of infected cells, affecting the expression of miRNA339 as well as that of lncRNA340,341. The analysis of SARS-CoV-2-infected bronchoalveolar stem cells identified multiple deregulated miRNAs capable of interacting with both host and viral RNAs to create a pro-viral environment and escape immune recognition342. An interesting observation regarding lncRNAs is the upregulation of MALAT1 [ref. 343], which was previously shown to promote inflammatory injury in lung tissue via negative regulation of IL-8 [ref. 344], and of MIR3142HG344,345, which is also induced by pro-inflammatory IL-1β and positively regulates secretion of IL-8 and CCL2 (neutrophil/monocyte chemoattractant) in lung fibroblasts345. The complex interactions between host and viral noncoding genes in SARS-CoV-2 infection is thus only starting to be explored, but the findings so far hint at diverse opportunities to target COVID-19 disease with RNA-based therapeutics.

**Box 3 | Targeting RNA viruses with RNA-based therapeutics**

The passenger instead of the guide strand into the RNA-induced silencing complex (RISC). Multiple approaches have been suggested to improve guide-strand RISC loading, including the incorporation of LNA s at the 5′-end of the passenger strand331,332, 2′-F-O-Me-phosphorodithioate modifications333, trimming334 or fragmentation335 of the passenger strand, its 5′-biotinylation336, and creation of so called Dicer-substrate siRNAs (longer siRNAs with asymmetrical termini and containing DNA nucleotides)337. Yet other off-target effects may stem from interactions of RNA therapeutics with proteins. Oblimersen (Genasense), a first-generation 18 nt phosphothiorate-modified ASO that targets BCL2 mRNA, was tested in several clinical
trials but showed limited efficacy\textsuperscript{21,140,141}. Molecular studies comparing oblimersen with BCL2-targeting siRNAs later showed several off-target effects, including upregulation of stress-response genes and downregulation of proliferation-associated genes\textsuperscript{142}, as well as downregulation of several apoptosis and glycolysis-related proteins\textsuperscript{143}. These effects were in part defined to be a class effect of phosphothiorate-modified ASOs, as the identical sequence formulated with a phosphodiester backbone did not show the same effect\textsuperscript{142,143}. Detailed molecular studies showed that phosphothiorate oligonucleotides such as oblimersen

\textbf{a} Small-molecule inhibitors of miRNA

\begin{itemize}
  \item Transcription
    \begin{itemize}
      \item Azobenzene-2
      \item \textit{\\downarrow} Pri-miR-21
      \item miR-122 inhibitor 2
      \item \textit{\\downarrow} Pri-miR-122
      \item Aza-flavanones
      \item \textit{\\downarrow} Pri-miR-4644
    \end{itemize}
  \item Nuclear processing
    \begin{itemize}
      \item Targaprimir-96
      \item \textit{\\downarrow} Pri-miR-96
      \item Drosha
      \item Internal loop
    \end{itemize}
\end{itemize}

\begin{itemize}
  \item Cytoplasmic processing
    \begin{itemize}
      \item Targapremir-210
      \item \textit{\\downarrow} Pre-miR-210
      \item Targapremir-18a
      \item \textit{\\downarrow} Pre-miR-18a
      \item BzDANP
      \item \textit{\\downarrow} Pre-miR-29a
      \item Linifanib
      \item \textit{\\downarrow} Pre-miR-10
      \item Proximity-enabled Dicer inactivation
      \item \textit{\\downarrow} Pre-miR-21
    \end{itemize}
\end{itemize}

\begin{itemize}
  \item Interaction element blocker
    \begin{itemize}
      \item IEB (NP-C86)
      \item \textit{\\downarrow} UPF1
      \item Nonsense-mediated decay
      \item GASS IncRNA
    \end{itemize}
  \item Structural element locker
    \begin{itemize}
      \item SEL
      \item \textit{\\downarrow} MALAT1 IncRNA
      \item Stabilizing secondary structure
      \item Disassembly of stabilizing secondary structure
      \item MALAT1 destabilization
    \end{itemize}
\end{itemize}

Fig. 2 | Modes of action of small-molecule inhibitors that target miRNAs and IncRNAs. \textbf{a} Small-molecule inhibitors of miRNA (SMIRs) may act at the transcriptional level or may affect the nuclear or cytoplasmic maturation steps of the microRNA (miRNA). The exact mechanism by which azobenzene-2, miR-122 inhibitor 2 and aza-flavanones inhibit the transcription of specific miRNA host genes to primary RNA transcripts (pri-miRNAs) is unknown. Targaprimir-96 binds to the internal loop of pri-miR-96 to prevent its processing by Drosha. Multiple small molecules interfere with Dicer processing, including targapremir-210, which binds to the Dicer cleavage site, and targapremir-18a, which binds to a 1 nt bulge present in three of the six miRNAs of the miR-17–92 cluster. BzDANP similarly binds to a C bulge present in miR-29a and miR-136, causing complex formation and slowing of Dicer processing. Linifanib inhibits the processing of pre-miR-10 via an unknown mechanism. Proximity-enabled Dicer inactivation makes use of two small molecules, a miRNA binder and a weak Dicer inhibitor that is active upon proximation. Use of a photocleavable linker can grant specific Dicer inactivation that can be terminated using light. \textbf{b} The first small molecules applied to modify long noncoding RNA (IncRNA) expression levels can be classified as interaction element blockers (IEBs) and structural element lockers (SELS). NP-C86 works as an IEB for GASS, blocking its interaction with UPF1, which normally results in nonsense-mediated decay of GASS, thus increasing the stability and half-life of GASS. Multiple SELs are being developed for MALAT1, which carries a stabilizing triple helix structure at its 3’ end. The SELs are aimed at disrupting the stabilizing triple helix, consequently resulting in MALAT1 destabilization and downregulation.
could directly bind and inhibit the mitochondrial channel protein VDAC, resulting in apoptosis. In accordance with an off-target effect, oblimersen also induced apoptosis in BCL-2-negative cell lines. An unexpected side effect was also noted in the clinical trial phase IB of the anti-miR-122 therapeutic RG-101, which was terminated owing to high levels of bilirubin in the blood (see Related links). Although the precise molecular mechanism causing impaired bilirubin transport is not yet clear, hepatocyte-specific loss of miR-122 in mice affected lipid, but not bilirubin, protein or glucose metabolism, indicating the absence of an undesired on-target effect. Liver-specific deletion of Dicer to abrogate global miRNA production also did not affect bilirubin metabolism in mice, arguing against a global impact on miRNA processing. The anti-miR-122 therapeutic miraviren showed a favourable safety profile with no reports of adverse immune effects or liver toxicity.

Undesired on-targeting is also common, especially when the RNA therapeutic is administered systemically. For example, the advanced tumour therapeutic MRX34 was shown to be taken up by white blood cells, and miR-34a plays major roles in T cells and macrophages. In vitro analysis of macrophages and T cells upon incubation with a miR-34a mimic showed changes in chemokine profiles that would be detrimental to tumour lysis. Whether this caused the low response rate (4%) or was involved in the adverse immune effects observed in the clinical trial of MRX34 remains to be elucidated. Similarly, the increase in chemotherapy-induced neuropathy in patients who received the XIAP-targeting ASO AEG35156 could be due to downregulation of XIAP in oligodendrocytes, glial or neuronal cells instead of tumour cells.

Another important consideration is the dosing of the RNA therapeutic. For example, it has been shown that off-targeting is more sensitive to siRNA concentration than on-target silencing. In addition, as many RNA therapeutics make use of the endogenous RNAi machinery, coarse overdosing can cause saturation of the system and prevent the function of endogenous miRNAs. The first report of this phenomenon indicated that strong shRNA overexpression in hepatocytes caused global downregulation of miRNAs, resulting in liver toxicity and mortality in mice. A large-scale screening study confirmed that global upregulation of miRNA target genes is a general phenomenon caused by the introduction of exogenous small RNAs.

When using miRNA mimics and antimiR approaches, dosing has additional implications. On the one hand, miRNAs are known to target multiple different miRNAs. It has for example been demonstrated that one cluster of miRs (the miR-15a–16-1 cluster) alone can affect (by direct and indirect targeting) about 14% of the entire transcriptome in leukaemic cells. This is both concerning, as it could lead to unwanted silencing of genes, and advantageous, as miRNAs can affect entire signalling pathways by synchronous silencing of different genes involved in that pathway. On the other hand, each mRNA target is regulated by multiple miRNAs and may possess different miRNA binding sites in close vicinity, resulting in binding competition. Therefore, the question that arises is how much the concentration of an overexpressed or downregulated miRNA affects the targetome (that is, the full spectrum of target genes silenced by a specific miRNA). Experimental approaches designed to identify the plethora of targets of a specific miRNA introduce an arbitrary amount of miRNA, not considering how the quality and quantity of targeted miRNAs could be affected. Let us consider the example of the oncosgenic miR-17–92 cluster, in which miRNAs may be upregulated 2- to 36-fold in a pathological setting (that is, in lymphoma cell lines or Burkitt lymphoma tissues). Transient transduction of miR-17–92 cluster mimics at concentrations commonly used in experimental settings caused not only a supraphysiological upregulation of up to 400-fold, but also nonspecific gene expression changes and accumulation of mutated and tailed mimic variants, whereas lower concentrations conversely did not result in specific gene knock-down. If miRNAs interact with their targets in a competitive manner, perturbation of the level of a single miRNA will affect the targeting of the mRNA by other miRNAs. In a mathematical model of such combinatorial regulation, miRNA overexpression or depletion could also lead to positive regulation of target miRNAs through changes in competitive interactions. Studies to address the effect of different miRNA concentrations on the targetome are thus warranted and seem the most logical basis for any future translational study with miRNA therapeutics. Unfortunately, dosing represents an important limitation in the field of miRNA therapeutics, and clinical studies are rarely able to determine the precise dose of an RNA therapeutic successfully delivered to the cell type of interest.

In summary, several issues persist with regard to specificity, including unexpected off-target effects and undesired on-target effects, and the relationship between dosing and specificity remains largely unexplored in the clinical setting. The following considerations may help in overcoming these issues in the future.

Potential solutions
Cell-specific miRNA modulation. One possible approach to limit the detrimental overexpression of a miRNA or antimiR in undesirable cells, is to express the RNA therapeutic using a suitable vector under the control of a specific promoter, which is overexpressed in the cells of interest. For instance, MYCN is a well-known driving oncogene in neuroblastoma. Therefore, more selective expression of a therapeutic miRNA in neuroblastoma cells could be achieved by cloning the miRNA into a vector under the promoter for MYCN. If this vector enters a non-neuroblastoma cell with little to no expression of MYCN, the expression of the miRNA will be minimal. In contrast, high expression will be achieved in MYCN-overexpressing cancer cells. In such a way, anti-miR-155 could be a suitable therapeutic in neuroblastoma, as it proved to contribute to resistance to chemotherapy through exosomal crosstalk between neuroblastoma cells and human monocytes in vitro. Such cell-specific miRNA modulation represents an alternative approach that has not been applied clinically. Dosing should be monitored closely when using this approach, as, for instance, shRNA expression from strong promoters caused neurotoxicity owing to saturation of the RNAi machinery. Conversely, expression of miRNA mimics from pri-miR-encoding plasmids resulted in miRNA expression at physiological levels and caused specific gene targeting.

Pseudo cell-specific targeting. In cases where a miRNA or IncRNA is specifically overexpressed in a diseased cell type but shows low expression in normal cell types, pseudo cell-specific targeting could be applied. This principle assumes that the downregulation of a lowly expressed transcript in normal cells will not produce strong adverse effects. This is the case for a large proportion of IncRNAs that are expressed at very low levels in normal cells, and therefore antisense therapy will have little to no effect (and consequently no toxicity) in normal cells. One such example is targeting of the human-specific IncRNAs from the category of pyknon transcripts.
These are noncoding transcripts that contain one or more very short distinct sequences (about 16 nt long) that are human and/or primate specific and work as sponges of miRNAs. Having at least 40 copies in the genome, each pyknon is quite poorly expressed in normal cells. Therefore, in the case of overexpressed pyknon transcripts in malignant cells, a nanodelivery system containing siRNA specifically targeting the transcript was used and achieved therapeutic efficacy in mouse models of colon cancer\(^{164,165}\). The clinical application of this approach requires careful preclinical investigation of the exact tissue expression patterns of the ncRNA, especially for such transcripts that are non-conserved and primate and/or human specific, as are many lncRNAs and some miRNAs\(^{166}\).

**Targeting precursor miRNAs.** Molecules that are able to interfere with miRNA genesis and function are a potent therapeutic strategy. Peptide nucleic acid (PNA) oligomers that target pre-miRNA are exploited to inhibit miRNA maturation. PNAs bind to their RNA targets with high specificity as even a single mismatch strongly influences the association constant of PNA–RNA complexes\(^\). The latter may be attributed to the dilution of these generally low transcribed but highly conserved, stem–loop structured (pre-miRNA), thereby disrupting its recognition by Dicer, which depends on the integrity of the stem–loop structure. Using PNAs, miR-210 maturation was successfully inhibited in K562 cells, blocking mithramycin-induced erythroid differentiation, and PNA-backbone modifications significantly improved cellular uptake and resistance to enzymatic degradation\(^{168-170}\). Interestingly, pre-miRNAs and their processing machinery including Dicer and Ago2 are also present in tumour-derived microvesicles\(^\), indicating that therapeutics targeting pre-miRNAs could also disrupt exosomal signalling between tumour cells and the microenvironment. SMIRs, as described above, may similarly be used to target precursor miRNAs.

**Using circular transcripts.** RNAs in circular form are much more stable than their linear counterparts. Multiple circular small RNAs have been synthesized, including dumbbell shaped (dbRNA)\(^{172-174}\), covalently closed\(^{175,176}\) and linker-circularized\(^{177,178}\) siRNAs. Interestingly, many of these circularized siRNAs were not only more stable but also mediated more efficient and prolonged RNAi effects\(^{179-181}\). dbRNA, for example, mediated stronger and longer RNAi effects at lower concentrations\(^\). A miR-34a-3p- and miR-34a-5p-containing dbRNA (db34a) inhibited angiogenesis in an in vitro model as well as in a zebrafish model\(^\). Covalent closure of only the passenger strand to prevent its RISC loading could significantly reduce off-target effects in vitro\(^\). In fact, circularization seems to overcome multiple hurdles experienced by linear siRNAs, showing less immune stimulation\(^{176,178}\), higher stability and bioavailability\(^{172,174,175,177}\), high efficacy\(^{172,173,176}\) and increased cellular uptake\(^\).

The discovery of circular lncRNAs (circRNAs) as miRNA sponges further broadens the possibilities for blocking the function of oncogenic RNAs. The first characterized circRNA was the ciRS-7 transcript, which is highly expressed in the human brain and contains more than 70 conserved miR-7 binding sites\(^\). It has been established that circRNAs are especially abundant in brain tissue\(^\), whereas they are mostly downregulated in cancer tissue\(^\). The latter may be attributed to the dilution of these generally low transcribed but highly stable transcripts in quickly proliferating tissues\(^\). CircRNA deregulation has been linked to several diseases, including cancer, diabetes and atherosclerosis (reviewed in\(^\)). Endogenous or synthetic circRNAs thus have the potential to be exploited as potent and highly stable miRNA sponge therapeutics. Foreign circRNAs were shown to cause an intron-mediated immune response through PAMP receptor RIG1 (REF\(^\)). N\(^\rightarrow\)-methyladenosine (m\(^\)A) modifications, as occur in endogenous circRNA, were able to alleviate immune stimulation\(^\). After further characterization of their immunogenic properties, synthetic circRNAs thus have great potential as miRNA sponge therapeutics. Synthetic circular miRNAs are further being explored to reconstitute protein expression and were shown to be less immunogenic (no RIG1 or TLR stimulation in vitro) and potentely translated\(^\).

**Ex vivo manipulations.** Although certainly not new, ex vivo manipulations may be more appreciated after successes such as the use of chimeric antigen receptor (CAR) T cells\(^\) and with new possibilities offered by the CRISPR–Cas9 system\(^{97,180}\). Most importantly, off-target effects can be monitored before re-implantation. The autologous tumour cell immunotherapeutic Vigil has shown promising results in advanced malignancies in phase I\(^\) and II\(^\) clinical trials. Vigil is generated from the patient’s tumour cells modified to express recombinant granulocyte–macrophage colony-stimulating factor (GM-CSF) and a bifunctional shRNA that targets furin. Furin inhibition results in downregulation of immunosuppressive cytokines TGFβ1 and TGFβ2, thereby supporting T cell responses in the tumour. Bifunctional shRNAs cause both mRNA degradation and inhibition of translation, causing stronger and prolonged RNAi effects and allowing lower dosing to reduce off-targeting\(^\). TGFβ1 and TGFβ2 were inhibited 90–100% in bi-shRNA furin-treated cells\(^\), and treatment of patients with advanced cancer with Vigil caused an increase in circulating activated T cells and significantly improved survival\(^{109,110}\). No long-term toxicity was reported in a 3-year follow up in patients with Ewing sarcoma who had received Vigil\(^\).

The CRISPR–Cas9 system\(^{97,180}\) offers new opportunities for therapeutic targeting of coding and noncoding genes. The system is capable of genome editing by active CRISPR–Cas9 or Cas12a (Cpf1)\(^\), gene interference or activation by catalytically dead (d)Cas9 fused to a repressive or activating effector domain\(^\), or RNA editing by the Cas13 variant\(^\). Clinical trials applying CRISPR–Cas for genome editing ex vivo are actively ongoing. For example, the disruption of the CCR5 gene, the predominant co-receptor for entry of the HIV virus into human T cells, in CD34\(^\)+ haematopoietic stem cells (HSCs) of HIV-infected subjects with haematological malignancies is being attempted (NCT03164135). The dCas9 system has so far been employed only in vitro, for example, to reactivate HIV long terminal repeat (LTR) transcription in latently infected cells to trigger their suicide death, presenting a potential curative treatment\(^\). Importantly, these HIV LTR-specific MS2-mediated single guide RNAs showed very few genome-wide off-target effects\(^\). Multiple lncRNAs have also been shown to activate (for example, MALAT1, HEAL, SAF) or repress (linncRNA-p21, NEAT1, GAS5, 7 SK, NRON, TAR-gag) HIV transcription, while lncRNAs NRON and uc002yug.2 contribute to HIV latency (reviewed in\(^\)). Furthermore, CRISPR–Cas9 is used to target the BCL11A gene enhancer, the disruption of which induced the expression of fetal haemoglobin ex vivo\(^\) in

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**NATURE REVIEWS | DRUG DISCOVERY**

**VOLUME 20 | AUGUST 2021 | 643**

**PERSPECTIVES**
CD34+ cells for treatment of β-thalassaemia (NCT03655678). The IncRNA UCA1 was recently shown to be crucially involved in human haem metabolism and erythrocyte differentiation of CD34+ HSCs219. This therefore shows that IncRNAs can expand the amount of therapeutic CRISPR–Cas9 ex vivo editing targets. For the CRISPR–Cas9 system to be applied in vivo, delivery issues especially regarding the large Cas proteins must be overcome and immunological responses (reviewed in200) must be carefully assessed.

The hurdle of delivery

Efficient delivery of RNA therapeutics not only to the organ and cell type of interest but also across the cell membrane to perform their intracellular functions is one of the greatest challenges in the field. Accordingly, the first and foremost reason for clinical trial termination regarding RNA therapeutics is lack of efficacy (TABLE 3). Efficient delivery of oligonucleotides is challenging owing to their instability, negative charge and hydrophilic nature preventing diffusion through cell membranes201. First- and second-generation chemical modifications improve stability and uptake by inducing resistance to nucleas degradation and increasing interaction with proteins129. Third-generation antisense technologies such as neutrally charged PMOs are highly stable but face difficulties with cellular uptake and require administration in high doses202. Different delivery systems are also being used, including lipid- and polymer-based vectors and ligand–oligonucleotide conjugate delivery systems. These delivery systems are taken up via different endocytosis mechanisms and consequently, endosomal escape of the RNA therapeutic must be facilitated to prevent its lysosomal degradation. Numerous strategies can be employed, such as the use of cationic lipids, viral or bacterial agents, or cell-penetrating peptides (CPPs), which are reviewed in detail elsewhere203,204.

Lipid nanoparticles (LNPs) can be easily modified, coupled to targeting moieties and are characterized by high biodegradability and biocompatibility with low immunogenicity. Surface modification with for example, hyaluronic acid (HA) or polyethylene glycol (PEG) reduces nonspecific uptake, cellular toxicity and aggregation, and improves characteristics such as tumour targeting and stability. Cellular uptake is mediated by endocytosis, and interactions between fusogenic cationic LNPs and anionic membrane lipids mediate the release of their cargo into the cytoplasm205. The FDA-approved siRNA therapeutic patisiran uses LNPs as a delivery method206, as did the miRNA mimic MRX34 [REF.17], and many more lipid-based RNA therapeutic delivery systems are being tested in clinical trials.

Polymers (synthetic or naturally based) are successful alternatives as miRNA delivery vectors. Polymer-based carriers are characterized over lipid carriers by their versatility in size, molecular composition and structure. Polymers such as polyethylene imine (PEI), polyactic-co-glycolic acid (PLGA), poly-amidoamine (PAMAM) and chitosan are being actively investigated in vitro and in vivo for the delivery of miRNA mimics or antisense oligonucleotides, either alone or in combination with chemotherapies for enhanced therapeutics. Many miRNAs have been delivered either systemically or locally through polymers, including miR-221/miR-222 [REF.207], miR-21 [REF.208,209], miR-150 [REF.116,211], miR-34a212, miR-145 and miR-33a213. Other studies address polymer combinations, such as PLGA–PEI LNPs modified with HA (for specific internalization by tumour cells) to deliver miR-145. After local administration of this construct to xenograft tumour-bearing mice, intact miRNA plasmid vectors were delivered, and stable expression of miR-145 was achieved following a significant antitumour effect214. PEI–PLGA–HA NPs were furthermore used to encapsulate miR-542-3p and doxorubicin for targeted delivery to triple-negative breast cancer cells215. An interesting development is also the LODER system, a miniature biodegradable polymeric matrix designed to slowly release siRNA against G12D-mutated KRAS upon local injection for the treatment of pancreatic cancer216,217. siG12D–LODER is currently being tested in a phase II clinical trial (NCT01676259).

The conjugation of oligonucleotides to various entities is being further explored for their delivery. Ligand conjugation is a major clinical strategy for delivery of RNA therapeutics to hepatocytes, offering selective delivery via receptor-mediated mechanisms218. N-acetylgalactosamine (GalNAc) is a high-affinity ligand for the hepatocyte-specific asialoglycoprotein receptor (ASGR), which mediates clathrin-mediated endocytosis. The endosomal release mechanism of GalNAc-conjugated RNAs is thought to be relatively inefficient and successful delivery is owed to the fast cycling of the ASGR219. Owing to its lysosomal degradation. Numerous linker-functionalized RNAs220. Although antibody conjugation is promising regarding target-organ delivery, for instance, a siRNA that targets myostatin was successfully delivered to skeletal muscle of mice by conjugation to an anti-CD71 antibody221, a major limitation is their entrapment in the endocytic compartment, which often results in limited efficacy222. The most promising developments regarding improved intracellular escape were seen with CPP conjugation. The efficacy of the splice-modifying PMO ASOs currently approved for treatment of Duchenne muscular dystrophy could be greatly enhanced upon conjugation to CPPs223. The CPP Pip8b2, for example, increased the uptake of DMD-targeting ASOs in vitro224, and HA2–ApoE(131–150) increased endosomal escape of nusinersen in mice225.

Other notable developments regarding delivery methods include, for instance, metal-based NPs, particularly gold NPs (AuNPs). Gold nanostructures show low toxicity profiles, may be conjugated to various compounds including carbohydrates, amino acids, peptides, proteins and oligonucleotides, and, most importantly, passively accumulate in tumour tissues226. The successful delivery of ss and ds oligonucleotides of up to 50 nucleotides or base pairs using AuNPs was first shown in vitro in a murine melanoma cell line227. Spherical nucleic acids (SNAs), that is, gold NPs densely coated with oligonucleotides228, have shown notable results as delivery vehicles to brain tumours. Systemic administration of miR-182 mimic-coated SNAs showed successful delivery to tumour in a murine glioblastoma model229. SNAs may also be combined with liposomal delivery, for instance, anti-miR-92b-loaded...
SNAs in apolipoprotein E (ApoE)-coated liposomes successfully crossed the blood–brain barrier in a murine glioblastoma model and reduced tumour cell viability232. Their clinical safety was proved in a phase 0 clinical trial in which SNAs coated with siRNAs targeting the Beclin1/LC3 oncogene were systemically administered to patients with glioblastoma (NCT03020017)233. The safety and proof of concept of adeno-associated virus 5 (AAV5)-mediated delivery is currently being tested for the potential treatment of Huntington disease (rAAV5-miHTT; NCT04120493). AAV5 has shown effective delivery to the brain in non-human primates234. With a huntingtin-targeting miRNA in a miR-451 backbone that is flanked by inverted terminal repeats to mediate genome integration, rAAV5-miHTT is intended as a one-time gene therapy235. Additional developments are being made with respect to the route of administration, such as inhalation (for example, pressurized dry powder inhalers, nebulizers, soft mist inhalers) that allow delivery of RNA-based therapeutics to specific regions in the pulmonary tract236,237.

In summary, LNPs, polymers, RNA conjugations, NP and virus-based approaches for delivery are already either used in clinically approved therapeutics or have entered clinical testing. Below, we highlight two alternative approaches that we think are promising candidates for clinical translation.

Potential solutions

**Exosome-mediated miRNA delivery.**

Cell-derived membrane vesicles (exosomes and microvesicles) have been reported to be effective delivery vectors. They are small sized (nanoscale) vesicles, secreted by various cells, and they contribute to intercellular communication through the delivery of different types of cargo, including proteins (cytoplasmic proteins, membrane proteins), lipids and nucleic acids (DNA, miRNAs and ncRNAs) involved in cell-to-cell communication180,239. Exosomes are thus considered natural carriers of miRNAs and may present an ideal delivery system owing to their negligible antigenicity, minimal cytotoxicity and their ability to bypass the endocytic pathway and circumvent phagocytosis240. The precise uptake mechanism for exosomes may differ depending on targeted cell type241. The exosomal transmembrane protein CD47 was shown to inhibit the phagocytosis of exosome delivery vehicles by monocytes, which increased their delivery to pancreatic tumours and the improved targeting of oncogenic KRAS242 (REF.243). The use of iExosomes (that is, engineered exosomes) loaded with anti-KRAS-G12D siRNA for the treatment of pancreatic cancer is currently being tested in a phase I clinical trial (NCT03608631).

Exosome biodistribution studies in animal models indicated accumulation in lung, liver and spleen upon intravenous injection, and in draining lymph nodes upon lymphatic injection244. However, a very recent study showed that genetic modification of the membrane proteins can vastly change the distribution patterns of exosomes245. Other work similarly suggests differences in biodistribution based on exosome subtypes246 and the preferential uptake of self-produced exosomes in parental cells247. These findings imply that exosome biodistribution can be tailored to specific needs based on their specific composition as well as the cells used for their production. Indeed, neuron-specific targeting of exosomes was achieved in mice through expression of exosomal membrane protein Lamp2B fused to the RVG peptide that specifically localizes to neurons248.

Exosomes separated from bone marrow-derived mesenchymal stem cells were induced to deliver LNA–anti-miR-142-3p to suppress the expression level of miR-142-3p to reduce the tumorigenicity of breast cancer cells. MiR-142-3p repression reduced the levels of miR-150 via derepression of miR-142-3p target gene adenomatous polyposis coli (APC) and consequent repression of Wnt–β-catenin signalling, which in turn induces miR-150 (REF.249). Furthermore, exosomes can be engineered to introduce multiple compounds, for example, an oligonucleotide and a chemotherapeutic agent, for a more efficient effect. For example, purified exosomes loaded with miR-21 and 5-fluorouracil (5-FU) were efficiently taken up by 5-FU-resistant HCT-116SFR cell lines. The combined delivery of miR-21 and 5-FU effectively reversed drug resistance of the cells and significantly elevated the cytotoxicity of cells compared with single treatment with 5-FU or miR-21 (REF.250).

Although using exosomes as a delivery platform is a promising strategy, large-scale production of exosomes is not easily available or commercially viable owing to the high cost. Production through chemically induced membrane blebbing has been suggested, which was shown to produce large numbers of highly homogeneous vesicles termed ‘extracellular blebs’251. In addition, food-derived exosomes have shown promising results in preclinical studies, including bovine milk-derived252 and grape-derived exosomes253. Grapefruit-derived exosomes coated with plasma membrane of activated leukocytes were successfully delivered to tumour inflammatory sites to attenuate tumour growth and colitis in mouse models. The homing of drug-loaded exosomes to inflammatory sites was mediated by chemokine receptors CXCR2 and LFA1 (REF.253). Furthermore, exosome production from delicate systems such as mesenchymal stem cells is being developed, employing, for example, 3D scaffolds, large-scale cell expansion methods and molecular modifications to increase exosome yield or genetic packaging (reviewed in244).

**Bacteriophage and bacterial minicell delivery vehicles.** Several studies have reported bacteriophages as a safe and effective delivery vehicle that can be loaded with RNAs or other drugs. For example, modification of MS2 bacteriophages to be used as virus-like particles (VLPs) could protect miRNAs against degradation. Self-assembly of MS2 by interaction between a specific MS2 cistron, a 19 nt sequence known as a pac site, and the bacteriophage coat protein allows the bacteriophage particles to be loaded with RNAs or other molecules. The covalent linkage of pre-miR-146a to modified MS2 bacteriophage particles significantly induced the expression of miR-146a in vitro and in vivo and resulted in target gene repression254. PP7 bacteriophages were similarly used to package pre-miR-23b. Insertion of a cell-penetrating peptide (CPP) into the AB loop of the PP7 bacteriophages facilitated the delivery of the pre-miR-23b into SK-HEP-1 hepatoma cells. Pre-miR-23b delivery using PP7 VLPs was followed by pre-miRNA processing to produce mature miR-23b, which consequently repressed its target gene cadherin and reduced cell migration255. Bacteriophages, like many other delivery agents and drugs, accumulate preferentially in liver and spleen as well as in solid tumours owing to the enhanced permeation and retention effect256,257. The modification of bacteriophage biodistribution is so far grossly understudied, but for example, splenic accumulation could be inhibited by PEG conjugation257 and their encapsulation by liposomes has improved oral administration258. Bacteriophages are currently being extensively explored in clinical trials as antibacterial agents, but not yet as delivery vehicles for RNA therapeutics. An important limitation could be the lack...
of naturally developed endolysosomal escape mechanisms. Achromosomal bacterial minicells are small size vesicles that are assembled by bacteria. Minicells with recombinant-plasmid DNA molecules are produced by transforming the parental bacterial strains with a plasmid of choice, before inducing production of minicells[26]. The ongoing MesosmiR-I phase I study (NCT02369198) is the first human study in which EnGeneIC Dream Vectors (EDV), that is, non-living bacterial minicells targeted with anti-EGFR antibody that encapsulate a tumour-suppressive miR-16 mimic (termed 'TargomiRs'), have been applied to patients with non-small cell lung cancer or mesothelioma[27,28]. This approach may be specifically beneficial for granting optimal intracellular delivery, as bacteria have a variety of natural endosomal escape mechanisms in place[29]. Minicells have shown promising results for specific delivery, including for example, EGFR-mediated delivery of doxorubicin-loaded minicells to the core of late-stage brain tumours in dogs[30], and the follic acid-mediated delivery of anti-VEGFA shRNA to murine xenograft tumours[31].

Outlook

The successful application of RNA-based therapeutics requires an unprecedented interdisciplinary approach, including technical advancements in molecular biology, immunology, pharmacology, chemistry and nanotechnology. An optimal RNA therapeutics should be extensively tested for immunogenicity, chemically modified to improve pharmacokinetics and pharmacodynamics, delivered under consideration of biodistribution patterns and intracellular escape mechanisms, specifically and potently interact with the intended target and be dosed at an optimal level to trigger the desired effect. Significant advancements in each of these areas has been made for RNA therapeutics individually, but their successful translation depends on further interdisciplinary development to improve tolerance, specificity and delivery. An example of such an interdisciplinary approach is the delivery of a miRNA inhibitor to heart tissue using an ultrasound and microbubble-targeted delivery system that suppressed cardiac hypertrophy in a mouse model[32]. Although challenging, with the many new, promising and creative developments that are emerging through valuable preclinical work, the challenges faced in the field of RNA therapeutics will ultimately be overcome.

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All authors contributed to the development of presented concepts and writing of the manuscript.

Competing interests

G.A.C. is one of the scientific founders of Ithax Pharmaceuticals. The other authors declare no competing interests.

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