Abstract: Neuromuscular disorders are a large group of rare pathologies characterised by skeletal muscle atrophy and weakness, with the common involvement of respiratory and/or cardiac muscles. These diseases lead to life-long motor deficiencies and specific organ failures, and are, in their worst-case scenarios, life threatening. Amongst other causes, they can be genetically inherited through mutations in more than 500 different genes. In the last 20 years, specific pharmacological treatments have been approved for human usage. However, these “à-la-carte” therapies cover only a very small portion of the clinical needs and are often partially efficient in alleviating the symptoms of the disease, even less so in curing it. Recombinant adeno-associated virus vector-mediated gene transfer is a more general strategy that could be adapted for a large majority of these diseases and has proved very efficient in rescuing the symptoms in many neuropathological animal models. On this solid ground, several clinical trials are currently being conducted with the whole-body delivery of the therapeutic vectors. This review recapitulates the state-of-the-art tools for neuron and muscle-targeted gene therapy, and summarises the main findings of the spinal muscular atrophy (SMA), Duchenne muscular dystrophy (DMD) and X-linked myotubular myopathy (XLMTM) trials. Despite promising efficacy results, serious adverse events of various severities were observed in these trials. Possible leads for second-generation products are also discussed.

Keywords: AAV; genetic neuromuscular disorders; gene therapy; clinical trials; toxicity; SMA; DMD; XLMTM

1. Introduction

Neuromuscular disorders are a group of heterogeneous rare diseases characterised by skeletal muscle dysfunction and caused primarily by motoneuron, peripheral nerve, motor end plate or muscle deficiencies. This family of pathologies encompasses a wide clinical spectrum, ranging from very weak and barely detectable clinical signs to extremely severe and life-shortening forms. Common symptoms include muscle-specific patterns of atrophy and weakness, occasionally associated with the involvement of additional organs, the most common complication being cardiac and/or respiratory failure. These diseases can be caused by many factors, notably autoimmunity; inflammation; poisoning; toxin accumulation; tumours; environmental agents; neurologic, metabolic or traumatic syndromes [1–3]; aging [1,4]; and genetic inheritance or spontaneous mutations in muscle or nerve-essential genes. The large majority of mutations are monogenic, with every nature of mutation and transmission mode possible. The classification of neuromuscular diseases based on their origins and phenotypical features is published every year at http://www.musclegenetable. The 2020 update
reports 1042 neuromuscular disorders caused by mutations in 587 different genes, classified in 16 groups, and many remain to be discovered [5]. The pathogenic mechanisms are very diverse, as they depend on the gene involved, and proteins with very different functions and subcellular localisations are affected (enzymes, structural proteins, metabolic key-players, etc). In this review, we will focus our interest on genetic neuromuscular diseases currently under interventional clinical trials with whole-body delivery.

2. Marketed Pharmacological Treatments

Before the 1990s, treatment options were limited to supportive therapies aiming at improving life comfort and lengthening lifespan. Anti-inflammatory drugs proved very efficient in preventing muscle degeneration and mortality in inflammatory myopathies [6]. Corticosteroids are also commonly used and show limited success in Duchenne muscular dystrophy (DMD, OMIM 310200), a very severe and the most common form of degenerative muscular pathology. Long-term clinical trials showed that prednisolone/prednisone or deflazacort corticosteroids reduce chronic muscle inflammation, stabilise muscle function, prolong ambulation and improve respiratory function and patients’ survival [7–9]. However, several side effects are associated with the prolonged usage of these immuno-modulators, the most severe being a drastic inhibition of the immune system’s functionality, occasionally leading to life-threatening opportunistic infections.

More recently, specifically targeted treatments were developed and approved for human applications by the regulatory agencies. The USA Food and Drug Administration (FDA) and the European Medicines Agency (EMA) approved Myozyme® (α-glucosidase, Sanofi-Genzyme, Cambridge, MA, USA) for long-term enzyme replacement therapy in Pompe patients, who suffer from a severe metabolic myopathy caused by mutations in the α-glucosidase-encoding gene (glycogen storage disease type 2 or Pompe Disease, OMIM 232300) [10–12]. The treatment proved particularly efficient in improving lifespan and muscle, respiratory and cardiac functions in classical infantile-onset Pompe disease (<1 year of age, with cardiomyopathy) [13,14], with more contrasted results in late-onset forms of the disease [15,16]. Occasional infusion-associated reactions and adverse events related to the treatment were reported. Nearly all were resolved with an interruption or reduction of the infusion rate or symptomatic treatment. In almost every case, repeated bi-monthly intravenous injections of the product led to the generation of α-glucosidase-specific antibodies [13–16], although seldom showed evidence of in vitro inhibitory activity. Some patients developed anaphylactic shock [15].

Exondys 51® (Eteplirsen, Sarepta Therapeutics, Cambridge, USA), a drug targeting the DMD pathology, was granted accelerated approval by the FDA in 2016 on the grounds of phenotype stabilisation, making it the first FDA-approved drug for DMD [17]. Severe Duchenne myopathy is caused by a variety of mutations in the dystrophin-encoding DMD gene [18]. The large majority are out-of-frame mutations resulting in the total loss of dystrophin, while the expression of shorter forms of dystrophin caused by in-frame DMD mutations leads to the milder Becker phenotype [19]. This observation constitutes the proof of concept that expressing shorter forms of dystrophin could be a therapeutic option for ameliorating, if not curing, DMD symptoms. Eteplirsen is a 30-nucleotide-long phosphorodiamidate morpholino oligomer (PMO) designed to skip DMD exon 51 and restore a shorter but functional reading frame. The weekly intravenous injection of this drug restores partial dystrophin expression in skeletal fibres [20], prevents muscle loss of function [21–23] and protects pulmonary and cardiac functions [24,25]. This drug offers a very good safety profile, probably due to its uncharged chemical nature. Of note, Translarna® (Ataluren, PTC Therapeutics, South Plainfield, NJ, USA), a read-through RNA interference molecule targeting non-sense mutations in Duchenne, showed a weak benefit in DMD ambulatory patients in clinical trials [26,27] and was granted conditional approval for ambulatory patients by the EMA in 2014, but was refused by the FDA. The treatment proved safe and delayed ambulation loss in longer-term studies compared with a historical cohort [28].

Spinraza® (Nusinersen, Biogen, Cambridge, MA, USA) was the first curative drug for spinal muscular atrophy (SMA) in paediatric and adult patients to be approved by the FDA in 2016 and the following year by the EMA [29,30]. SMA, the most common motoneuron degenerative disease
and the leading genetic cause of infant mortality, is due to hereditary bi-allelic mutations in the SMN gene [31]. Spinraza® is an antisense oligonucleotide interfering with the splicing of an alternative form of the gene (SMN2) and leading to the production of a functional SMN protein. Repeated intrathecal injections result in an increase in SMN proteins and meaningful improvement in motor development and function with the associated survival of the patients [32,33]. Just recently, in August 2020, the FDA approved Evrysdi® (Risdiplam, Genentech/Roche, San Francisco, CA, USA) as the first oral and at-home treatment for all SMA patients from 2 months of age [34]. Similarly to Spinraza, this SMN2 splicing modifier increases the levels of SMN proteins and shows clinically meaningful improvements in survival and motor and respiratory functions in SMA patients [35–37]. However, while Spinraza requires four administrations in the spinal cord a year, Evrysdi is taken orally for systemic distribution once a day, widening the field of application to patients excluded from intrathecal injections because of scoliosis.

Even though these drugs ameliorate the patient’s life and prognosis, they do not cure the diseases and necessitate constant re-dosing, a burdensome shortcoming for patients with an already altered quality of life. Long-term adverse events due to constant drug re-administration are also an important issue, especially as an immune response towards the treatment often develops with time, impeding its efficacy. Moreover, these personalised medicine treatments are generally highly specific for the targeted disease and mutation. Because of their wider range of application, a very intense research field is focused on developing gene replacement approaches. These strategies, which take advantage of the natural capacity of viruses to infect specific human cells, consist of inserting therapeutic genes in place of viral sequences in vectors devoid of replicative capacity. They offer the advantage of being usable regardless of the mutation type and position, at least for pathologies caused by losses of function. After a long period of difficulties linked mostly to the route of administration and to the production of the therapeutic vectors, the last ten years finally saw the translation of several proofs of concept into promising clinical trials. The vector favoured for the delivery of genes in neuromuscular tissues is derived from the adeno-associated virus (AAV). In the last ten years, several AAV-based treatments have been approved for human usage. In 2012, Glybera® (alipogene tiparvovec, UniQure, Lexington, KY, USA) was the first to be accepted by the EMA for the correction of a rare inherited metabolic disorder, substantiating AAV innocuousness and long-term efficacy [38,39]. Luxturna® (Voretigene Neparvovec, Spark Therapeutics, Philadelphia, PA, USA) was later approved for the local treatment of a rare retinal disease [40–42]. Very recently, regulatory agencies granted full (FDA) and conditional (EMA) approval to Zolgensma® (onasemnogene abeparvovec-xioi, AveXis/Novartis, Bannockburn, IL, USA), the first AAV-based treatment for the whole-body correction of SMA [43–45], paving the way for other myopathies.

3. The Therapeutic Toolbox for Muscle Gene Therapy

3.1. About Wild-Type AAV

The AAV virus is a 25 nm-diameter non-enveloped human parvovirus, with a simple architecture composed of a single-stranded 4.7 kb linear DNA genome encapsidated within an icosahedral protein capsid. The DNA bears four open-reading frames (ORFs) coding, respectively, for the four non-structural Rep proteins involved in the viral cell cycle (Rep 78, 68, 52 and 40); the three structural Cap proteins VP1, VP2 and VP3, assembling in a 1:1:10 ratio to constitute the 60 monomers of the capsid; the assembly-activating protein (AAP), promoting capsid assembly [46]; and the recently described membrane-associated accessory protein (MAAP) [47]. The ORFs are framed by two highly structured 145 bp palindromic inverted terminal repeats (ITRs) acting in cis as structural signals to drive AAV replication and genome packaging. AAV can infect both dividing and quiescent cells [48].

Various AAV serotypes of human and primate origin (AAV1 to AAV13) and more than a hundred natural variants have been identified (AAV1 [49], AAV2 and AAV3 [50], AAV4 [51], AAV5 [52], AAV6 [53], AAV7 and AAV8 [54], AAV9 [55], AAV10 and AAV11 [56], AAV12 [57] and AAV13 [58]).
Based on VP1-capsid composition, the AAVs were phylogenetically classified into six clades, regrouped together according to genetic relatedness [59]. Although many display a broad tissue tropism, they generally show preferential infections of specific organs. The cell tropisms depend on many parameters, but subtle differences in the capsid’s amino acid sequence and structure are one essential feature driving tissue targeting [55,60,61]. Once disseminated in the blood stream, AAVs have to overcome several barriers to deliver their DNA within host cells’ nuclei. First, AAVs can be neutralised by pre-existing neutralising antibodies (NAbs), as seroprevalence resulting from natural infections with wild-type AAV is common in the general human population, with a high cross-reactivity between serotypes [62–64]. Second, AAVs have to attach to specific receptors before being internalised within host cells. AAV capsids were shown to interact with specific glycan moieties of host membrane proteoglycans: heparan sulfate for AAV2 [65]; heparin for AAV3 and AAV6 [66]; sialic acid for AAV1, AAV4, AAV5 and AAV6 [66–69]; and galactose for AAV9 [70–72]. Transmembrane receptors such as PDGF for AAV5 [73] and the 37/67 kDa laminin receptor LamR for AAV8 [74] were also reported to be surface receptors. Their in vivo biodistribution correlates with and could account for virus tropism. Efficient virus endocytosis requires secondary binding events with membrane co-receptors. For AAV2, the most widely studied AAV, the hepatocyte growth factor receptor c-Met [75], αVβ5 integrin [76] and fibroblast growth factor receptor 1 (FGFR1) [77] were demonstrated to increase AAV infectiosity and proposed as co-receptors. However, using a candidate approach based on genetic deletion and supplementation, Pilay et al. demonstrated the existence of a co-receptor common to all the tested serotypes (AAV1, 2, 3B, 5, 6, 8 and 9), the previously uncharacterised type I transmembrane protein KIAA0319L, renamed AAVR [78,79]. Within the cell cytoplasm, AAV undergoes intracellular trafficking via the microtubule network to reach the nucleus [80] and achieves endosomal escape, nuclear entry and capsid unfolding. The AAV lytic cycle needs co-infection with a helper virus such as adenovirus [49], herpesvirus [81] or cytomegalovirus [82] for replication to occur. In the absence of this helper virus, the AAV enters a latent state. Several reports have evidenced the preferential integration of the AAV genome into the transcriptionally active environment of the AAVS1 locus in the q13.4-Ter region of host chromosome 19 genomic DNA [83–87]. Other hotspots were evidenced in chromosome 5p13.3 (AAVS2) and chromosome 3p24.3 (AAVS3) [88]. However, these studies were performed in cell culture, and it was recently evidenced in vivo that AAV mainly persists as transcriptionally active episomal forms and sometimes integrates randomly in the host genome [89]. Clonal integration in six oncogenes in liver tissue associated with hepatic tumorigenesis was also identified [89,90]. No specific enrichment was found in major AAV targets previously identified in cell lines [89,90].

### 3.2. Of the Usage of Recombinant AAV for Central Nervous System (CNS) and Muscle-Specific Targeting

For neuromuscular diseases, AAV vectors stand out as the most promising tools for driving body-wide muscle gene expression, as their wild-type counterparts have not been associated with a pathologic condition, they target myocytes and they are relatively poorly immunogenic. Nonetheless, several factors limit their application, mainly their low packaging capacity (<5 kb), especially as many neuromuscular genes are larger. Another issue is the targeting specificity, as specific gene delivery is desirable to reduce the risk of toxic off-target effects.

AAV recombinant vectors (rAAVs) derived from wild-type viruses are devoid of viral genetic elements, apart from the two ITRs in between which the transgene of interest is inserted. Plasmids encoding Rep, Cap and a helper are brought in trans within an appropriate production cell line to achieve DNA packaging [91]. AAV2-based recombinant genomes have been packaged in many different capsid types, resulting in a wide collection of “pseudotyped vectors” (rAAV2/X, where X stands for the capsid serotype). In the absence of the Rep gene, in both murine models and cell lines, the rAAV genome mostly concatamerises and forms circular, transcriptionally active episomes unable to divide when host cells cycle [92–94], or integrates at a very low rate in the host genome, randomly [48,95–97] or in preferential regions: near chromosomal instability points or in CpG islands, active genes and regulatory sequences [98–101]. In cell lines, chromosomal rearrangements were observed near the
AAV-host genome breaking points [102,103]. Importantly, in murine hepatocytes, rAAVs were also reported to integrate at a low frequency into chromosome 12, at the Rian locus (RNA imprinted and accumulated in the nucleus), upregulating neighbouring non-coding RNAs and genes [104,105]. This integration, suggested to participate in murine hepatocellular carcinogenesis, seems specific to neonate animals and to some genetic backgrounds, and was not seen in adult mice [106].

More than 20 years ago, rAAV2s were the first vectors to prove their efficacy for the efficient and persistent transduction of post-mitotic neuromuscular cells [95,107]. The local brain delivery of a reporter transgene placed under the control of a ubiquitous strong promoter resulted in neuron and, to a lower level, glial cell transduction in rodents [107]. The long-term transduction of muscle fibres was observed after the intramuscular injection of a reporter gene in wild-type mice and rhesus monkeys, pointing out the inter-species tropism of this vector [95]. However, rAAV2s have a preference for slow-twitch muscle fibres, which might restrict their therapeutic benefits [108]. Additionally, of all the serotypes identified to date, AAV2 is the most common target of pre-existing NAbs in human populations [62,63], which could potentially prevent effective transduction in most of the putative patients [109]. Finally, a side-by-side comparative study of rAAV1 to 9 carried out with a ubiquitously driven luciferase reporter transgene evidenced that rAAV2 is amongst the lowest for general and muscle-specific transduction after intravenous injection, the optimal administration route for myopathy [110]. Today, rAAV2 is mainly used for tissue-specific gene therapy, such as local brain injection in clinical trials aimed at CNS delivery for Parkinson’s disease [111].

Recombinant AAV1, 7, 8 and 9 showed higher muscle transduction than rAAV2 after local injection in mice [54,55,112–114] and dogs [115]. Muscle targeting was also achieved, though with lower efficacy with AAV5 [113] and AAV6 [114].

Apart from very few diseases in which a specific group of muscles are affected and can be targeted by local delivery, whole-body muscle transfer has to be achieved for myopathy treatments, and the delivery is usually performed by systemic administration, or specific cerebro-spinal fluid delivery in the case of CNS-specific pathology. The body-wide intravascular delivery of rAAV packaged with reporter genes confirmed the widespread dissemination and highest muscle tropism of rAAV1, 7, 8 and 9 in mice [54,110,116–121], dogs [115,122,123] and monkeys [119,124]. Conflicting publications report on rAAV9’s preferential tropism for fast fibres [120] or slow fibres [114], but as they were performed with different promotors and different murine genetic backgrounds, general conclusions cannot be drawn.

The vascular endothelium is a major barrier for rAAV tissue distribution. Its permeation through the use of vascular endothelium growth factor (VEGF) was once demonstrated to enhance tissue transduction with rAAV6, largely inefficient by the intravascular route [125], though this effect is lost at high doses of the vector, and ensuing attempts to use it failed [116]. Muscle ischemia, a feature associated with some myopathies [126], was also shown to improve muscle targeting, partly for the same reasons [127].

Skeletal muscles are composed of long-lived mature post-mitotic fibres and of satellite cells, a population of progenitors crucial for muscle regeneration. Ideally, for expression persistence, therapeutic vectors should target myofibres and satellite cells, but unfortunately, rAAVs are inefficient for satellite cell transduction [128]. However, even though the episomal DNA can be diluted by successive cell divisions during muscle growth or regeneration, the transgene genome was shown to be stable for years in terminally differentiated myocytes, leading to continuous transgene expression [95,124,129].

Heart targeting is essential for treating neuromuscular diseases with cardiomyopathic features. Interestingly, several reports showed that rAAV9s lead to the highest levels in heart muscle [110,118,119,130,131], although rAAV6’s superior cardiac efficacy was once reported [132]. Differences in the vector doses and systemic routes of administration are likely to account for this discrepancy. The cardiotropic properties of rAAV9s might originate, at least partly, from their specific binding to galactose receptors [70,71]. Interestingly, the intravascular delivery of rAAV9 is also an appealing strategy for CNS targeting, as this serotype is the most efficient for crossing the blood–brain
barrier. Indeed, motoneurons and glial cells were transduced in the spinal cords and brains of mice, cats and non-human primates [118,133–138]. This strategy is a safer alternative to local CNS delivery. This unique feature of rAAV9s could come from increased vascular permeability and/or from their attachment to specific receptors distinct from those of other serotypes, such as the galactose receptor [70–72]. Indeed, the crystallographic structure of AAV9 revealed the specificity of the capsid in regions associated with receptor attachment that could account for its unique cellular tropism [61].

Finally, serotypes AAV1–9 transduce the liver with very high efficacy in mice, dogs and primates [110,116–120,122,129,135,136], but apart from the transient elevation of the aminotransferase enzyme related to the expression of the GFP transgene [135], no serious adverse events (SAEs) related to the capsid were reported during the biodistribution studies. Capsid-specific NAbs commonly developed with both local and systemic injection, though the levels varied with the dose and route of administration, but no major immunotoxicity was evidenced [95,110,112,115,135,136].

3.3. Restricting Expression by Muscle and CNS-Specific Promoters

Apart from capsid choice, a careful selection of the transgene regulatory elements, especially the promoter, is essential for specific expression. Viral promoters, such as the cytomegalovirus (CMV) or the Rous sarcoma virus (RSV), have generally been used for proofs of concept in early muscular gene therapy development, as they allow broad and powerful transgene expression [139–142]. The CMV promoter is currently used in several clinical trials for Duchenne and Becker muscular dystrophies, sporadic inclusion body myositis and Pompe disease (NCT02354781, NCT01519349, NCT00428935 and NCT00976352) [143–147]. However, it is now known that eukaryotic cells progressively silence transgene expression driven by viral promoters as a result of an immune mechanism to shut off viral expression, limiting their use for gene therapy applications where the long-lasting expression of the transgene is crucial [148–151]. An alternative to limit transgene silencing is the use of eukaryotic constitutive promoters, such as the elongation factor 1α (EF-1α), phosphoglycerate kinase (PGK), ubiquitin C (UBC) or hybrid promoters such as the chicken β-actin promoter coupled with the CMV early enhancer (CAG promoter), which shows high levels of transgene expression [152]. Interestingly, the CAG promoter is currently being used in a gene therapy clinical trial aiming at treating SMA type 1 patients: it shows success in driving appropriate expression levels in target tissues, as clinically meaningful benefits are achieved [153–156]. Nevertheless, constitutive transgene expression, notably in antigen presenting cells (APCs), was reported to induce an immune response [157–159].

In order to minimise ectopic transgene expression, muscle-specific promoters such as muscle creatine kinase (MCK) [160], desmin (Des) [161] or α-myosin heavy chain (α-MHC) [162] have been developed, showing higher muscle specificity compared to constitutive promoters [161]. Transgene expression efficacy driven by the Des promoter was successfully demonstrated in 2014 in preclinical studies performed in murine and canine models of X-linked myotubular myopathy (XLMTM) [163] and has recently shown promising results in a clinical trial with meaningful improvements in neuromuscular and respiratory functions (NCT03199469) [164].

Despite specific gene expression, muscle-specific promoters usually do not allow a high level of transgene expression in muscle cells and have a large size, limiting the packaging capacity for the transgene. Therefore, different laboratories have developed truncated muscle-specific promoters by selecting specific regulatory sequences to optimise both promoter strength and muscle-specific expression [165–168]. In 2008, Wang et al. designed compact muscle-specific promoters by combining an 87 bp proximal basal MCK promoter with a double (dMCK) or a triple (tMCK) tandem of the modified MCK enhancer [169], leading to highly efficient shorter promoters of 509 bp and 720 bp, respectively [170]. These two hybrid promoters demonstrate high transgene expression in skeletal muscles (except for the diaphragm), with no expression in the brain or liver. Interestingly, the dMCK and tMCK promoters are not active in the heart, which could be an advantage for the gene therapy of muscular diseases without cardiomyopathies. Following a successful proof of principle of gene transfer efficacy using tMCK in mouse models of Charcot-Marie-Tooth neuropathy type 1A [171] and limb
girdle muscular dystrophy (LGMD) type 2D [172], this promoter has moved forward to clinical trials for these diseases (NCT03520751, NCT01976091 and NCT00494195) [173,174]. However, both dMCK and tMCK were reported to show fast-twitch myofibre preferences, which could limit treatment efficacy depending on the pathology [170]. Inversely, the MHCK7 promoter (770 bp), based on the assembly of specific enhancer and promoter regions of MCK and α-MHC, was shown to direct high levels of transgene expression specifically in the skeletal and cardiac muscles, with the advantage of being expressed in both fibre types [175], and proved more efficient for muscle expression than MCK1 in a murine model of Pompe disease [176]. This promoter was shown to direct robust micro-dystrophin expression in a systemic gene replacement clinical trial for Duchenne muscular dystrophy [177]. Compact muscle-specific promoters were also designed by assembling multiple copies of myogenic regulatory elements of natural muscle promoters and enhancers. The synthetic muscle-specific C5.12 promoter was reported to present a 6- to 8-fold expression increase over the CMV promoter [178].

The use of tissue-restricted promoters has also revealed their ability to evade undesirable adaptive immune responses directed against the transgene product. A possible explanation for these results is the inhibition of transgene expression in transduced professional APCs. Cordier et al. previously showed that inserting the muscle-specific C5.12 promoter instead of the ubiquitous CMV promoter enables human γ-sarcoglycan expression in mice, probably impairing the anti-transgene immune response [179]. The same observation was made with α-sarcoglycan driven by the C5.12 promoter [180] or α-galactosidase A driven by the DC190 liver promoter for treating Fabry disease [181]. Another hybrid promoter, also based on the MCK enhancer and coupled with the SV40 promoter (MCK/SV40), resulted in the long-term sustainability of the transgene expression with a minimal cellular and humoral immune response compared to the ubiquitous CMV and CAG promoters, suggesting benefits for gene therapy applications with immunogenic transgenes [168].

Liver targeting is important to promote tolerance to the transgene product in order to lead to stable muscle expression [182,183]. In this context, Colella et al. designed a new tandem promoter enabling the expression of the transgene in the targeted muscle cells for treatment efficacy, as well as in hepatocytes to trigger immune tolerance to the transgenic protein [184]. To combine muscle-specific and hepatic transgene expression, both the apolipoprotein E enhancer (ApoE) and the human alpha-1 anti-trypsin promoter elements, known to allow tolerogenic transgene expression in the liver, were multiplexed with the muscle-specific C5.12 promoter. This approximately 1 kb hybrid promoter efficiently promotes transgene expression in muscles and prevents transgene immunity.

4. Translating Preclinical Studies into Clinical Trials

The achievable skeletal muscle, heart and CNS-specific targeting, together with the apparent safety of capsids, paved the way for the preclinical assessment of AAV-driven therapies for myopathies. Dozens of proofs of concept were made, but we will focus this discussion on the strategies that were translated into clinics for the whole-body treatment of SMA, DMD and XLMTM congenital myopathy. Nonetheless, Table 1 provides a general overview of the main clinical features and SAEs observed in all body-wide and CNS-targeted AAV-driven interventional clinical trials ongoing for neuromuscular disorders.
| Disease | AAV Serotype | Promoter | Transgene | Name | Administration/Dose | Clinical Trial ID (Study Name) | Sponsor/Collaborator | Study Phase/Status | Study Timelines (Clinical Follow-Up) | Age, Gender, Actual or Estimated Planned Number of Participants Enrolled | Serious Adverse Events |
|---------|--------------|----------|-----------|------|---------------------|------------------------------|----------------------|------------------|--------------------------------------|------------------------------------------------|---------------------|
| DMD     | AAV9         | CK8      | Micro-dystrophin | SCT-001 | Intravenous 2 doses | NCT03885742 (IGNITE DMD) Solid Biosciences, LLC | Phase 1/2 active, not recruiting | 2017–2024 (2 years) | 4 to 17 years, males, n = 16/same as current | Complement activation kidney failure, platelet count drop (n = 1 at 5 × 10^{13} vg/kg) + cardiopulmonary insufficiency (n = 1 at 2 × 10^{13} vg/kg) [185,186] |
| DMD     | AAVrh74      | MHCK7    | Micro-dystrophin | SRP-9001 | Intravenous | NCT0375164 Sarepta Therapeutics, Inc. | Phase 1/2 active, not recruiting | 2018–2021 (3 years) | 3 months to 7 years, males, n = 4/12 | No serious adverse events [177] |
| DMD     | AAVrh74      | MHCK7    | Micro-dystrophin | SRP-9001 | Intravenous 1 dose | NCT0379116 Sarepta Therapeutics, Inc. | Phase 2 active, not recruiting | 2018–2026 (5 years) | 4 years to 7 years, males, n = 41/24 | - |
| DMD     | AAV9         | Human muscle-specific | Mini-dystrophin | PF-06939926 | Intravenous 1 × 10^{14} vg/kg 3 × 10^{14} vk/kg | NCT0362502 Pfizer | Phase 1B active, not recruiting | 2018–2026 (5 years) | 4 years and older, males, n = 30/12 | Antibody response, complement activation, acute kidney injury, haemolysis, thrombocytopenia (n = 1 at 3 × 10^{14} vg/kg) [185] |
| SMA     | AAV9         | Hybrid CMV enhancer/chicken β-actin promoter | Human SMN | AVXS-101 | Intravenous 6.7 × 10^{13} vg/kg 2 × 10^{14} vg/kg | NCT02122952 AveXis, Inc. | Phase 1 completed | 2014–2017 (2 years) | Up to 6 months of age, males and female, n = 15/9 | Elevated serum aminotransferase levels (>10× normal level) [14] |
| SMA     | AAV9         | Hybrid CMV enhancer/chicken β-actin promoter | Human SMN | AVXS-101 | Intravenous therapeutic dose | NCT0388277 (STRIVE) AveXis, Inc. | Phase 3 completed | 2017–2019 (18 months of age) | Up to 6 months of age, males and females, n = 22/15 | - |
| SMA     | AAV9         | Hybrid CMV enhancer/chicken β-actin promoter | Human SMN | AVXS-101 | Intrathecal 6 × 10^{13} vg 1.2 × 10^{14} vg 2.4 × 10^{14} vg | NCT03881729 (STRONG) AveXis, Inc. | Phase 1 suspended (on clinical hold pending further discussions regarding pre-clinical findings) | 2017–2021 (15 months) | 6 to 60 months of age, males and females, n = 51/27 | SAE mainly related to the disease itself (n = 7). Transaminits events probably related to treatment (n = 2). [188] |
| Disease | AAV Serotype | Promoter | Transgene | Name | Administration/Dose | Clinical Trial ID (Study Name) | Sponsors/Collaborators | Study Phase/Status | Study Timelines (Clinical Follow-Up) | Age, Gender, Actual or Estimated Number of Participants Enrolled | Serious Adverse Events |
|---------|--------------|----------|-----------|------|----------------------|------------------------------|----------------------|------------------|--------------------------------------|---------------------------------------------------------------|---------------------|
| SMA     | AAV9         | Hybrid CMV enhancer/chicken β-actin promoter | Human SMN | AVXS-101 | Intravenous | NCT03461289 (STRIVE-EU) AveXis, Inc. | AveXis, Inc. (STRIVE-EU) AveXis, Inc. | Phase 3 completed | 2018–2020 (18 months of age) | Up to 6 months of age, males and females, n = 33/30 | - |
| SMA     | AAV9         | Hybrid CMV enhancer/chicken β-actin promoter | Human SMN | AVXS-101 | Intravenous | NCT03505099 (SPRINT) AveXis, Inc./PRA Health Sciences | AveXis, Inc./PRA Health Sciences | Phase 3 active, not recruiting | 2018–2021 (18 and 24 months of age) | Up to 42 days, males and females, n = 30/44 | - |
| SMA     | AAV9         | Hybrid CMV enhancer/chicken β-actin promoter | Human SMN | AVXS-101 | Intravenous single dose | NCT03837184 AveXis, Inc./PRA Health Sciences | AveXis, Inc./PRA Health Sciences | Phase 3 active, not recruiting | 2019–2021 (18 months of age) | Up to 6 months of age, males and females, n = 2/6 | - |
| XLMTM   | AAV8         | Dos      | Human MTM1 | AT132 | Intravenous | NCT03199469 (ASPIRO) Audentes Therapeutics | Audentes Therapeutics | Phase 1/2 active, not recruiting (FDA placed on clinical hold since June 2020) | 2017–2024 (5 years) | Up to 5 years, males, n = 24/12 | Progressive liver dysfunction, hyperbilirubinemia, death from sepsis or gastrointestinal bleeding (n = 3/17 at 3 × 10^14 vg/kg) [189] |
| Pompe   | AAV2/8       | Liver-specific promoter | hGAA | ACTUS-101 | Intravenous 2 doses | NCT03330673 Asklepios Biopharmaceuticals, INC./Duke University and National Institute of Arthritis and Musculoskeletal and Skin Diseases (NIAMS) | Asklepios Biopharmaceuticals, INC./Duke University and National Institute of Arthritis and Musculoskeletal and Skin Diseases (NIAMS) | Phase 1/2 recruiting | 2018–2022 (52 weeks) | 18 years and older, males and females, n = 8/6 | - |
| Pompe   | AAV          | Liver-specific promoter | hGAA | SPK-3006 | Intravenous dose escalation | NCT04935304 (RESOLUTE) Spark Therapeutics | Spark Therapeutics | Phase 1/2 Recruiting | 2020–2023 (32 weeks) | 18 years and older, males and females, n = 20/same as current | - |
| Pompe   | AAV8         | Hybrid liver/desmin promoter | hGAA | AT845 | Intravenous 2 doses | NCT04174105 (FORTIS) Audentes Therapeutics | Audentes Therapeutics | Phase 1/2 Recruiting | 2020–2027 (5 years) | 18 to 81 years, males and females, n = 8/same as current | - |
| Danon   | AAV9         | CAG      | hLAMP2B | RP-A501 | Intravenous 2 doses | NCT03882437 Rocket Pharmaceuticals Inc. | Rocket Pharmaceuticals Inc. | Phase 1 recruiting | 2019–2023 (3 years) | 8 years to 14 years and 15 years and older, males, n = 24/same as current | - |
Table 1. Cont.

| Disease         | AAV Serotype | Promoter | Transgene | Name  | Administration/Dose | Clinical Trial ID (Study Name) | Study Phase/Status | Study Timelines (Clinical Follow-Up) | Age, Gender, Actual or Estimated, Planned Number of Participants Enrolled | Serious Adverse Events |
|-----------------|--------------|----------|-----------|-------|---------------------|-------------------------------|---------------------|--------------------------------------|-------------------------------------------------------------------|-----------------------|
| LGMD2E          | scAAV/ rh24  | MHCK7    | SGCb      | SRP-9003 | Intravenous 5 × 10^{13} vg/kg | NCT010652259 Sarepta Therapeutics, Inc. | Phase 1/2 active, not recruiting | 2018–2020 (3 years) | 4 to 15 years, males and females, n = 6/9 | Elevated liver enzymes associated with transient increase in bilirubin (n = 1) [190] |
| Batten disease  | AAV2         | CU       | hCLN2     | -     | CNS administration 3 × 10^{12} vg | NCT010151216 Weill Medical College of Cornell University/Nathan’s Battle Foundation | Phase 1 completed | 2004–2019 (18 months) | 3 to 18 years, males and females, n = 10/11 | - |
| Batten disease  | AAVrh.10     | CU       | hCLN2     | -     | Direct CNS administration 9 × 10^{13} vg/2.85 × 10^{11} vg | NCT01014985 Weill Medical College of Cornell University | Phase 1/2 completed | 2010–2017 (18 months) | 3 to 18 years, males and females, n = 8/16 | - |
| Batten disease  | AAVrh.10     | CU       | hCLN2     | -     | Direct CNS administration 9 × 10^{13} vg/2.85 × 10^{11} vg | NCT01014576 Weill Medical College of Cornell University/National Institute of Health | Phase 1 active, not recruiting | 2010–2032 (18 months) | 2 to 18 years, males and females, n = 25/16 | - |
| Batten disease  | scAAV9       | CB       | CLN6      | AT-GTX-501 | Intrathecal | NCT01225080 Amicus Therapeutics | Phase 1/2A active, not recruiting | 2016–2021 (24 months) | 1 year and older, males and females, n = 13/6 | - |
| Batten disease  | scAAV9       | P546     | CLN3      | AT-GTX-502 | Intrathecal 2 doses | NCT01072572 Amicus Therapeutics | Phase 1/2A active, not recruiting | 2018–2023 (36 months) | 3 to 10 years, males and females, n = 7/same as current | - |
| GSD1a           | AAV8         | Native promoter | G6Pase  | DTX401 | Intravenous 3 doses | NCT010317085 Ultragenyx Pharmaceutical INC | Phase 1/2 recruiting | 2018–2020 (52 weeks) | 18 years and older, males and females, n = 18/9 | No treatment-related serious adverse events reported to date |

DMD: Duchenne muscular dystrophy; SMA: Spinal muscular atrophy; XLMTM: X-linked myotubular myopathy; LGMD2E: Limb girdle muscular dystrophy type 2E; GSD1a: Glycogen storage disease type 1a.
4.1. SMA Trial

The ubiquitous SMN protein plays a key role in RNA regulation, and its deficiency in SMA is associated with cell-specific pre-mRNA splicing defects, possibly accounting for the tissue selectivity of the pathology [191]. Indeed, lower motoneurons are the cells primarily affected by degeneration in SMA, but other tissues, in particular, the heart, are also occasionally affected [192]. Local CNS delivery performed by the intrathecal administration of an rAAV9-hSMN vector proved efficient in correcting the motoneurons pathology in mice at doses in the $10^{13}$ vg/kg range [193]. Widespread distribution of the transgene was also demonstrated in non-human primate (NHP) spinal cord and brain motoneurons. However, local delivery might be clinically risky, and limits body-wide vector distribution and the correction of extra-CNS symptoms. Interestingly, taking advantage of the ability of AAV9 to cross the blood-brain barrier after systemic administration, three independent laboratories reported the preclinical safety and efficacy of an rAAV9-human SMN vector in different animal models [194–197]. All these studies used a self-complementary (sc) vector, which bears a DNA construct enabling the shunting of the transcription of the second DNA strand and hence leads to quicker gene expression than conventional single-stranded vectors [198]. A remarkable rescue of the phenotype was observed in mouse and cat models of SMA receiving intravenous doses ranging from $3 \times 10^{13}$ to $3.3 \times 10^{14}$ vg/kg of body weight (see Table 2 for details). The treatments rescued survival and all the major clinical manifestations, such as muscle atrophy and weakness, respiratory distress, weight loss and paralysis. The correction of murine cardiomyopathy was also reported [195]. An extensive motoneuron distribution was confirmed in cynomolgus macaques [194]. The product used by Barkats and collaborators has similar effects to the one used by Kaspar and collaborators at a 10-fold lower AAV dosage [194,196] (Table 2). This might be due to the codon-optimised enhancement of the transgene expression and/or by way of promoter regulation. Widespread motoneuron transduction was observed in the spinal cord, and the heart, skeletal muscles and liver were also highly transduced. Apart from the necrosis of the tails and ears seen in long-term survivors and attributed to the lack of SMN in these tissues [194,196,197], no safety issues were evidenced, and a clinical trial was initiated in 2014 [44]. Fifteen 0.9- to 7.9-month-old patients were treated intravenously with an rAAV9-hSMN product controlled by the hybrid CMV enhancer/chicken-$\beta$ actin promoter (product referred to as AVXS-101), either with $6.7 \times 10^{13}$ vg/kg (three patients) or $2 \times 10^{14}$ vg/kg (twelve patients) (AveXis/Novartis, NCT02122952) [44]. To this day, all the patients are alive and show significant amelioration of motor, respiratory and nutritional functions [44,153]. The improvements are substantial as seen from a comparison with a natural history cohort [154]. The effect is dose-dependent and related to the time of initiation: the earlier the injection, the more efficient the treatment [156]. This treatment proved more efficient than Spinraza® [155]. It is also longer-lived and safer, as Spinraza® necessitates constant re-administration by the risky intrathecal route [199]. Additional benefits might come from the widespread correction of SMN-related defects in other organs, the most important being the cardiac tissue. To date, fifty-six SAEs have been reported, amongst which two are deemed treatment-related [44]. They are limited to elevated serum aminotransferase levels reaching more than 10 times the normal range, without any other liver enzyme abnormalities or clinical manifestations. This important elevation of hepatic enzymes was rescued by a short course of glucocorticoids in the first patient, treated with a low dose (prednisolone, 1 mg/kg/day for 30 days, starting one day before AAV injection), which was thereafter administered systematically one day before the treatment administration to prevent liver-related toxicity. Granted these excellent results, the FDA approved AVXS-101 for usage in SMA patients in the USA in May 2019 [43]. This new drug goes by the name of Zolgensma and is the third AAV-based gene therapy approved to date for genetic diseases.
Table 2. Preclinical animal studies in mice models of SMA.

| Reference | Promoter | Codon Optimisation | Dose vg/per Mouse | Dose vg/kg of Body Weight | Expression in CNS | Mean Survival (Days) | Adverse Events |
|-----------|----------|--------------------|-------------------|---------------------------|------------------|----------------------|----------------|
| [196]     | PGK      | Yes                | $4.5 \times 10^{10}$ | $3 \times 10^{13}$       | SC: 80–140% of WT levels | 160 d (in 100% mice) | - Hyperactivity     |
|           |          |                    |                   |                           | Brain: low        |                      | - Tail necrosis    |
|           |          |                    |                   |                           |                  |                      | - Ear necrosis     |
|           |          |                    |                   |                           |                  |                      | - Bilateral cataract|
| [194]     | CBA      | No                 | $5 \times 10^{11}$ | $3.3 \times 10^{14}$     | SC: 42% of WT levels | >250 d (n = 4, 1 death at d 97) | - Necrotic pinna |
|           |          |                    |                   |                           |                  |                      |                 |
| [197]     | CMV      | Yes                | $1 \times 10^{11}$ | $6.7 \times 10^{13}$     | Lumbar SC: 66.5% MN | 69 d (in 80% of mice) | - Short tail       |
|           |          |                    |                   |                           | Thoracic SC: 45% MN|                      | - Ear necrosis     |
|           |          |                    |                   |                           | Cervical SC: 55% MN|                      | - Moderate eyelid inflammation |

All mice were from the SMN delta 7 strain (SMN2<sup>+/−</sup>, SMNA7<sup>+/−</sup>, smn<sup>−/−</sup>) and received an intravenous injection in the facial vein at p1 of an AAV9 solution. The doses expressed in vg/kg were calculated using a mouse body weight at p1 of 1.5 g (the average weight of a p1 SMN-Delta7 pup [193]).
Intriguingly, a preclinical report published after this trial’s initiation demonstrates the acute toxicity of a closely related product composed of an identical CAG-hSMN cassette packaged in the rAAV9 variant AAVhu68 and injected into wild-type NHPs and piglets at $2 \times 10^{14}$ vg/kg (the highest dose in the SMA trial) [200]. The biological abnormalities did not resolve in one out of three injected NHPs, leading to euthanasia at Day 5. No piglets died. The vector genome copy number was roughly 1000-fold higher in the liver than in other tissues. Intense dorsal root sensory neuron degeneration was evidenced in both the NHPs and piglets, with additional acute hepatocellular injury and liver failure, systemic inflammation and internal haemorrhage in monkeys. Because of the acute time course (abnormal parameters at Day 4–5), the toxic effects are not thought to be related to the activation of an adaptive immune response to the capsid or transgene and destruction of hepatocytes. They are more likely to result from the activation of an intracellular cellular stress pathway linked to genome or capsid overload in hepatocytes, together with the activation of systemic inflammation and the associated coagulopathy. In line with these findings, another AAV9-derived vector, AAV-PHP.B, coding for an unrelated GFP transgene and injected in NHPs at a slightly lower dose of $7.5 \times 10^{13}$ vg/kg (nearly identical to the lower dose of $6.7 \times 10^{13}$ vg/kg in the SMA trial) led to similar toxic effects on the liver and to thrombocytopenia and haemorrhage [201]. Here again, the time course of the acute symptoms is not consistent with an undesirable activation of the adaptive immune system. It is unclear whether liver damage or coagulopathy is the primary defect, but it is worth mentioning that the liver damage induced by some viral infections participates in lowering the platelet number, although this mechanism was not reported for AAV vectors [202]. Whether the toxic effects are related only to the dose or to the capsid used remains unclear. AAVhu68 and AAV PHP-B are closely related to AAV9 (two and seven amino acids of variation, respectively), but that could substantially change vector entry and processing. It would seem so, as high doses of rAAV9 ranging from $7.5 \times 10^{13}$ vg/kg [201] to $1–3 \times 10^{14}$ vg/kg [138] did not lead to toxicity in NHPs. The toxicity might also be related to the species used (NHPs and piglets) and the health status (wild-type animals) and will not necessarily translate into human toxicity in patients, as vector processing might be substantially different. The toxicity could also relate to the un-unified mode of vector purification and contaminants, especially as the ratio of empty/full capsids varies according to protocols. Thus, the therapeutic window is probably quite narrow in the SMA trial, as high levels of vector are necessary to achieve therapeutic benefit.

4.2. DMD Trials

DMD is a devastating and the most common muscle degenerative pathology, and as such, it is the subject of many therapeutic attempts. Respiratory and cardiac complications are common, and patients’ lifespans are severely reduced. DMD is the largest human gene ($\approx 14$ kb cDNA, NM_004006.2), which impedes its encapsidation within an AAV. Dystrophin is composed of an N-terminal actin-binding domain, 24 spectrin-like repeats articulated by four hinge regions and, at the C-terminal extremity, a cysteine-rich domain and a specific C-terminal domain. This highly flexible molecule interacts with a membrane-bound molecular complex (DAPC for dystrophin-associated-protein complex) and with sarcomeric actin, ensuring the plasticity of the muscle structure and resistance to contraction-induced injury. The observation that the deletion of a large part of the central domain leads to a very mild phenotype in patients set the ground for a large number of therapeutic trials aiming at expressing dystrophin forms shortened in the spectrin-like region [19]. Two main paths have been followed: shortening the natural gene by inducing the skipping of specific exons and restoring the reading frame (mutation-specific therapies) or bringing in trans a reduced version of the gene (a therapy amenable to all forms of dystrophinopathies). A founder paper of Chamberlain’s team established the importance of the different domains by investigating the phenotypes of transgenic mice in which various forms of shortened dystrophin, named micro (<30% of the full-length coding sequence)- or mini-dystrophin, were expressed [203]. Exon-skipping feasibility and efficacy was demonstrated by bringing the adequate oligonucleotide within myofibres using a U7-driven AAV [204–209]. The gene transfer of micro-dystrophin offers more versatile options and
proved very efficient in improving muscle force, protecting muscle from contraction-induced lesions and improving heart function in murine and canine models of the disease using the rAAV serotypes 6 [125,210–212], 8 [213–216] and 9 [139,141,217–221] delivered by the intravascular route. The most commonly used micro-dystrophins are variants of the ΔR4-R23ΔCT form, but the inclusion of the R16-R17 spectrin-like domains, involved in linkage to the membrane-bound cell metabolism regulator nitric oxide synthase NOS [222], was proposed to have additional therapeutic benefits [140,218]. Based on these preclinical proofs of concept, three clinical trials using micro-dystrophin gene transfer have been in progress since 2017 (Pfizer [New York, NY, USA], NCT03362502; Sarepta Therapeutics: NCT03375164; Solid Biosciences [Cambridge, MA, USA], NCT03368742). The three trials use muscle-specific promoters, rAAV9 or rAAVrh74 serotypes, and high and comparable doses of vector (1 × 10^{14} to 3 × 10^{14} vg/kg) (see Table 3). AAVrh74 was chosen by one group (Sarepta’s trial) because of its simian origin, which should decrease the likelihood of pre-existing immunity. Indeed, in a population of DMD patients, AAVrh74 sero-prevalence was shown to be low (measured in fewer than 20% of the patients tested) [223], and the average titres were also amongst the lowest [224]. Quite surprisingly, this seems to be a specific feature of DMD, as higher titres of antibodies are measured in non-DMD children, possibly owing to the small size of the population or to a disease-specific effect on AAV biology [224]. Another study even showed higher levels of antibodies against rAAVrh74 than against other serotypes in a healthy child population, probably because of cross-reactivity with serotypes present in humans [225]. Furthermore, AAVrh74’s safety has been demonstrated in a preclinical dose-escalation study in Duchenne’s model mice and in NHPs [226,227], as well as in humans in a clinical trial targeting LGMD, though the doses used were 100-fold lower than in the current DMD trial (1 × 10^{12} or 3 × 10^{12} vg/kg in the LGMD trial versus 2 × 10^{14} vg/kg in the DMD trial) [228]. The minimal effective dose was defined as 2 × 10^{14} vg/kg in mdx mice, a DMD model, and safety was confirmed in NHPs at doses reaching up to 6 × 10^{14} vg/kg [226,227,229]. One year after a single injection of 2 × 10^{14} vg/kg of AAVrh74-MHCK7-coΔR4-R23ΔCT (SRP-9001) in four patients, the first results are encouraging in terms of safety [177]. No SAEs were reported, and 18 mild or moderate events were deemed treatment-related. As previously observed in haemophilia [230,231] and SMA [44] clinical trials, liver enzymes peaked and diminished with a glucocorticoid course (n = 3). No adverse immune responses occurred, and, as expected, a transitory T cell response and the development of stable titres of antibodies against AAVrh74 were observed. The product was highly expressed, as seen in biceps brachii biopsies. Whether the treatment has any beneficial effects remains to be assessed more closely, although a clinically meaningful improvement of 2.2 to 7 points on the NorthStar Ambulatory Assessment score (NSAA) multi-parametric scale (maximum score of 34) suggests motor function improvement. A comparison with a historical cohort of untreated patients and longer time of treatments is needed to draw more definite conclusions. These encouraging results preclude dose escalation, and a new randomized, placebo-controlled clinical trial with a much larger sample size is under way (NCT03769116).
Table 3. Micro-dystrophin clinical trials in DMD patients. NSAA: NorthStar Ambulatory Assessment score, SM: Skeletal muscle.

| Trial Promoter/Product Name/Reference | Vector | Promoter | Micro-Dystrophin Domains | Dose vg/kg of Body Weight | Expression in SM | NSAA | Serious Adverse Events |
|--------------------------------------|--------|----------|--------------------------|---------------------------|------------------|------|------------------------|
| Sarepta SRP-9001-101 [177]            | AAVrh74| MHCK7 (SM and cardiac) | coΔR4-R23/ΔCT             | 2 × 10^{14}               | 95.8% of normal  | 5.5 points increase after 1 year | |
| Pfizer PF-06939926                    | AAV9   | Human muscle specific | -                         | 1 × 10^{14} 3 × 10^{14}   | 23.6% of normal 29.5% of normal | 2 points increase after 1 year | In 1 patient at 3 × 10^{14} vg/kg: complement activation, acute kidney failure, thrombocytopenia |
| Solid SGT-001                         | AAV9   | CK8      | ΔR2-R15/ΔR18-R22/ΔCT     | 2 × 10^{14} 2 doses       |                  |      | Complement activation, acute kidney failure, thrombocytopenia (2 SAEs in 6th patient) |
However, at very close doses, two other products composed of an rAAV9, a muscle-specific promoter and a micro-dystrophin transgene led to product-related SAEs. In one trial (product PF-06930026, Pfizer), the six participants included to date have shown a mean of ≈40% dystrophin-positive fibres at 1 × 10^{14} vg/kg and ≈70% at 3 × 10^{14} vg/kg in a bicep biopsy taken two months after injection, corresponding altogether to ≈24 to 30% of normal dystrophin expression [187]. The NSAA score increased by 4.5 points after one year in two participants treated with the lowest dose. However, one child treated with 3 × 10^{14} vg/kg developed a rapid antibody response with complement activation, acute kidney injury, haemolysis and thrombocytopenia. A transient 2-fold elevation of liver serum enzymes was observed, though it was not considered significant enough to indicate hepatic failure. Suspected complement-mediated nephropathy resulted in a protocol-driven pause of enrolment. Haemodialysis together with a course of complement inhibitor solved the problem in fifteen days. In the third trial (product SGT-001, Solid Biosciences), which differs slightly by the construct used (product SGT-001, different promoter and integration of the nNOS-binding domain in the transgene), similar treatment-related toxic events were seen in two patients at doses of 5 × 10^{13} and 2 × 10^{14} vg/kg. To date, six patients have been included, three at low and three at high doses. The preliminary results showed weak dystrophin expression in the three patients who received low dosages. The first patient injected at 5 × 10^{13} vg/kg developed complement activation, kidney failure and platelet count drops without signs of liver damage. The clinical hold [186] was lifted in 2018 after full symptom resorption following treatment with a modified course of steroids and a complement inhibitor and a change in the study design (the inclusion of an intravenous glucocorticoid administration in the first weeks following drug injection). A second patient dosed at 2 × 10^{14} vg/kg developed the same symptoms together with cardiopulmonary decline, leading to a second FDA hold of the trial. The SAEs fully resolved, but the clinical trial remained on hold on the grounds of remaining questions related to the mode of production of the product [185,232] and was finally allowed to continue in October 2020 [233]. A dose-finding study in a canine model of the pathology did not evidence any safety issue for this product at doses reaching 5 × 10^{14} vg/kg [221,234]. These SAEs could be related to the AAV9 capsid, though no severe side effects were observed in the SMA trial with this serotype at an equivalent dosage. The genetic background might account for the different effects between the SMA and DMD trials, whether for vector processing or the immune response. In the absence of liver injury, an immune response-mediated platelet drop, complement activation and ensuing nephropathy might be a reasonable hypothetical pathogenic mechanism. This could also be in line with the incidence of the age of the patients, as younger children are included in the only trial without SAEs, and the immune system is immature at a younger age [235]. The information on the three DMD trials is summarised in Table 3.

4.3. XLMTM Trial

XLMTM is a very rare congenital centronuclear myopathy caused by mutations in the MTM1 gene, affecting 1/50,000 boys [236]. Skeletal and respiratory muscles are deeply affected, and many patients decease before one year of age, mainly from respiratory failure. The MTM1 gene encodes a lipid phosphatase, myotubularin, involved in PI3P dephosphorylation and membrane remodelling [237,238]. The myotubularin cDNA, together with the regulatory elements, can be packaged in an AAV, and two very good murine and canine models of the disease recapitulate the main features of the pathology, noticeably, histological defects specific to centronuclear myopathies, generalised muscle hypotrophy and weakness, and lifespan reduction [239–241]. With these tools in hand, Buj-Bello and collaborators established a very convincing proof of concept, first by intramuscular injection with an rAAV2/1-CMV-mtm1 product [242], and next using the whole-body delivery of an rAAV2/8-Des-mtm1 product in mouse and canine models [163]. In both models, a single intravenous injection of a dose of ≈3 × 10^{13} vg/kg led to an important improvement of muscle and respiratory functions, and survival was largely extended. Importantly, therapeutic effects were also observed, though to a lesser extent, in older mice, showing that pathology reversal, essential in patients presenting the symptoms at
birth, could be achievable. In 4-year-old, long-term survivor dogs, gait, respiratory and neurological functions remained comparable to the ones of wild-type, age-matched controls, despite a progressive decline in the vector copy number in muscles, which reached a plateau after three years of age, and a diminution of muscle force [243]. A dose study carried out in the canine model established the dose-dependency of the therapy, with a significant correction achieved from $2 \times 10^{14}$ vg/kg, a quasi-normalisation of the phenotype at $5 \times 10^{14}$ vg/kg and no significant side-effects, apart from the expectable humoral immune response towards the vector and a thickening of the heart septal wall without functional consequences [244]. In this protocol, muscle expression defects evidenced by a transcriptomics approach were corrected by the mid-dose of $2 \times 10^{14}$ vg/kg [245]. Considering that the doses reversing the pathology are in the $10^{14}$ vg/kg range and challenge vector production, an additional efficacy study was carried out in three infant NHPs [246]. Eight weeks after intravenous injection, a dose of $8 \times 10^{14}$ vg/kg did not lead to significant treatment-related adverse events and produced MTM1 protein expression at levels 8- to 20-fold higher than endogenous levels in target skeletal muscles [246]. Importantly, despite a high vector copy number in the liver, the myotubularin protein level remained normal, and serum markers of liver damage did not peak significantly. Altogether, these results led to the initiation of a clinical trial in 2017 on XLMTM infants. The ASPIRO phase 1/2 trial aims at treating ventilatory-assisted patients aged less than 5 years with ascending doses $(1 \times 10^{14}$ vg/kg or $3 \times 10^{14}$ vg/kg) of an rAAV2/8-Des-hMTM1 vector (AT132 product, Audentes Therapeutics [San Francisco, USA], NCT03199469). Until very recently, the results were strikingly positive. To date, twenty-three patients have been treated, six at $1 \times 10^{14}$ vg/kg and 17 at $3 \times 10^{14}$ vg/kg; the CHOP-INTEND (Children’s Hospital Of Philadelphia INfant Test of Neuromuscular Disorders) has improved by various levels, the limb and trunk strength have increased, and new developmental skills have been achieved, such as controlling head movement, rolling over or sitting unassisted [247–249]. Respiratory function has improved significantly resulting in patients being weaned off ventilators completely. One SAE possibly related to the product occurred and was resolved by a course of intravenous steroids and supportive care. However, since the 5 May 2020, three patients treated with the highest dose have died. All three patients had progressive liver dysfunction characterised by hyperbilirubinemia starting a few weeks after dosing. Preliminary findings suggest that two children died from sepsis and one from gastrointestinal bleeding. The FDA put the trial on hold on the 29 June [189]. This tragic event remains hard to rationalize, as 14 out of 17 children treated with the high dose have not developed complications to date. The common features of the three deceased children were an older age (the boys were at the higher end of the age cut-off), a heavier weight and a pre-existing hepatobiliary disease of an unknown severity, although one can assume it to be mild, as hepatic disorders were an exclusion criterion. This condition might have facilitated liver toxicity due to the large doses of vectors. This toxicity is reminiscent of the one observed in NHPs [200,201], and the activation of complement through the formation of vector–antibody complexes, which have been implicated in lethal systemic inflammation with an adenovirus vector [250], has been hypothesized [251]. Of note, some children dosed at $1 \times 10^{14}$ vg/kg also had pre-existing liver disorders and did not develop the complications, despite being years out from treatment.

5. Improvement of the Therapeutic Toolbox

5.1. Towards Safer Next-Generation Muscle and CNS-Restricted AAVs

It is becoming increasingly evident that AAVs should be chosen carefully for every clinical application, considering specificities such as the patient’s genetic background, age, disease progression, sex, immunological state and targeted tissues. Capsid engineering is commonly used to develop safer next-generation AAV variants. These methods rely either on rational design in which capsids are tailored by targeted modifications, or on directed evolution, consisting of recovering new capsids from randomly generated high-complexity libraries after selective pressure on a tissue of interest.
For neuromuscular disorders, the improvement of muscle transduction; reduction of off-targeting, especially in the liver; and development of vectors escaping the immune response are major endeavours.

AAV2.5, obtained by replacing five residues in the AAV2 capsid with corresponding orthogonal residues of AAV1 [146] and several other variants generated by variable combinations of 32 capsids’ amino acids [252], improved muscle transduction compared with parental serotype 2 or 1 but were not assessed for whole-body distribution. Three AAV2 variants, AAV2i8, a chimeric capsid obtained by replacing a receptor-binding hexapeptide motif in the AAV2 capsid with corresponding residues in the AAV8 capsid [253], and two variants obtained by peptide insertions in a hypervariable loop [254,255] showed equivalent or improved targeting in skeletal muscles, with an important reduction in the liver in comparison with AAV2. AAV2i8 was also shown to be less likely to be serum-neutralized than the parental capsid [253]. The ratio of skeletal muscle/liver transduction was also better than for AAV9 in mice [256] but not in NHPs [257]. An additional insertion of a galactose-binding footprint on AAV2i8 did not improve the ratio further in mice [256].

Three other variants proved even more efficient than AAV9 for improving the muscle/liver transduction ratio: (1) AAV-9.45 is an AAV9 variant obtained by the random integration of amino acids and showing reduced liver expression and identical muscle and heart transduction when compared with AAV9 [258]. (2) AAVpo1 is a natural pig isolate that transduces muscles and the heart to a slightly lower level than AAV9 but presents the advantage of being completely detargeted from the liver [259]. (3) AAV-B1 is a chimeric AAV isolated from a shuffled library consisting of 11 parental serotypes and displaying reduced liver transduction and at least 10-fold higher muscle and CNS tropism than AAV9 [260].

A series of mutations on surface phosphorylatable residues of the AAV1 and AAV9 capsid improved vector stability and led to 3 to 10 times lower transduction in the liver than in muscles [261]. AAVM41 was isolated from a chimeric AAV1 and AAV9 capsid’s shuffled library and reduced both skeletal muscle and liver targeting while preserving heart transduction compared with AAV9, suggesting that this serotype could be of interest for rescuing cardiac pathologies [262,263]. Tyrosine-specific modifications of the AAV6 capsid can improve vector muscle entry [264].

Several variants demonstrated interesting characteristics regarding immune evasion. Bat AAV serotype 10HB transduced muscle with a higher muscle/liver ratio than primate AAV and showed a reduced sensitivity to antibody neutralisation [265]. A method consisting of purifying new AAV2 variants by rabbit antibody-specific affinity chromatography resulted in the identification of several antibody-resistant clones, though neither the relevance to human sera nor variant biodistribution were assessed [266]. The AAV1 variant CAM130 isolated through multiple rounds of neutralizing-antibody escape from several species evaded neutralizing antibodies, even at high concentrations, in mice, NHPs and human sera, while maintaining the tissue tropism of the parental AAV1, suggesting it could be suitable for clinical trials in large populations, as seropositivity is a common exclusion criterion [267]. Finally, after applying the double selection of variants resistant to human-serum neutralization and selected after local muscle transduction, the AAV mutant MuSi2 was isolated and showed immune response escape together with the preservation of muscle tropism, although transduction was largely reduced by the intravenous route in comparison with AAV9 [268]. Considering that this variant transduces muscle differently according to the route of administration, low vascular permeability was hypothesised. Future protocol improvement could aim at selecting new variants after intravascular injection.

Altogether, these new vectors have the potency to improve targeting efficiency and reduce the off-target effects and immune response. Their respective characteristics are summarised in Table 4.
Table 4. Next-generation recombinant AAVs for improved muscle targeting, reduced liver targeting and/or a reduced immune response.

| Reference | AAV Name | Parental AAV | Method                                                                 | Compared with | Receptor | Muscle Transduction | Heart Transduction | Liver Transduction | Immune Response |
|-----------|----------|--------------|----------------------------------------------------------------------|---------------|----------|---------------------|---------------------|-------------------|-----------------|
| [253]     | AAV2i8   | AAV2         | Rational design: replacement of receptor-binding hexapeptide with corresponding residue in AAV8 | AAV2/AAV8     | Not HS   | =AAV8 >AAV2         | =AAV8 >AAV2         | <AAV2 and AAV8 (40-fold lower) | Lower cross reactivity to AAV2 antibody |
| [257]     |          |              |                                                                      | AAV9 (rhesus monkey) |         | <AAV9 (122-fold lower) | <AAV9 (46-fold lower) | <AAV9 (11-fold lower) | ND              |
| [256]     | AAV2i8G9 | AAV2i8       | Rational design: graft galactose-binding footprint of AAV9 in VP3 AAV2i8 | AAV2i8/AAV9    | Not HS   | Glycan =AAV2i8 >AAV9 | AAV2i8 < AAVal8 < AAVal9 | AAVal8 < AAVal8G9 < AAVal9 (5-fold lower) | ND              |
| [146]     | AAV2.5   | AAV2         | Rational design: AAV2 capsid with 5 mutations from AAV1             | AAV2          | HS       | >AAV2 (2- to 5-fold) | NA                  | NA                | -No cellular immune response to capsid. -Lower cross reaction to AAV2 NAb |
| [254]     | AAV2.5   | AAV2         | Rational design: insertion of muscle-targeting peptide in AAV2 capsid | AAV2          | Not heparin | ≥AAV2 (2-fold) | >AAV2 (7-fold) | <AAV2 (2.5-fold) | ND              |
| [258]     | AAV9.45  | AAV9         | Directed evolution: random mutagenesis of surface-exposed regions of AAV9 | AAV9          | ND       | =AAV9 <AAV9         | =AAV9 <AAV9 (10- to 25-fold lower) | <AAV9 (≈140-fold) | ND              |
| [259]     | AAV pol  | NA           | Natural pig isolate                                                | AAV9          | ND       | <AAV9 (≈24-fold) | <AAV9 (≈3-fold) | <AAV9 (≈140-fold) | ND              |
| [261]     | AAV9-V731F | AAV9   | Rational design: tyrosine mutations Other mutants (no AAV of reference) | AAV5          | ND       | >AAV5 (1.5-fold) | <AAV5 (10-fold) | <AAV5 (≈125-fold) | No pre-existing immunity No cross-neutralisation by antisera against all common AAVs |
| [255]     | AAV2-VNSTRLP | AAV2 | Directed evolution: from AAV2 display peptide library with in vitro selection for heart tropism | AAV2          | ND       | =AAV2 <AAV9         | >AAV2 (10-fold) | <AAV2 (≈10-fold) | <AAV2 (≈10-fold) |
| [263]     | AAVM41   | AAV1/6/7/8  | Directed evolution by shuffling the capsids of AAV1 to AAV9 and in vivo selection on skeletal muscle | AAV6          | ND       | <AAV6 >AAV9         | >AAV9 (up to 13-fold) | <AAV9 <AAV6 | Lower cross reactivity |
| [260]     | AAVB1    | AAV8        | Directed evolution of DNA shuffled library and selection on brain tissues | AAV9 Not SA | Not Galactose | >AAV9 (10- to 26-fold higher) | >AAV9 (14-fold higher) | <AAV9 (3.6-fold higher) | Modestly more resistant to neutralisation than AAV9 |
| [252]     | AAVC4    | AAVC7       | Ancestral reconstruction from NHP and human AAV by combinatorial variation of 32 amino acids and selection on muscle cells | AAV1 Not SA | Not galactose | >AAV1 (10-31-fold higher) | NA | NA | Not resistant to neutralisation with IVIG |
Table 4. Cont.

| Reference | AAV Name | Parental AAV | Method | Compared with | Receptor | Muscle Transduction | Heart Transduction | Liver Transduction | Immune Response |
|-----------|----------|--------------|--------|---------------|----------|---------------------|--------------------|--------------------|-----------------|
| [265]     | AAV10HB  | NA           | Isolation from bat faecal and intestinal tissues | AAV2 AAV8 ND | ND | Ratio muscle/liver = 8.8 | Ratio muscle/liver < 1 for AAV2 and AAV8 | Reduced neutralisation with IVIG = AAV2 |
| [266]     | Several variants: r2.4/r2.15 AAV2 | | Directed evolution: random mutagenesis and selection of heparin binding or neutralising serum binding | AAV2 Not heparin ND ND ND | Reduced neutralisation with serum/AAV2 |
| [268]     | Musi2 | Capsid shuffled library | Directed evolution: shuffled library selected on patients’ sera and amplified in vivo in mouse muscle | AAV1/AAV2/AAV2.5/AAV6/AAV8/AAV9 ND | IM = AAV6 = AAV9 IV < AAV9 | | | Immune escape |
| [267]     | CAM130 | AAV1 | Directed evolution: rational mutagenesis on AAV1 capsid residues in contact with antibodies, library generation and evolution on vascular endothelial cells | AAV1 ND ND | >AAV1 (2-fold) =AAV1 | Neutralisation escape to murine, NHP and human sera |

IVIG: intravenous human IgG; HS: heparan sulfate; SA: sialic acid; NA: not applicable; ND: not determined.
5.2. Enhancing the Repertoire of Muscle and CNS-Restricted Promoters

With the development of in silico analysis technologies, a multistep, genome-wide data-mining strategy was performed to identify conserved skeletal muscle-specific cis-regulatory modules (Sk-CRMs) in highly expressed muscle-specific genes. Sk-CRM4, containing binding sites for the E2A, CEBP, LRF, MyoD and SREBP transcription factors, boosted transgene expression driven by Des or C5.12 promoters in heart and skeletal muscles (up to 400-fold), with a significant improvement of the mdx mouse phenotype [269]. Similarly, a 1030 bp modular muscle hybrid (MH) promoter composed of two enhancers (from the Des and Mck genes, respectively), a proximal promoter and an intron (modified from the Mck gene and core promoter) was more efficient than the Des promoter in skeletal and cardiac muscles, with limited expression in non-muscle tissues compared with the CMV promoter, showing a high potential for muscular gene therapy [270].

The presence of large promoters limits the size available for the transgene in the cassette, which proves problematic for several muscle genes. Promoterless cassettes were recently tested for liver expression. In this strategy, the transgene, flanked by homology arms, is brought into the cells by rAAV and integrates by nuclease-free homologous recombination downstream of the native promoter, where it is regulated like the endogenous gene is. Despite promising results in hepatocytes [271,272], the promoterless strategy might be limited for muscle application, as muscle cells are mostly quiescent and homologous recombination is restricted. Nonetheless, it might prove interesting for satellite cell targeting, provided that muscle progenitor targeting can be achieved.

5.3. Detargeting with miRNA-Based Elements

While regulatory elements such as introns, polyA signals or the Woodchuck hepatitis virus post-transcriptional regulatory element (WPRE) can be added to improve global transgene expression, miRNA-based sequences can mitigate tissue-specific transgene expression [273]. MiRNAs are small (approximately 22-nucleotide-long) non-coding RNAs post-transcriptionally silencing gene expression in plants and animals. Once bound to complementary target sites (TS) in mRNA, they either reduce its stability or inhibit its translation, which results altogether in the reduction of protein expression [274]. While the number of identified miRNAs has constantly increased since their discovery in Caenorhabditis elegans in 1993 [275], the miRBase database reports 1917 annotated hairpin precursors and 2654 mature sequences in the human genome [276]. Some miRNAs present a tissue-specific pattern of expression, with expression detectable only in a particular tissue or at least 20-fold higher than in other tissues [277]. Amongst tissue-specific miRNAs are found the MyomiRs, a family of miRNAs expressed in both cardiac and skeletal muscles, namely, miR-1, miR-133a, miR-122a, miR-124a, miR-208b, miR-499 and miR-486, with the exception of miR-208a and miR-206, which are specifically expressed in the heart and skeletal muscles, respectively [278].

One strategy to improve the specificity of AAV-mediated gene delivery, overriding the broad tissue tropism of AAV vectors and/or promoter leakage in non-targeted tissues, is based on the miRNA-mediated post-transcriptional regulation of the transgene. Indeed, the insertion of miRNA TS into the 3′UTR of a gene expression cassette limits transgene expression in tissues expressing the corresponding miRNA [279]. Due to the small size of miRNAs, it is therefore feasible to insert different miRNA TS in the 3′UTR of the expression cassette to detarget specific cell types depending on the application. For neuromuscular disorders, this strategy was applied for the reduction of expression in the heart [280], liver [281,282] and APCs [283–287].

The control of heart transgene expression is of utmost importance, because even if no specific cardiac toxicity has been reported to date in clinical trials, preclinical reports have evidenced the danger of transgene cardiac overexpression when the heart is not the primary target [280] or even when it is [288]. The insertion of the cardiac-specific miR-208a TS in the cassette was shown to prevent cardiac transgene expression and rescued the cardiac toxicity resulting from transgene overexpression in this organ, while maintaining the efficient expression of the transgene in skeletal muscles [280].
MiR-122 is highly expressed in the liver. The insertion of miR-122 TS in the 3′UTR of a reporter gene was able to prevent protein expression in the liver after rAAV9 intravenous administration without interfering with cardiac protein expression [281,282]. The level of transgene repression was related to the number of repetitive miRNA TS used. However, the recent paper of Kraszewska et al. challenges this apparently safe approach. Indeed, in some genetic backgrounds, transgene expression was completely repressed not only in the liver, but also in the cardiac muscle, linked with the presence of miR-122 in these animals’ hearts. MiR-122 was also shown to be present in the human cardiac tissues of patients with cardiomyopathy and in human iPSC-derived cardiomyocytes. The cardiac expression showed high variability between different mouse strains, sexes and human individuals [289]. This publication challenges the liver-specificity of miR-122 and warns against miRNA inter-individual variability.

As previously mentioned, preventing transgene expression in APCs may avoid undesirable adaptive immune responses directed against the transgene product. A miR-based approach aiming at inhibiting transgene expression in APCs by inserting four targets of the endogenous miR-142-3p (exclusively expressed in the hematopoietic lineage) at the 3′ end of the transgene coding sequence [283] allowed escaping a deleterious adaptive immune response after gene delivery with either lentiviral [284] or AAV vectors [285–287].

Most importantly, it is crucial to verify during preclinical studies that miRNAs are not reduced by the miRNA TS and that their natural targets are not misregulated, as that could induce detrimental side effects. To our knowledge, no clinical trial has used miRNA TS in the cassette to date. If it ever happens, checking beforehand the mean level of the targeted miRNA in the treated population will be essential, as miRNA expression can substantially vary between individuals, sexes or pathologies [289–291].

6. From Preclinical Studies to Clinical Trials… and Back: General Point of View

The ongoing clinical trials summarised herein show spectacular results in terms of efficacy, especially in SMA and XLMTM, two very severe conditions characterised by generalised muscle weakness and respiratory deficiency often leading to infant deaths in the first years of life. The DMD trials need further investigation. However, this very beautiful landscape has lately been obscured by SAEs in two DMD and in the XLMLM trials, leading to three children’s deaths in the last case. It is hard to find common features in the two situations, as, apart from the doses, which are very close, the genetic background, the age of the patients and the vector capsids are different. DMD-related adverse events have been proposed to be caused by adverse immune reactions driving acute kidney failure, while XLMTM fatal hepatotoxicity is not associated with obvious immunotoxicity. Future investigations will undoubtedly document these side-effects and help with the design of next-generation products, but for now, with the current state of our knowledge, a lot of effort has to be put into designing the safest therapeutic strategies for future trials, especially as some diseases are not prone to being good candidates in terms of the benefit/risk ratio. Several factors have to be considered in the “ideal” trial design (see Figure 1).
Figure 1. Of rAAV and men: the ideal journey.
6.1. Defining the Product

The transgene definition is obviously central and often evident, except in large proteins, where shorter forms have to be assessed with extreme care in preclinical studies to choose the best product possible. However, improvements can be made even with a full-length transgene. For example, codon optimisation leading to better protein expression has proven beneficial [196,292]. A thoughtful choice of promoter and other regulatory elements, such as WPRE addition, which proved efficient in enhancing transgene expression [293], can also improve the efficacy of the products. As for capsids, the choices are widening deeply, with new attractive vectors improving specific targeting. However, these vectors have to be assessed more closely in terms of safety before they can be considered for clinical usage. All these choices are crucial for specific tissue targeting and the reduction of off-target effects. Fundamentally, a very good knowledge of the levels of the protein to replace in every tissue of the healthy population is necessary to design the best targeted product. The addition of miRNA TS to detarget the liver [281], heart [280] and APCs [294] proved efficacious in reducing ectopic expression, but has not been tried in neuromuscular-deficient patients to date, probably because the underlying risks have to be assessed more closely. It is also worth mentioning that even with perfect targeting, the overexpression of transgenic protein in the targeted organs can also lead to toxic effects [295].

6.2. Manufacturing AAV

Vector production is a critical process for gene therapy success and safety. Importantly, the methods used for vector production and titration are not standardised, complicating the comparability of different clinical trials. Indeed, it was previously shown that the same production of rAAV8 led to significant variations in titres when dosed in 16 different laboratories [296]. In the absence of consensual methods, a common standard used for all clinic-intended rAAV production could help to correct the titres amongst trials.

Moreover, according to the method of production/purification used, various quantities of toxic contaminants can be found in AAV production. Indeed, endotoxins are known to be able to activate the human immune system and lead to SAEs and often contaminate AAV production. It is therefore crucial to reduce their load in the final product [297]. Their safety limit is defined as 5.0 International Units of endotoxin per kilogram of body mass by the FDA and the European Pharmacopeia for intravenous usage in humans [298,299], but could be raised further depending on the patient status (age, disease severity, etc). Of note, other contaminants, not known or tested, could also play a role in the safety and/or efficacy of the product. Notably, the presence of empty capsids in the final product was shown to reduce transduction efficiency and may participate in side effects [300].

6.3. Choosing the Best Preclinical Models

While proof-of-concept and preclinical studies aim at determining the minimal effective dose of the product, toxicology studies evaluate its safety/toxicity. Although using relevant animal models mimicking the human disease in proof-of-concept and preclinical studies seems obvious, the choice of animal models in toxicology studies remains unclear. The assessment of chemical drug toxicity is traditionally performed on wild-type animals such as rats, dogs or monkeys, as they are relevant for phase 1 clinical trials aiming at determining the safety in the general population. However, gene therapy cannot ethically be tested in healthy individuals. The crucial question is the relevance of extrapolating toxicology findings to human clinical trials. Vector entry relies on cell surface receptors and co-receptors, and its internal traffic requires components of the host cells. As these processes are not well deciphered in humans, inter-species variability could preclude toxicity results in some species. In line with this idea, piglets did not show liver failure and haemorrhage when administered the same dose of rAAVhu68 as NHPs [200]. Additionally, NHPs are not necessarily the best species as assumed, since toxicology studies in XLMTM at $8 \times 10^{14}$ vg/kg did not detect coagulation defects or acute toxicity [246], while three children died at $3 \times 10^{14}$ vg/kg. This is not completely surprising,
as membrane and cytoskeletal remodelling likely alter vector processing in neuromuscular diseases, possibly significantly modifying AAV efficacy. In line with these considerations, the International Council for Harmonisation of Technical Requirements for Pharmaceuticals for Human Use (ICH) defines a relevant species as “one in which the test material is pharmacologically active” and states that if “no relevant species exists, the use of relevant transgenic animals expressing the human receptor or the use of homologous proteins should be considered. [. . .] In certain cases, studies performed in animal models of disease may be used as an acceptable alternative to toxicity studies in normal animals” [299].

6.4. Defining the Dose

For intravenous administration, the AVV dose administered is proportional to body weight, regardless of the age, gender, genetic background and disease severity of the individual treated. However, any of these factors could influence AAV efficacy and toxicity. The relative weight of organs is not proportional to body weight during development. For example, the liver/body ratio is higher in children than in adults [301], which could lead to variable levels of transduction and influence vector biodistribution at different ages. The immune systems of young children are not fully mature [235] and could therefore facilitate AAV transfer. Although it is not the only difference, the two DMD trials showing toxicity events potentially linked to immune responses were performed in older patients than the trial without any SAEs. Sex was also shown to impact AAV transfer in hepatic tissue but not in other tissues, with the male liver being more transduced than females’ [302]. Finally, the disease itself can modify the structure and function of various organs, with high variability between individuals of the same age. Indeed, in the XLMTM trial, the three deceased patients had pre-existing hepatobiliary diseases and were older than the other infants without SAEs treated at the same dose. Although it is challenging, finding a better and universal normalisation method for dose calculation could probably improve clinical trial standardisation and safety.

6.5. Circumventing Immune Response

As previously mentioned, the adaptive immune response directed against a viral-derived vector restricts the full therapeutic potential of in vivo gene therapy [303]. Thus, in the first clinical trial showing safe and efficacious liver targeting with an rAAV2 vector carrying the human factor IX transgene (under the control of a liver-specific promoter), transgene expression was only transient [109]. A decline in expression starting at four weeks was associated with transient liver transaminases and the detection of CD8+ T cells directed against the AAV2 capsid. This unexpected deleterious cellular immune response was vector-dose-dependent and, in the absence of preclinical animal models, is still poorly understood. Nevertheless, a short prednisolone treatment quickly given in response to liver injury is often sufficient to stabilise transgene expression and has been used since then [230,231]. As a result, anti-AAV neutralizing antibodies are one of the most important remaining barriers, either impairing the efficacy of gene transfer in a set of patients with a cross-reactive pre-existing immunity against wild-type AAV, or precluding the redosing of patients developing a rapid and strong humoral response after the first vector injection. In preclinical models, numerous strategies targeting the host have been proposed such as plasmapheresis [304], direct tissue injection or isolated organ perfusion and immunosuppression combining rituximab (anti-CD20 depleting monoclonal antibody) and others drugs, or synthetic particles encapsulating rapamycin [305,306]. Other strategies target the rAAV vector itself, such as the use of alternative and less-prevalent serotypes, empty decoy capsids [307], exosome-enveloped AAV vectors or the generation of novel AAV capsids with optimised biodistributions and transduction efficacy, as well as the capacity to evade NAbs, as discussed above [47,305]. The most promising approach to including patients non-eligible to date was recently reported with the use of an IgG-cleaving endopeptidase from Streptococcus pyogenes (IdeS) [308]. The IdeS enzyme very rapidly (in a few hours) cleaves human IgG into F(ab′)2 and Fc fragments, and is safe and efficient in patients with donor-specific antibodies undergoing kidney transplantation [309]. In both mouse and NHPs, Leborgne et al. reported that IdeS treatment was able to decrease pre-existing
anti-AAV antibodies to a level sufficient to enable efficient liver gene transfer, even in the setting of vector re-administration [308]. Equivalent properties were demonstrated with IdeZ, a homolog of IdeS [310]. In July 2020, Sarepta Therapeutics announced an agreement with Hansa Biopharma to develop and promote imlifidase (the commercial IdeS) as a pre-treatment for DMD and LGMD gene therapy.

6.6. Assessing Long-Term Efficacy

Since the clinical trials using whole-body delivery in neuromuscular diseases are quite recent, the long-term assessment of efficacy will be made available in the next few years. For CNS-targeted treatments, a relative stability of the treatment is to be expected, as neurons are the longest living cells of the body. However, muscles, while being in a post-mitotic state, are remodelled during growth and following exercise, which could dilute the therapeutic effect. Targeting the treatment to satellite stem cells could hence be useful to help maintain long-term efficacy. Unfortunately, AAV-driven attempts to target satellite cells have failed. While reinjection might prove complicated, it might be worth using lentiviral vectors to target stem cells, as it was demonstrated efficient in transducing satellite cells in vivo [311], or new rAAV-rDNA integrating vectors, which proved their efficacy for directed integration in dividing and quiescent cells [312]. Importantly, it was recently shown that the AAV virus is found in episomal and randomly integrated transcriptionally active forms in human samples of liver tissues, and while it is impossible to know the time of infection, considering the large number of samples, it certainly suggests long-term persistence of the virus [89]. Whether this is also true for rAAV genomes remains to be determined.

6.7. Assessing Long-Term Toxicity

Serious concerns about the long-term safety of rAAV vectors were raised after several genotoxic studies performed in mice. Recombinant AAV2, 8 and 9 vectors were shown to integrate in the Rian locus on chromosome 12, irrespective of viral transgene, mouse genotype, sex or genetic background [104,105]. This insertion upregulates non-coding RNA and genes proximal to the Rian locus and is associated with an increased rate of hepatocellular carcinoma (HCC). The trans-regulatory elements carried by the vectors influence genotoxicity: sometimes, insertion is seen, but adjacent oncogenes are not overexpressed and HCC does not develop. Capsid-specific properties may also influence genotoxicity. These results were all obtained in neonatal mice, and neither integration in the Rian locus nor HCC were ever observed in older animals [106]. Other results obtained with sc vectors evidenced insertion within proto-oncogenes injected in young adult mice [313]. As no liver tumours have been seen to date after rAAV treatment in humans, the risk of insertional mutagenesis is probably very low, if it ever exists. Nonetheless, patients should be followed longitudinally to monitor long-term effects.

Figure 1 summarises the main steps necessary to push forward an AAV-based gene therapy medicinal product from preclinical studies to clinical trials

Without minimising the importance of the tragic toxic events seen in the current clinical trials, it is worth emphasizing that AAV-mediated gene therapy is the only treatment that led to highly significant disease improvement in severely affected human patients. It is now necessary to go back to bench work in order to decipher the pathogenic mechanisms underlying AAV-linked toxicity and design safer next-generation therapeutic cassettes. Indeed, AAV therapy remains the main source of hope for patients affected by neuromuscular disorders, and there are many more diseases to treat. Importantly, our laboratory is planning a new rAAV-based clinical trial using micro-dystrophin transfer in DMD patients, in partnership with Sarepta [314]. The baseline study, aiming at collecting data on the natural disease course in DMD male subjects aged from 5 to 9 years of age, is currently ongoing (GNT-014-MDYF, NCT03882827), and the interventional gene therapy trial should start in early 2021.

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**Abbreviations**

NHP: non-human primate; SMA: spinal muscular atrophy; DMD: Duchenne myopathy disease; XLMTM: X-linked myotubular myopathy; CNS: central nervous system; AAV: adeno-associated virus; SMN: survival motor neuron; FDA: Food and Drug Administration; EMA: European Medicines Agency; SAE: serious-adverse event; miR: microRNA; PMO: phosphorodiamidate morpholino oligomer; ITR: inverted-terminal-repeat; LGMD: limb-girdle muscular dystrophy; NAbs: neutralizing antibodies; Sk-CRMs: skeletal muscle-specific cis-regulatory modules; WPRE: woodchuck hepatitis virus post-transcriptional regulatory element; APC: antigen presenting cells; sc: self-complementary.

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