Amyotrophic lateral sclerosis is a progressive and devastating neurodegenerative disease. Despite decades of clinical trials, effective disease-modifying drugs remain scarce. To understand the challenges of trial design and delivery, we performed a systematic review of Phase II, Phase II/III and Phase III amyotrophic lateral sclerosis clinical drug trials on trial registries and PubMed between 2008 and 2019. We identified 125 trials, investigating 76 drugs and recruiting more than 15 000 people with amyotrophic lateral sclerosis. About 90% of trials used traditional fixed designs. The limitations in understanding of disease biology, outcome measures, resources and barriers to trial participation in a rapidly progressive, disabling and heterogenous disease hindered timely and definitive evaluation of drugs in two-arm trials. Innovative trial designs, especially adaptive platform trials may offer significant efficiency gains to this end. We propose a flexible and scalable multi-arm, multi-stage trial platform where opportunities to participate in a clinical trial can become the default for people with amyotrophic lateral sclerosis.
Clinical trials in Amyotrophic Lateral Sclerosis

Biological understanding
Lack of sensitive outcome measures and biomarkers
Challenges in drug selection

Disease factors
Rapidly progressive
Disabling
Heterogenous

Challenges

Trial design
Restrictive criteria
Restrictive protocols
Limited sample sizes
Limited resources

Opportunities for future trials
Innovative trial designs
Advances in biological understanding
Better outcome measures

Efficient
Flexible
Scalable
Inclusive
Accessible
Introduction

Amyotrophic lateral sclerosis is a rapidly progressive disease with a median survival of 2–3 years. While there has been some progress in non-pharmacological interventions, such as non-invasive ventilation and gastrostomy, and symptomatic pharmacological treatments in amyotrophic lateral sclerosis, trials in the last 25 years have largely failed to identify effective disease-modifying drugs. Riluzole, approved in 1995, is the only globally licenced disease-modifying drug and prolongs survival by just 2–3 months on average. Edaravone has been approved in some countries including USA, Canada, Japan and South Korea following a positive trial in a highly selected population. Sodium phenylbutyrate-taurisodiol (AMX0035), masitinib and methylcobalamin have recently emerged as promising candidates following trials in subsets of people with amyotrophic lateral sclerosis (pwALS). However, evidence for generalizable and substantial effects on survival for edaravone, AMX0035, masitinib and methylcobalamin is limited.

Limitations of pre-clinical models and incomplete knowledge of the underlying biology of amyotrophic lateral sclerosis are some of the factors hindering translational success. However, the last decade has seen substantial progress in our understanding of the genetic and molecular pathobiology of amyotrophic lateral sclerosis and related disorders. This improved mechanistic insight has, in turn, led to the identification of many promising therapeutic targets that justify not only further experimental study but also, in many instances, formal clinical trials. Given that at present only a small fraction of pwALS participate in clinical trials, despite a prognosis similar to that for many cancers where trial participation rates are considerably higher, there is an opportunity to learn from innovations in cancer medicine. This includes re-thinking historical approaches to drug selection and evaluation in standalone two-arm trials.

Here, we evaluate previous trial designs and consider specific methodological challenges. These include the rapid pace of decline and accumulating disability in amyotrophic lateral sclerosis, the clinically heterogeneous nature of amyotrophic lateral sclerosis and a range of trial design and conduct issues. We review emerging innovative trial designs and propose a flexible multi-arm, multi-stage trial platform model that can incorporate new candidate drugs as they are identified.

Materials and methods

We conducted a systematic search of trial registries including clinicaltrials.gov, World Health Organization International Clinical Trials Registry Platform (ICTRP), European Union Clinical Trials Register (EU CTR), and PubMed on 9 April 2019 to identify Phase II, Phase II/III and Phase III Clinical Trials of an Investigational Medicinal Product (CTIMPs) assessing potential disease-modifying drugs in amyotrophic lateral sclerosis. We searched clinicaltrials.gov for all interventional trials of ‘amyotrophic lateral sclerosis’ or ‘motor neuron disease’. We searched ICTRP and EU CTR for interventional trials of ‘amyotrophic lateral sclerosis’ using ‘Phase II’ and ‘Phase III’ filters. We searched PubMed for ‘(“motor neuron disease” OR “amyotrophic lateral sclerosis”) AND (Clinical Trial [ptyp])’. Trials registered, completed or published between 1 January 2008 and 9 April 2019, excluding extension trials, were included. No language restrictions were applied.

Statistical analysis

We performed a narrative synthesis on all included trials to summarize trials according to phase, study status, study duration, number of participants, investigational medicinal products (IMPs), eligibility criteria and primary outcome measures. For studies with entries on multiple registries, data from ClinicalTrials.gov were used as the primary data source. For studies registered on registries with results published in journals, publications were used as primary source of data. To review sample size considerations including recruitment and retention, we summarized trials for completed trials with efficacy-based primary outcome measures according to number of study arms, number of participants recruited, number of participants recruited per arm, withdrawal rates and reasons for withdrawal.

Data availability statement

Data for this review are available in the Supplementary material.

Results

Characteristics of trials

A total of 1152 records were identified [see Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) diagram in Fig. 1]: 344 were duplicates, and 683 were excluded because they did not meet our eligibility criteria. Total of 58 records on PubMed corresponded with trials identified from the registry search. In total, 125 trials, with a total of 15 647 participants, were identified and included in the analysis. Characteristics of these trials are summarized in Table 1. A total of 73 trials were recorded as completed. Only 6 of the 125 trials were terminated prior to planned completion: 2 of these were trials with novel designs terminated for futility; 1 was terminated due to adverse events; 1 was terminated after a participant had rapid progression; the reasons for termination for two studies were not stated. Trial duration ranged between 4 and 95 months (median 25 months).
Eligibility criteria
About 73% of trials specified amyotrophic lateral sclerosis disease duration within their inclusion criteria. Median permitted maximum disease duration was 36 months [interquartile range (IQR) 24–36 months]. Amyotrophic lateral sclerosis participants at King’s stage 4 (requiring enteral and/or respiratory support) were excluded from 119 trials. Total of 53 trials stipulated use of riluzole as part of their inclusion or exclusion criteria.

IMP s
A total of 76 IMPs were evaluated in these 125 trials (Supplementary Tables 1 and 2), of which 10 IMPs were tested in 3 or more trials. The most frequently tested intervention was lithium, tested in 10 trials (6 Phase II, 2 Phase II/III and 2 Phase III), involving a total of 1414 pwALS. A total of 82 trials used oral forms of IMPs, but only 7 of these specified forms, such as liquid, soluble and powder preparations which could be delivered through enteral feeding tubes, an important consideration raised in engagement with pwALS. Two IMPs have been licenced since riluzole was approved in the USA in 1995 and the European Union (EU) in 1996. Edaravone was licenced in Japan in 2015 and in the USA in 2017 after a study showed improvement in the Amyotrophic Lateral Sclerosis Functional Rating Scale Revised (ALSFRS-R, a multi-domain functional rating scale) in a small subset of pwALS characterized by early amyotrophic lateral sclerosis, functional independence with no respiratory insufficiency and a decrease in ALSFRS-R of 1–4 points during a 12-week observation period.4
patient stratification leading to loss of generalizability, aspects of study conduct and the handling of missing data. More recently, Food and Drug Administration (FDA) and EMA have granted orphan drug designation for AMX0035 following a positive Phase II trial in which AMX0035 reduced rate of decline of ALSFRS-R by 0.42 points/month over 24 weeks compared to placebo. Although there was no difference in survival between treatment groups during the primary analysis, subsequent analysis of long-term open-label extension results showed a significant improvement in survival [hazard ratio (HR) = 0.56, P = 0.023] in participants originally randomized to AMX0035 compared to participants originally randomized to placebo. However, it is notable that overall sample size was small, only 20% of participants remained in the extension trial at the time of publication, and the cohort was highly enriched using strict eligibility criteria and thus, treatment effect may not be generalizable to the wider amyotrophic lateral sclerosis population. In April 2021, FDA expressed an interest in seeing data from an additional placebo-controlled trial prior to receiving a New Drug Application for AMX0035.

In a post hoc analysis of a long-term Phase II/III randomized controlled study, ultra-high-dose methylcobalamin treated pwALS prolonged time to death or ventilation support and reduced decrease in ALSFRS-R score compared to placebo for pwALS diagnosed and entered into trial within 12 months of symptom duration. However, the primary analysis for the trial did not show any significant difference between treatment groups. Nuedexta (dextromethorphan/quinidine) was approved for treatment of pseudobulbar affect in amyotrophic lateral sclerosis in 2011 by the FDA and 2013 by EMA. Following reports from pwALS noting an improvement of their bulbar symptoms on Nuedexta, a Phase II randomized, blinded, crossover study of 60 participants was performed. Nuedexta was associated with significantly improved bulbar Function Scale (CNS-BFS), a self-reported measure of swallowing, salivation and speech, compared to placebo over a treatment period of 28–30 days. However, it is unclear if this reduction translates into a clinically meaningful effect. Longer term larger studies are required to establish its effects on disease progression. Of note, the marketing authorization for Nuedexta has been withdrawn in the EU on request of the manufacturer due to commercial reasons.

### Trial design

About 90% of the included trials used traditional fixed designs where each trial evaluated one active IMP (including dose-ranging studies testing one active IMP at multiple doses) and were designed, conducted then analysed in a consecutive manner, with no flexibility to adopt changes, which may become desirable over the course of the trial. A total of 12 trials used ‘novel’ or ‘alternative’ trial designs (Table 2). In the Airlie House amyotrophic lateral sclerosis clinical trial guidelines, any significant difference between treatment groups.

### Table 1 Phase II, Phase II/III and Phase III CTIMP assessing potential disease-modifying treatment in amyotrophic lateral sclerosis registered, completed or published from 1 January 2008 onwards excluding excluding

| Description | Number of trials |
|-------------|------------------|
| By status  |                  |
| Active     | 35               |
| Completed, has results | 61           |
| Completed, no results   | 12              |
| Not yet recruiting   | 5               |
| Terminated | 6                |
| Unknown   | 4                |
| Withdrawn | 2                |
| Total     | 125              |
| By phase   |                  |
| Phase II  | 97               |
| Phase II/III | 8              |
| Phase III | 20               |
| Total     | 125              |
| By trial design |        |
| Non-randomized, open label | 26         |
| Randomized | 96               |
| Parallel: 87  |                |
| Crossover: 7  |                |
| Factorial: 1  |                |
| Assignment not specified: I | 1          |
| Parallel, randomization not specified | 22          |
| Not specified | 2                |
| Total     | 125              |
such designs include adaptive studies, seamless Phase II/III designs, enrichment designs and futility designs.

Next, we reviewed sample size considerations including recruitment and retention. To do this, we analysed studies with efficacy-based primary outcome measures (Table 3). Results were available for 49 (32 Phase II, 5 Phase II/III and 12 Phase III) of the 89 trials with efficacy-based primary outcome measures. A total of 37 studies evaluated two arms, 5 were single-arm (including 1 using historical controls29), 6 trials evaluated three arms (5 studies of one IMP each at different doses and 1 multi-arm trial of two active IMPs) and a single trial had four arms (testing one IMP at three doses against placebo). The total number of participants recruited per trial ranged from 23 to 943. The median number of participants per arm was 43 (IQR 23–107) (Fig. 2). Only 26 of 49 trials (20 Phase II, 4 Phase II/III, 2 Phase III) recruited fewer than 100 participants in total, and only 4 Phase III trials recruited more than 500 participants. Excluding trials with adaptive designs, which were terminated early due to predefined futility criteria, 87% of the studies achieved at least 90% of their target enrolment. However, attrition significantly affected studies: 40% of trials had an attrition rate (calculated as the proportion of recruited patients who did not complete studies for reasons other than death) of more than 20% (range 0–70%, median 16%). In these studies, median percentage of participants withdrawing due to adverse events was 26% (IQR 0–45%, range 0–100%). Other reasons for withdrawal included participant request, protocol violation, loss to follow up, perceived lack of efficacy, non-compliance, disease progression felt to be due to study drug, burden of participating, physician or investigator’s decision and reaching protocol-defined stopping point other than death.

Outcome measures

Total 24 of the 49 studies used ALSFRS-R, while 10 trials used survival, and 2 used compound scores combining survival and ALSFRS-R as primary outcome measures.30,31 Other primary outcome measures included alternative functional rating scales, muscle strength testing, disease staging, respiratory function and biofluid biomarkers (Table 3). The substantial variation in primary endpoints used reflects the lack of consensus on what is considered a relevant and clinically important improvement for confirmatory trials, and the lack of sensitive and reliable outcome measures to detect potential treatment effects in exploratory trials.

ALSFRS-R scores were reported as mean rate of decline or slope of decline, or as an absolute or percentage change from baseline score, measured over durations ranging between 8 and 90 weeks. Trials using ALSFRS-R were planned to detect a range of treatment effects,

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**Table 2 Novel trial designs for Phase II, Phase II/III and Phase III CTIMPs evaluating potential disease-modifying treatment of amyotrophic lateral sclerosis**

| Novel trial designs | Completed with results | Conclusive results on efficacy/futility | Early stopping due to predefined futility criteria | Number of trials |
|---------------------|------------------------|----------------------------------------|-----------------------------------------------|-----------------|
| Multi-arm studies*  |                        |                                        |                                               |                 |
| NCT01257581: Phase II study comparing Creatine 30 mg versus tamoxifen 40 mg versus tamoxifen 80 mg | Y | N | N/A | 2 |
| NCT00355576: Phase II study comparing minocycline and creatine versus celecoxib and creatine | Y | N | N/A |  |
| Multi-phase study   |                        |                                        |                                               | 1               |
| NCT00349622: Phase I-III study of ceftriaxone | Y | Y | N |  |
| Sequential/adaptive design |                        |                                        |                                               | 6               |
| NTR 1448: Phase II study of lithium | Y | Y | Y |  |
| NCT00136110: Phase III study of sodium valproate | Y | Y | Y |  |
| NCT00690118: Phase II study of pioglitazone | Y | Y | Y |  |
| NCT00243932: Phase II study of coenzyme Q10 | Y | Y | Y |  |
| NCT00818399: Phase II/III study of lithium in combination with riluzole | Y | Y | Y |  |
| NCT02238626: Ibudilast (MN-166) in subjects with amyotrophic lateral sclerosis (IBU-ALS-1201) | N | N/A | N/A |  |
| Phase II futility design |                        |                                        |                                               | 1               |
| EudraCT 2014–005367-32: Phase II study of guanabenz | N | N/A | N/A | 1 |
| Historical controls, primarily virtual data collection |                        |                                        |                                               | 1               |
| NCT02709330: amyotrophic lateral sclerosis reversals—Lunasin regimen | Y | Y | N/A |  |
| Adaptive seamless Phase II/III |                        |                                        |                                               | 1               |
| NCT00706147: Phase II/III randomized, Placebo-controlled trial of arimoclomol in SOD1 positive familial amyotrophic lateral sclerosis | Y (closed after Phase II due to slow recruitment) | N | N |  |
| Total                |                        |                                        |                                               | 10/7 (70%) | 5/7 (71%) | 12 |

*For multi-arm studies, trials specified evaluated more than one active investigational medicinal product; studies including concomitant use of riluzole and dose-ranging studies were not included here.
including a reduction in the rate of decline of anything from 10% to 50%, or a 3-point difference in ALSFRS-R scores over 24–40 weeks.

Survival results were reported in several ways including number of deaths, proportion alive at a specified time-point, Kaplan–Meier curves, time-to-event including death or proxies for survival such as tracheostomy or the initiation of permanent assisted ventilation. Follow-up periods for survival ranged from 12 to 18 months.

Biomarkers were specified as outcome measures in 42 (34%) of trials: 27 trials included biofluid biomarkers, 3 measured imaging biomarkers and 13 measured neurophysiological biomarkers.

### Proportion of pwALS in clinical trials

To determine the approximate proportion of pwALS participating in clinical trials, we evaluated reported recruitment data for trials identified in our systematic search and compared this to amyotrophic lateral sclerosis epidemiology data (Supplementary Table 3). We identified 66 studies, which recruited in the UK and the USA and extracted the total number of participants from these countries. This was based on published data where these were available; and where not, by taking the proportion of recruiting centres in the UK or USA multiplied by the total number of participants. Applying this approach to those studies where the country of recruitment was known gave estimated recruitment within 1% (2887 versus 2916 observed) of the true value. We calculated trial participation rates using an estimated annual incidence of amyotrophic lateral sclerosis of 2/100 00032 and population data from national statistics. We estimate that fewer than 5% of pwALS in the UK and 8% of pwALS in the USA participated in a CTIMP during this period. Globally, we estimate that the 15 647 pwALS who participated in these 125 CTIMPs between 2008 and 2019 represent around 2.5% of the total number of new cases of amyotrophic lateral sclerosis during this period.

### Discussion

Our analysis of trials between 2008 and 2019 found 76 IMPs evaluated in 125 trials, predominantly with two-arm trial designs with a range of sample sizes and a variety of outcome measures. We estimate that fewer than 5% of newly diagnosed pwALS entered a clinical trial. Limitations of this review include incomplete reporting of

| Type of primary outcome measure                          | Number of trials |
|----------------------------------------------------------|------------------|
|                                                          | Phase II | Phase II/III | Phase III | Total |
| **Efficacy**                                             | 63       | 8            | 18        | 89    |
| Functional rating scales                                 | 39       | 4            | 8         | 51    |
| ALSFRS-R                                                 | 36       | 4            | 7         | 47    |
| AALSRS                                                    | 1        | 0            | 1         | 2     |
| CNS-BFS                                                   | 1        | 0            | 0         | 1     |
| Penetration aspiration scale                              | 1        | 0            | 0         | 1     |
| Combined survival/functional outcome measure e.g. Joint rank scales/CAFS | 1        | 1            | 2         | 4     |
| **Staging**                                              | 3        | 1            | 0         | 4     |
| ALS-MITOS                                                | 1        | 0            | 0         | 1     |
| Loss of self-sufficiency                                 | 1        | 0            | 0         | 1     |
| Time to tracheostomy/death                               | 0        | 1            | 0         | 1     |
| Not specified                                             | 1        | 0            | 0         | 1     |
| **Survival**                                             | 8        | 1            | 6         | 15    |
| Respiratory function (SVC)                               | 2        | 0            | 1         | 3     |
| **Muscle strength**                                      | 3        | 1            | 1         | 5     |
| Manual muscle testing (MMT)                              | 1        | 1            | 1         | 3     |
| Isometric arm strength                                   | 2        | 0            | 0         | 2     |
| **Biomarkers**                                           | 7        | 0            | 0         | 7     |
| Cerebrospinal fluid                                      | 2        | 0            | 0         | 2     |
| Blood                                                     | 1        | 0            | 0         | 1     |
| Neuroradiology                                            | 1        | 0            | 0         | 1     |
| Neurophysiology                                           | 3        | 0            | 0         | 3     |
| Pharmacodynamics/pharmacokinetics                        | 4        | 0            | 0         | 4     |
| **Safety and tolerability**                              | 30       | 0            | 2         | 32    |

AALSRS = Appel amyotrophic lateral sclerosis rating scale; ALSFRS-R = amyotrophic lateral sclerosis functional rating scale revised; ALS-MITOS = amyotrophic lateral sclerosis Milano–Torino staging; CAFS = combined assessment of function and survival; CNS-BFS = Centre of Neurologic Study Bulbar Function Scale; MMT = manual muscle testing; SVC = slow vital capacity.
completed trials, which may reflect reporting bias. A total of 73 trials were recorded as completed, but for 12, the results were not available in a manner consistent with EU or FDA regulations. Of these 12, 2 did not provide completion dates, 2 were completed within the preceding 12 months and 8 were completed more than 12 months prior, noting that FDA and EU regulations mandate reporting of results within 12 months of completion. Three of these trials published results in either journals or on trial registries between April 2019 at the time of writing, while three had published conference proceedings but these did not contain the full dataset.

Figure 2 Number of pwALS participating in trials. (A) (top) shows number of pwALS recruited per trial by phase and year for trials with efficacy-based outcome measures. Published results were used as the source where available (denoted by circle), and registry data were unavailable (denoted by triangles). The 10 trials with highest number of pwALS/trial are labelled with the intervention tested—these trials were two-arm trials with the exception of the following dose-ranging trials: tirasemtiv Phase III (4-arm), masitinib (3-arm), CK-2127107 (4-arm) and talampanel (3-arm). The dashed horizontal lines reflect our sample size calculations using PROACT data: the red line indicates \( n = 200 \) (100/arm) required for a 2-arm RCT with randomization of 1:1, powered at 85% to detect a 25% difference in rate of ALSFRS-R decline over 12 months with a test conducted with a two-sided 20% significance level. The blue line indicates \( n = 500 \) (250/arm) required to evaluate one IMP in a two-arm RCT with randomization of 1:1, powered of 90% to detect a HR of 0.65 for survival over 2 years with a two-sided 5% significance level. (B) (bottom) shows a box and whiskers plot of the number of pwALS recruited per study arm according to CTIMP phase for Phase II (32 trials), Phase II/III (5 trials) and Phase III (12 trials) CTIMPs evaluating potential disease-modifying treatment of amyotrophic lateral sclerosis.
required by regulations. Furthermore, there was inconsistency of data across data sources for a few trials, where conflicting data were recorded on different trial registries or in trial publications. For our review, we analysed data from the primary source as specified in our methods. Ideally, we would group studies according to their aims for our analysis. We would divide studies into exploratory versus confirmatory studies and subdivide exploratory studies according to their main aims including safety, dose-ranging or efficacy. However, most trial registries do not currently include such information. Thus, we limited our analysis to Phase II and Phase III studies, using the primary outcome measure presented to infer the primary aim of each study.

Here, we discuss challenges in designing, delivering and conducting trials in amyotrophic lateral sclerosis, and review the possible causes of low proportion of pwALS entering CTIMPs.

**Understanding of disease biology**

While our understanding of amyotrophic lateral sclerosis disease biology, especially in genetics and molecular biology, has improved over recent years, there remains significant gaps in our knowledge of mechanistic targets and networks in amyotrophic lateral sclerosis.\(^9\) This impedes improvements in disease models, drug development and selection for clinical trials.\(^9,35\) Furthermore, the limitations in disease understanding also poses challenges in developing more reliable, objective, sensitive and specific biomarkers of disease progression and prognosticatiation. Notwithstanding recent advances in biomarker development, these are yet to be established and validated as primary efficacy-based outcome measures in clinical trials.\(^36\)

Thus, we continue to rely on clinical rating scales and survival to evaluate efficacy and narrow inclusion criteria to reduce heterogeneity in trial cohorts. This, in turn, has implications on sample sizes, trial duration, access to trials and generalizability of results as discussed further below.

**Disease-related factors**

Heterogeneity of clinical presentation, site of onset and rate of progression often result in diagnostic delay, with an average time from symptom onset to diagnosis of 12–15 months.\(^37\) Combined with the multiple manifestations, rapidly progressing, disabling and short survival of the disease, this results in a short-time window to identify, screen and recruit pwALS. Furthermore, adherence with trial protocols often becomes challenging with increasing disability; movement, ambulation, speech, swallowing and respiration are commonly affected, the latter causing some pwALS to become dependent on enteral feeding and ventilatory support. Data from Clinical Audit Research Evaluation for Motor Neurone Disease (CARE-MND),\(^38\) the highly curated Scottish MND register, showed that a quarter of pwALS in Scotland between 2015 and 2019 had a gastrostomy and 17% used non-invasive ventilation. This is particularly problematic for trials where invasive procedures, such as intrathecal delivery of IMPs or lumbar punctures, or maintained ability to swallow oral medications are required. Distant travel to trial centres to attend face-to-face appointments is likely to be physically demanding and burdensome for pwALS with accumulating disability. Cognitive and behavioural changes are well recognized and worsen with disease progression,\(^39–41\) forming another barrier to trial participation, including issues around informed consent, and retention.

**Narrow versus broad trial inclusion criteria**

In addition to practical and logistical considerations, phenotypic heterogeneity in amyotrophic lateral sclerosis adds complexity to participant selection. This includes variation in survival, which is well recognized in amyotrophic lateral sclerosis.\(^42–45\) In a Scottish cohort (2015–16), \(~10%\) pwALS are long survivors, defined as survival beyond 8 years, with median survival from onset of 15.6 years.\(^42\) Other sources of heterogeneity with prognostic implications include age at onset, site of presentation, respiratory involvement, subtypes of amyotrophic lateral sclerosis, genetic heterogeneity and diagnostic certainty.\(^1\)

To create a more homogenous cohort, increase trial protocol adherence and exclude long survivors, investigators often use restricted eligibility criteria as demonstrated in our review as well as a review of amyotrophic lateral sclerosis clinical trials between 2000 and 2017 by van Eijk et al.\(^10\) However, employing strict eligibility criteria inevitably affects the generalizability of results. Furthermore, it reduces the proportion of eligible participants in a condition where prognosis is grave and treatment options are severely limited. Van Eijk et al.\(^10\) found that on average 60% of pwALS were ineligible to participate in clinical trials, with more than 20% of pwALS excluded because of El Escorial criteria stipulations alone. Along with restrictions of symptom/disease duration, this has particular implications on recruitment of patients of certain phenotypes,\(^46\) such as progressive bulbar palsy, flail arm or flail leg syndrome, primary lateral sclerosis and progressive muscular atrophy. Notably, the complexity and poor test-retest reliability of the revised El Escorial criteria has been highlighted in a recent study,\(^47\) leading to the proposal of the Gold Coast criteria\(^48\) where diagnostic criteria for amyotrophic lateral sclerosis is simplified and the categories of possible, probable and definite amyotrophic lateral sclerosis in the El Escorial criteria collapsed into a single entity. In our review, 95% of trials excluded pwALS who were enterally or ventilator supported while 42% either mandate riluzole co-treatment or exclude pwALS on concomitant riluzole. Of
note, 45% pwALS in Scotland were started on riluzole between 2015 and 2020, 15% of whom subsequently discontinued riluzole. Indeed when applying the eligibility criteria of the edaravone 2017 trial to an incident cohort, only 10% pwALS would have been eligible. Nevertheless, the authors found that strict eligibility criteria only minimally reduced heterogeneity in trial endpoints. An alternative approach is to adopt broad inclusion criteria recruiting all subtypes of amyotrophic lateral sclerosis including long survivors, supported by protocol innovations to allow participation of pwALS with more advanced disease. Examples include liquid IMP preparations and home/telephone/video assessments. However, this approach requires larger sample sizes to enable a sufficiently powered trial to account for a potentially reduced effect size in a more heterogeneous cohort, or a prospective analysis plan that excludes ALSFRS-R (but not survival) data, from participants who are beyond 8 years of disease onset at trial entry.

The recent development of personalized risk prediction models based on clinical and genetic data, including the presence of a ‘C9orf72’ repeat expansion, could be used for stratification in future trials and might achieve a better balance between endpoint heterogeneity and exclusion rates. Van Eijk et al. calculated that incorporating the European Network for the Cure of Amyotrophic Lateral Sclerosis risk model for participant selection in the edaravone 2017 trial could increase proportion of pwALS eligible nearly fivefold while maintaining similar power.

Figure 3 Example sample size requirements according to trial designs. (A): traditional 2-arm Phase II trial with randomization of 1:1 to evaluate 1 IMP against placebo or control with sample size calculated based on 85% power for a 20% two-sided significance level test to detect a 25% difference in rate of ALSFRS-R decline at 12 months calculated using data from PROACT. (B): a seamless Phase II/Phase III multi-arm multi-stage trial with randomization of 1:1:1 with interim analysis where arms can be stopped if predefined efficacy and safety criteria are not met.
Designing clinical trials to evaluate clinically meaningful change

The tension between practical considerations of feasibility of recruitment, compliance and retention, versus adequate power for evaluation of meaningful outcome measures is apparent throughout the amyotrophic lateral sclerosis trial literature. Indeed, what constitutes a clinically meaningful change on these endpoints is difficult to define. The range of primary outcome measures and primary endpoints reflects the absence of widely available, robust and sensitive biomarkers. There is, however, an encouraging trend towards the use and development of biomarkers in more recent trials. Out of 29 trials including biomarkers as outcome measures with trial start dates available in registries, 21 (72%) were conducted within the past 5 years. Meanwhile, there remains a dependency on clinical ordinal scales, in particular the ALSFRS-R. ALSFRS-R was the most common primary outcome measure used (41% of Phase II studies and 35% of Phase III studies). Survival was the second most common primary outcome measure used in Phase III studies (30%).

Although ALSFRS-R is widely used and validated, a Rasch analysis identified that ALSFRS-R total score lacks unidimensionality, meaning investigators cannot be confident that the same total ALSFRS-R score in two pwALS unidimensionality, meaning investigators cannot be confident in the total ALSFRS-R score in two pwALS reflects comparable clinical conditions. The use of total ALSFRS-R score is particularly problematic where differences in treatment responses between subscales increase and may dilute treatment effects. Furthermore, as ALSFRS-R plateaus and small reversals are common, stable ALSFRS-R scores over short intervals may not represent treatment effect and has implications for trial duration. Notwithstanding these issues, the sample size required to detect a clinically meaningful change in ALSFRS-R is considerable. Using simulation models based on the Pooled Resources Open-Access Amyotrophic Lateral Sclerosis Clinical Trials (PROACT) database, we calculate that ~200 participants (100/arm) are required for a two-arm randomized controlled trial (RCT) with randomization of 1:1, to provide 85% power to detect a 25% difference in rate of ALSFRS-R decline over 12 months with a test conducted with a two-sided 20% significance level (Fig. 3A). We believe this statistically liberal approach is appropriate in Phase II. Similarly, noting that survival is considered the gold-standard outcome for Phase III and using natural history data obtained from CARE-MND, we estimate that >500 patients are required to evaluate one IMP in a two-arm RCT with randomization of 1:1 and 2 years follow up, to achieve a power of 90% to detect a HR for death of 0.65 with a two-sided 5% significance level. A larger sample size would be required to account for the increased heterogeneity should trials use broader inclusion criteria.

These sample sizes demonstrate a major challenge in amyotrophic lateral sclerosis trials. More than half of the studies in our review with efficacy-based primary outcome measures recruited fewer than 100 participants, notwithstanding allowances for missing data, dropouts and death. Thus, with hindsight, these studies were highly unlikely to answer—in a statistically robust manner—whether the proposed intervention was clinically effective or warranted further evaluation in a Phase III trial. The inability to determine drug efficacy or futility in a single, definitive and timely trial, may lead to re-evaluation of the same IMP in subsequent trials. This is inefficient in time, cost and patient resource.

Regulatory variation across countries

Another important challenge in amyotrophic lateral sclerosis trials is the variation in regulatory requirements and complex administrative processes across different countries and jurisdictions. These often hinder and delay multinational clinical trials as demonstrated in other diseases and may detract pharmaceutical companies from running trials in some countries. This is a particular challenge to conducting sufficiently powered, definitive trials in diseases like amyotrophic lateral sclerosis where prevalence is low. Furthermore, as shown in the case of edaravone, regulatory authorities vary considerably in the level and extent of evidence for efficacy considered sufficient for drug approval. This reflects one of many ethical dilemmas in amyotrophic lateral sclerosis clinical trial regulations. On one hand, regulatory bodies exist to protect and safeguard public health, and does so by ensuring that only drugs with acceptable efficacy and safety profiles are approved. On the other, pwALS have strongly advocated for access to treatments, stating that they are willing to accept greater risks that drugs are harmful or ineffective in the face of a fatal incurable illness. Indeed, more than 50 000 people signed a petition by the amyotrophic lateral sclerosis Association and I AM amyotrophic lateral sclerosis to this effect, requesting that FDA make AMX0035 available based on the Phase II trial results. Some argue that so called ‘real-world evidence’ from post-marketing observational studies can substitute for high quality evidence from large RCTs. While such observational studies are useful in detecting rare adverse events and (possibly) large, unexpected beneficial effects, the potential for such studies to determine moderate efficacy or safety signals accurately is limited by inherent biases. There are well established licencing pathways, such as Fast Track (FDA) or PRIority MEdicines—EMA, to accelerate evaluation of IMPs, and programmes such as expanded access (FDA), compassionate use (EMA) or Early Access to Medicines Scheme (UK), which allow access to IMPs for pwALS who are unable to access these in the context of clinical trials. One strategy, therefore, would be for trialists to increase...
their probability of trial success by recruiting more homogenous cohorts through narrower inclusion criteria, with pwALS who do not meet these criteria able to access these drugs via expanded access programmes. However, regulators will usually only provide a marketing authorization for the population in which the IMP has shown efficacy in RCTs. The processes through which amyotrophic lateral sclerosis stakeholders (industry, trialists, clinicians and pwALS) navigate clinical development and regulatory approval is therefore complex, and much would be gained from harmonization and increasing clarity and consistency in national trial regulations, their interpretation and implementation. The Organisation for Economic Co-operation and Development has set out stratified recommendations to this end.63

Proposal for future trials

While efforts to harmonize national regulations are underway, considering the small proportion of pwALS entering interventional clinical trials and the increasing emergence of candidate medicines, there is an opportunity to improve trial design in ALS. This requires innovation in how drugs are selected and how trials are undertaken. The emerging trends in drug selection encompass four broad approaches; machine learning assisted systematic evaluation of the pre-clinical and clinical literature,64 phenotypic high-throughput drug screening platforms adopting human induced pluripotent stem cell modelling of brain cells,65,66 expert panel evaluation of candidate drugs, and more target led discovery of novel putative medicines. Moreover, the emergence of gene-based treatment approaches for monogenetic causes of amyotrophic lateral sclerosis has already led to genetically stratified treatment trials that bring additional considerations including the nature of the control arm.67,68 Regardless of how drugs are selected, the pace of growth in both our biological understanding of amyotrophic lateral sclerosis and linked new technologies for accelerated drug screening and discovery heralds an era where multiple biologically plausible putative medicines will be identified.

Innovative trial designs pioneered in cancer medicine69,70 and successfully adopted in a small number of amyotrophic lateral sclerosis trials may offer a new standard. In our review, 7 out of 10 of these trials, where results were available, delivered conclusive answers on lack of efficacy of IMPs (Table 2). Five15,16,71–73 out of seven trials with adaptive designs and predefined stopping criteria were able to identify futile treatments and stop trials early, reducing exposure of pwALS to ineffective treatments. An adaptive multi-phase approach in evaluating ceftriaxone enabled timely, seamless transition between phases, removing the need for multiple submissions for funding and regulatory approvals.74

Another class of trial design that offers promise for amyotrophic lateral sclerosis is the platform trial. These are trials, which evaluate multiple treatments simultaneously using a single master protocol, typically against a shared control or standard-of-care arm. Platform trials are especially effective where there are several promising medicines without any a priori evidence to favour one over the other.75,76 In this regard, such trials offer significant gains in efficiency by using a single master protocol as opposed to running multiple and inevitably consecutive conventional two-arm studies. Recently, such designs have enabled trial lists to identify the efficacy of dexamethasone in COVID-19 at speed.77 Platform trials may include adaptive features, such as in multi-arm multi-stage (MAMS) platform trials (Fig. 3B). In these, interim analyses are performed at pre-specified timepoints, with treatment arms being discontinued if they do not reach predetermined activity or safety outcomes. Seamless Phase II into III transition can also be prospectively incorporated into the trial protocol and activated if a given arm meets predefined criteria. With appropriate statistical approaches, it is possible to use outcome information gathered from participants in Phase II in the Phase III part of the trial. Further new treatments can be introduced and tested within the protocol.

The time and cost efficiencies gained by stopping ineffective arms early, while being able to introduce new arms in what is in effect a continuous trial platform without initiating further independent RCTs, are significant. It is estimated that a multi-arm study evaluating two active arms within the same trial against control halves the cost of separate traditional two-arm RCTs to evaluate two drugs.76 The Systemic Therapy in Advancing or Metastatic Prostate cancer: Evaluation of Drug Efficacy trial is an example of a successful MAMS trial. Total of 10 treatments have been evaluated over 20 years, resulting in change in standard-of-care three times.78–80 If standard two-arm RCTs had been used, it has been estimated that evaluating the same number of treatments would have required more than 40 years.78–80 Building on this success, similar adaptive trials have been initiated in melanoma, glioblastoma multiforme and Ebola.81

Notwithstanding the benefits of MAMS platform, they present significant operational and statistical challenges. One operational challenge is the need for substantial and sustained funding to enable the creation of an integrated and long-term administrative, project and statistical trial infrastructure.82–84 Statistical analysis plans for MAMS can be complex and include the need to calculate the type I error, or the choice of stopping boundaries, on how to adjust boundaries when the variance of a normally distributed endpoint is unknown, on the impact of adding a treatment arm during a MAMS trial or on whether additional patients should be allocated to the control group in such circumstances.85–88 The organization of interim analyses must also be efficient, with data collection, monitoring and statistical analysis undertaken against tight deadlines, particularly when aiming for a seamless transition from Phase II to Phase III.
Accordingly, the development of platform trials in amyotrophic lateral sclerosis is gathering momentum with three declared initiatives: USA-based HEALEY amyotrophic lateral sclerosis Platform Trial88 (ClinicalTrials.gov registration number NCT04297683), European-based Treatment and Research Initiative to Cure amyotrophic lateral sclerosis ‘MAGNET’ platform trial68 and UK-based Motor Neurone Disease—Systematic Multi-Arm Randomized Trial90 (ClinicalTrials.gov registration number NCT04302870). Each will have a master protocol and is multi-arm, although they vary in terms of inclusion criteria, genetic stratification, selection method of IMPs and the use of trial adaptation. Specifically, the HEALEY trial is designed to evaluate multiple drugs under a master protocol with sharing of participants on placebo across treatment regimens and frequent interim analyses to allow early stopping for success or futility of individual regimens. Participants will be randomized firstly to a treatment regimen, followed by 3:1 randomization to either active treatment or placebo. Current listed arms are zilucoplan, verdiperstat and CNM-Au8. The MAGNET trial is a Phase III adaptive platform trial, which will use a personalized prognostication model to determine eligibility. The first drug to be evaluated in MAGNET is lithium carbonate in UNC13A homozygous patients. MND-SMART is initially testing two IMPs (trazodone and memantine) against a single placebo arm with a seamless adaptive Phase II/Phase III multi-arm, multi-stage design, incorporating measures to maximize accessibility to pwALS including using broad inclusion criteria, liquid IMP and video/telephone assessments.

Conclusion

Phenotypic heterogeneity, disease-related disability and the lack of sensitive and reliable outcome measures are some of the challenges in designing and conducting amyotrophic lateral sclerosis clinical trials. With improved understanding of amyotrophic lateral sclerosis disease biology and increasing number of promising candidate medicines being identified for evaluation, innovative trial designs can be instrumental in making trials more efficient, flexible, scalable and accessible. Together these advances will bring closer a new default where every pwALS has the opportunity to participate in a clinical trial.

Supplementary material

Supplementary material is available at Brain Communications online.

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Competing interests

M.M. and C.J.W. were members of the MHRA Commission for Human Medicines at the time that the mastinib application was declined by the EMA. C.J.W. is an independent data monitoring committee member for clinical trials of AB Science (marketing authorization holder for mastinib) for which University of Edinburgh receives payment. C.J.W. and R.A.P. were supported in this work by NHS Lothian via Edinburgh Clinical Trials Unit. In the last 3 years, J.C. has received support from the Efficacy and Evaluation (EME) Programme, a Medical Research Council (MRC) and National Institute for Health Research (NIHR) partnership and the Health Technology Assessment (HTA) Programme (NIHR), the UK MS Society, the US National MS Society and the Rosetrees Trust. He is supported in part by the National Institute for Health Research, University College London Hospitals, Biomedical Research Centre, London, UK. He has been a local principal investigator for a trial in MS funded by the Canadian MS society. A local principal investigator for commercial trials funded by: Actelion, Biogen, Novartis and Roche; has received an investigator grant from Novartis; and has taken part in advisory boards/consultancy for Azadyne, Biogen, Celgene, Janssen, MedDay, Merck, Novartis and Roche.

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