Research paper

Novel Selenium-based compounds with therapeutic potential for SOD1-linked amyotrophic lateral sclerosis

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ABSTRACT

Background: Amyotrophic lateral sclerosis (ALS), also known as motor neuron disease as well as Lou Gehrig’s disease, is a progressive neurological disorder selectively affecting motor neurons with no currently known cure. Around 20% of the familial ALS cases arise from dominant mutations in the sod1 gene encoding superoxide dismutase1 (SOD1) enzyme. Aggregation of mutant SOD1 in familial cases and of wild-type SOD1 in at least some sporadic ALS cases is one of the known causes of the disease. Riluzole, approved in 1995 and edaravone in 2017 remain the only drugs with limited therapeutic benefits.

Methods: We have utilised the ebselen template to develop novel compounds that redeem stability of mutant SOD1 dimer and prevent aggregation. Binding modes of compounds have been visualised by crystallography. In vitro neuroprotection and toxicity of lead compounds have been performed in mouse neuronal cells and disease onset delay of ebselen has been demonstrated in transgenic ALS mice model.

Finding: We have developed a number of ebselen-based compounds with improvements in A4V SOD1 stabilization and in vitro therapeutic effects with significantly better potency than edaravone. Structure-activity relationship of hits has been guided by high resolution structures of ligand-bound A4V SOD1. We also show clear disease onset delay of ebselen in transgenic ALS mice model holding encouraging promise for potential therapeutic compounds.

Interpretation: Our finding established the new generation of organo-selenium compounds with better in vitro neuroprotective activity than edaravone. The potential of this class of compounds may offer an alternative therapeutic agent for ALS treatment. The ability of these compounds to target cysteine 111 in SOD may have wider therapeutic applications targeting cysteines of enzymes involved in pathogenic and viral diseases including main protease of SARS-Cov-2 (COVID-19).

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1. Introduction

Amyotrophic lateral sclerosis (ALS), also referred to as motor neuron disease (MND) and Lou Gehrig’s disease, occurs in approximately 2 per 100,000 individuals worldwide, causing loss of motor neurons in the cerebral cortex and spinal cord resulting in severe muscle weakness, paralysis and fatal respiratory failure within 2–5 years from disease onset [1]. For some sporadic (sALS) and twenty percent of the familial (fALS) cases, the disease has its origins in superoxide dismutase whose stability as a dimer is compromised leading to a ‘gain of function’ that results in aggregation of protein [2]. Among the 25 genes contributing to fALS [1], genetic defect in sod1 located on chromosome 21q22.11 was the first one to be discovered and is thus best studied with over 180 different known mutations [3], some of which are more common in certain populations. In North America, half of sod1-related fALS cases arise from mutation of Ala4 to Val [4], with one of the most severe forms of fALS with the shortest survival

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Research in Context

Evidence before this study

Toxic inclusion of superoxide dismutase 1 (SOD1) mutant in neuron cells is one of the causes of Amyotrophic lateral sclerosis (ALS), a progressive neurological disorder. There are only two approved drugs, riluzole and edaravone, for ALS treatment with limited effectiveness. Discovery and development of new drugs is urgently needed for ALS patients. Ebselen, a synthetic organo-selenium drug with potent anti-oxidant and cytoprotective effects, is considered to be a potential template for developing therapeutic molecules that can stabilise dimers of ALS-causing SOD1 mutant via binding to cysteine 111 which could reduce aggregation and delay disease progression.

Added value of this study

In this work, we established a group of organo-selenium compounds using benzoisoselenazolone warhead of ebselen with potent SOD1 mutant stabilisation effect. We elucidated binding modes of compounds at the dimer interface of SOD1 mutant by high resolution crystal structures which are beneficial for describing structure-activity relationship. Some of these compounds exhibited better in vitro neuroprotection in mouse neuronal cells than edaravone. In vivo disease onset delay by ebselen has been demonstrated for the first time in transgenic ALS mice model.

Implications of all the available evidence

Our findings suggest ebselen-based compounds are a valuable class of ALS therapeutic agents with superior in vitro neuroprotection than edaravone and with acceptable safety characteristics. In vivo disease onset delay of ebselen also strengthens the potential of ebselen-based compound to be progressed to next stage of drug development programme with potential for therapies that could be offered to patients as alternative treatments alone or in combination with other approved ALS drugs in future.

time [5]. To date, there are only two drugs approved for clinical use in ALS patients: riluzole and edaravone [6]. Riluzole modestly extends life expectancy by 2–3 months in the final stage of ALS but has poor positive effect at earlier stages of the disease [7]. Edaravone has had limited success and acceptance as a drug worldwide but is more effective than riluzole in slowing disease progression, with a limited application time. Riluzole modestly extends life expectancy by 2–3 months in the final stage of ALS but has poor positive effect at earlier stages of the disease [7]. Edaravone has had limited success and acceptance as a drug worldwide but is more effective than riluzole in slowing disease progression, with a limited application time.

2. Materials and methods

2.1. Ethics statement

The experiments using genetically modified animals and organisms were approved by the Animal Care and Use Committee and the recombinant DNA experiment committee of Nagoya University (approval numbers #19269 and #143, respectively). The mice were treated in accordance with ARRIVE guideline and the animal use guidelines of the Animal Care and Use Committee, Nagoya University.

2.2. Synthesis of lead compounds

Details for the synthesis of ebselen analogues are included in the supplementary information.

2.3. Recombinant SOD1 production

The pET303C plasmids containing A4V and double-point mutated A4V C6S human SOD1 genes were generated by site-directed
mutagenesis from wild-type gene using primers shown in Table S1 and transformed into E. coli strain BL21(DE3). The SOD1 transformants were cultured at 37 °C in ampicillin-supplemented LB broth until optical density at 600 nm reaches 0.6–0.8. Then, SOD1 expression was induced by the addition of 0.5mM isopropyl β-d-1-thiogalactopyranoside (IPTG) and 0.3mM ZnSO4 followed by incubation at 18°C for 16 hours. Recombinant proteins were purified by two different methods previously described [28]. A4V<sup>CSS</sup> SOD1 mutant used in DSF assays was purified using affinity-exchange chromatography on diethylaminoethyl (DEAE) sepharose and eluted in a stepwise increasing concentration of NaCl. Protein fractions eluted in 5–100mM NaCl were combined and loaded to a Superdex 20 16/600 column (GE Healthcare) in Tris-buffered saline (TBS) pH 7.4. Recombinant A4V SOD1 used in crystallographic study was purified by precipitation cut using 2.5M ammonium sulphate followed by hydrophobic interaction chromatography on Phenyl sepharose (GE Healthcare) eluted in a stepwise decreasing concentration of ammonium sulphate. A4V SOD1 obtained in 1–2M ammonium sulphate fractions were dialysed against Tris-buffered saline (TBS) pH 7.4 at 4 °C overnight followed by gel filtration on a Superdex 200 16/600 column (GE Healthcare) eluted with the same buffer. The purified proteins were snap-frozen in liquid nitrogen and stored at -80 °C prior to experiments. The concentration of all SOD1 proteins was determined by ultraviolet absorption at 280nm using a molar extinction of 5500M<sup>-1</sup>cm<sup>-1</sup>.

2.4. Differential scanning fluorimetry (DSF)

All DSF experiments were carried out using a StepOnePlus Real-Time PCR machine (Life Technologies). 100 μM A4V<sup>CSS</sup> SOD1 was pre-incubated with 500 μM compounds diluted from 250mM stock solution at 4°C overnight. The mixtures were buffer-exchanged against fresh TBS using micro-centrifugal ultrafilter (Vivaspin, MWCO 50kDa) to remove excess compound. 70 μM buffer-exchange protein was mixed with 10 × concentration of SYPRO Orange dye (Life Technologies) which was used as a fluorescent probe. Fluorescent intensities were monitored from 25 °C to 95 °C at a ramp rate of 0.3 °C/min. The melting temperature (T<sub>m</sub>) was calculated based on the Boltzmann equation using the T<sub>m</sub>Tool<sup>TM</sup> software (Life Technologies).

2.5. Crystallisation and structure determination

All compounds were prepared for 250mM stock solution by dissolving in DMSO. 0.2mM of A4V SOD1 was incubated with 1mM compound being at 4 °C overnight before concentrated to 10mg/ml SOD1. 3 μL of A4V SOD1 solution was set hanging crystallisation drops by mixing with 3 μL of reservoir solution containing 100mM sodium acetate pH 4.7, 150mM NaCl and 2.5–2.8M ammonium sulphate. The crystallisation drops were incubated at 19°C for 2–3 days allowing vapour diffusion against corresponding reservoir solutions before macro-seeding using a few A4V SOD1 crystals in P2<sub>1</sub> space group previously grown in the same condition. Mature crystals were obtained within a week after seeding and soaked in Paratone oil (Hampton Research) as cryo-protectant before freezing in liquid nitrogen. Diffraction data was collected at 100K on i03 (compounds 3, 6 and 9), i04 (compounds 1 and 5), i04-1 (ebselen, compounds 2 and 10) and i24 (compound 13) at Diamond Light Source, UK, integrated using DIALS [29] and scaled using Aimless [30] in CCP4 suite. Phase problem of all structures was solved by molecular replacement using an A4V SOD1 structure (PDB: 1UXM; chain A and B) as a starting model in MolRep [31]. Structure models were initially refined by rigid body refinement in Refmac5 [32] and then manually modified in COOT [33] followed by cycles of restrained refinement in Refmac5. Ligand models of ebselen-based compounds were produced in Jlignand [34] and manually added into corresponding electron density.

2.6. Cell culture, transfection and MTS assay

Mouse neuroblastoma Neuro2a cells (RRID: CVCL_0470) were cultured and measured their viability as described previously [35]. Briefly, the cells seeded at 5.0 × 10<sup>4</sup> /well on a poly-D-lysine coated 96 well plate were transfected to express mutant human SOD1 species using Lipofectamine 2000 (Thermo Fisher Scientific). After 6 h of transfection, the growth medium (Dulbecco’s Modified Eagles’ Medium (DMEM) containing 4.0 g/L glucose, 10% (v/v) fetal bovine serum (FBS) (both from Thermo Fisher Scientific)) was replaced with a differentiation medium (DMEM containing 1.0 g/L glucose, 2% (v/v) FBS and 2 mM N6,2-0-dibutyryladenosine 3’,5’-cyclic monophosphate (Nacalai Tesque, Kyoto, Japan)) with the indicated ebselen-derived compounds at the indicated concentration. All the compounds were first dissolved into DMSO at 10 mM and stored at -30 °C until use. The cells were incubated for 48 h, and the differentiation medium was replaced with a fresh one. Then, the cell viability was measured using MTS assay (Celltiter 96<sup>®</sup> AQueous One Solution Cell Proliferation Assay, #G3580, Promega Biosciences, San Louis Obispo, CA, USA) according to the manufacturer’s instructions with an absorption spectrophotometer Infinite F50 (TECAN Group Ltd., Männedorf, Switzerland). To determine LC<sub>50</sub> of the compounds, the cell viability at different compound concentrations was fitted using the least squares method to a sigmoid curve with the following formula: 

\[
\text{Cell Viability} = \left( 1 + 10^{[\text{logIC50}-\text{log}\text{Concentration}]} \right)^{-1}
\]

2.7. Animals

Transgenic mice expressing SOD1<sup>G93A</sup> (B6.Cg-TgSOD1<sup>G93A</sup> I1Gur/J) (RRID: IMSR_JAX:004435) were obtained from the Jackson Laboratory (Bar Harbor, ME, USA). Genotyping for the SOD1G93A mice and determination of the times for disease onset and end-stage were previously described [36]. In brief, the disease onset was determined by the time when the mice reached maximal body weight. The mice were randomly divided into two groups that fed with a powdered CE-2 diet (CLEA Japan Inc., Tokyo, Japan) in the presence or absence of 0.016 % ebselen (#E0946, Tokyo Chemical Industry Co., Ltd., Tokyo, Japan) using a feeder for powder diet provided by Shinano Seikakusho Inc. (#SN-950) (Tokyo, Japan) from 70 days of age to the end-stage. The diet was replenished three times per week, and the mice had free access to their assigned diet and deionized water. We estimated the amount of ebselen taken by mice from the food intake, and confirmed that about 300 μg/kg/day ebselen was taken during all over the cohort. The mice were housed in the specific pathogen-free (SPF) environment (with a 12 h light–dark cycle at 23 ± 1°C) and treated in compliance with the requirements of the Animal Care and Use Committee, Nagoya University.

2.8. Statistics

For animal experiments, G93A SOD1 mice were grouped in a random manner. Measurements of motor function and body weights were carried out in a non-blinded manner. Survival times were analyzed with a log-rank test, and no sample size calculation was performed in this study. All the data of MTS assays were analyzed by one-way ANOVA followed by the post-hoc Tukey’s multiple comparison t-test. All the statistical analyses were carried out by using GraphPad Prism 8 software (GraphPad Software, La Jolla, CA).

2.9. Role of funding source

Compound synthesis, DSF screen and crystallography were supported by the ALS Association grant (WA1128). Cell-based and animal model studies were granted by Fostering Joint International Research (19KKn0214) from the Ministry of Education, Culture, Sports, Science and Technology (MEXT), Japan. Funders did not play any part in study
3. Results

3.1. Generation of novel ebselen analogues with good physicochemical properties that promote SOD1 thermal stability

Ebselen has been demonstrated to bind cysteine and stabilise dimer of A4V SOD1, which is severely prone to monomerisation and aggregate leading to acute form of ALS [17,26]. Based on benzoisoleselenazolone core of ebselen, functionalised N-aryl and methylene linker moieties were incorporated into the organoselenium core to improve dimer stabilisation of SOD1 and drug-like properties (Figure S1). To rank stabilising effect of ebselen-based small molecules, DSF screen against A4V SOD1 with C65 mutation background (A4V$^{C65}$ SOD1) has been used to evaluate performance via different melting temperatures of compound-treated and native protein ($\Delta T_m$). C65 background was designed to eliminate the destabilisation of A4V SOD1 in solution due to occasional binding of ebselen to free thiol of cysteine which results in undeterminable melting curve of DSF assay [26]. A greater positive $\Delta T_m$ indicates improved thermal stability of A4V$^{C65}$ SOD1 affected by compound binding. The compounds with a methylene linker imparted greater thermal stability than N-aryl derivatives. The structures of A4V SOD1 bound with ebselen and its methylene linked derivative (compound 2) derived from C2 space group crystals reveal distinct binding poses of ligands (Figure S2), which could describe different outcomes in $\Delta T_m$ values. Ebselen, an N-aryl compound, interacts with loop VI of SOD1 (residues 101–109), whilst compound 2 poses perpendicularly to ebselen placing its aromatic tail into the gap between Thr2 and Ile151. Therefore, functionalising the aryl ring of the methylene linker scaffold that enables further interactions may provide better stabilisation of SOD1 dimer. This offers greater substrate scope to be incorporated into the methylene linker series compared to the N-aryl scaffold analogues [26].

Herein, a series of methylene linker (compounds 1–10 and 14) and N-aryls (compounds 11–13) compounds incorporating a variety of functional groups were synthesised and their $\Delta T_m$ values in A4V$^{C65}$ SOD1 were measured by DSF assays. Compounds 1, 3 and 9 have stronger stabilising effect on A4V$^{C65}$ SOD1 above other methylene linked compounds as shown in Table 1. Furthermore, incorporation of a carbonyl morpholine moiety with benzoisoleselenazolone core (compounds 12–14) seems to establish outstanding $\Delta T_m$ values at the same level or better than ebselen. Apart from increasing stability of SOD1 dimer, potent candidates need to have suitable drug-like properties that allow central nervous system (CNS) access and desirable ADME properties to ensure the compound is CNS optimised. For the compounds with a methylene linker, we have successfully co-crystallised six of this class of compounds giving good opportunity to see the influence of Cl and OMe substituents on the phenyl or pyridyl tail group. Starting with chlorobenzyl ebselen, compounds 2 and 5 have a highly similar ligand pose observed in overlaid crystal structures in Fig. 2a. Unlike N-aryls compounds, A4V SOD1–compounds 2 and 5 structures display longer distance between seleno-aromatic rings (Fig. 2b). Thus, the interaction between compounds is weaker than N-aryls compounds. Moreover, meta–Cl substituent shifts the benzyl group away from Thr2 and Ile151 impairing hydrophobic contacts between N- and C-termini (Fig. 2b). This may cause weaker SOD1 stability of compound 5 than compound 2. In the superimposed structures of A4V SOD1–compounds 3 and 6, only one monomer appears perfectly overlaid compound molecules like compounds 2 and 5 (Fig. 2c). Compound 6 at another monomer acts as a bridge between monomers by localisation of its meta-methoxybenzyl tail closer to the main chain of Asp109 of another monomer, while compound 3 behaves identically to compounds 1, 2 and 5 whose tail picks up hydrophobic interactions with Thr2 and Ile151. Cross-monomer interaction of compound 6 seems to enhance dimer interface whereas $\Delta T_m$ of compound 6 is lower than compounds 1 and 3, which have closer contact to N- and C-termini (Fig. 2c). This indicates that the interaction between Thr2 and Ile151 is probably more important for dimer stability than the interaction between benzyl tail and neighbour Asp109. On the other hand, this concept is not applicable to compounds 9 and 10 that pyridyl or para-methoxypyridyl rings were replaced in the scaffold of compound 1. Strong stabilising profile was observed in both compounds, especially compound 9 with greater $\Delta T_m$ than ebselen. The binding pose of compound 9 appears in the same way as compound 6 but compound 10 performs similarly to compound 3 (Fig. 2d). This contrasting effect may occur because
higher polarity of the pyridyl ring that can establish stronger cross-
monomer linking than compound 6 via dipole-dipole interaction
with the main chain of neighbour Asp119 (Fig. 2d) resulting in the
best stabilisation among methylene linker series. Based on an inte-
grated knowledge of the binding mode visualisation in crystallo-
graphic structures, DSF assays and MPO score analysis, we noted that
compounds 9 & 10 from methylene linker series and compounds 12 & 13 from N-aryl series are front runners from this class with excel-

Table 1

| Structure | \( \Delta T_m \) (°C) | A4V\textsuperscript{CAs8} SOD1 dimer stability (\( \Delta T_m \)), MPO score and each physicochemical parameter that contributes to the overall score. 1-3 are from [26]. A traffic light system was employed to demonstrate each value contribution; Green – good/desirable; Amber – ok; Red – poor/undesirable. ClogP: calculated logarithm of partition coefficient of a compound between n-octanol and water, ClogD: calculated logarithm of distribution coefficient of a compound between n-octanol and water at pH 7.4, MW: molecular weight, tPSA: topological polar surface area, CNS MPO: central nervous system multiparameter optimisation. | ClogP | ClogD | tPSA (Å\(^2\)) | MW | HBD | pK\(_a\) | MPO |
|-----------|---------------------|------------------------|--------|--------|----------------|-----|-----|------|--------|
| Ebselen   | +8.77               | 3.70                   | 2.62   | 20.31  | 274.19        | 0   | -0.4| 4.4  |
| 1         | +8.76               | 3.73                   | 2.76   | 20.31  | 288.22        | 0   | -0.9| 4.3  |
| 2         | +6.36               | 4.44                   | 3.21   | 20.31  | 322.66        | 0   | -1.1| 3.7  |
| 3         | +8.73               | 3.65                   | 2.53   | 29.54  | 318.25        | 0   | 0.8 | 4.9  |
| 4         | +6.70               | 3.87                   | 2.76   | 20.31  | 206.20        | 0   | -1.0| 4.2  |
| 5         | +3.95               | 4.44                   | 3.21   | 20.31  | 322.66        | 0   | -1.1| 3.7  |
| 6         | +7.77               | 3.65                   | 2.53   | 29.54  | 318.25        | 0   | 0.8 | 4.9  |
| 7         | +6.85               | 3.87                   | 2.76   | 20.31  | 206.20        | 0   | -1.0| 4.2  |
| 8         | +0.17               | 4.76                   | 3.45   | 29.54  | 372.51        | 0   | -1.0| 3.8  |
| 9         | +9.62               | 2.23                   | 1.52   | 32.67  | 289.20        | 0   | 4.6 | 5.6  |
| 10        | +8.08               | 3.05                   | 1.49   | 41.90  | 319.22        | 0   | 2.9 | 6.0  |

3.3. In vitro neuroprotection and toxicity of lead compounds in mouse neuronal cells

A phenotypic screening of ebselen, edaravone and nine lead com-

pounds (1, 2, 5, 6, 9, 10, 11-13) has been undertaken in mouse N2a
neuronal cells transfected with human wild-type and G93A SOD1, which is the mutant used in representative SOD1-ALS transgenic mice [27]. The biophysical study of G93A SOD1 revealed that the mutation at residue 93 destabilises SOD1 via opening β-barrel.
at the concentrations of 0.1, 1 and 10 in SOD1-ALS transgenic mice. All compounds were dosed to N2a cells against neuronal cells expressing G93A SOD1 is still critical for verifying dimer stabilising compound activity to cells in this experimental condition. We found that transcontrol, because wild-type SOD1 does not induce any obvious dam-
matter stability of wild-type by only 2 fold and G93A SOD1 by just over 5 fold in contrast to the 60 fold increase for A4V. G93A SOD1 was not used as a model to find dimer stabilising compound via DSF assay. However, in vitro screen against neuronal cells expressing G93A SOD1 is still critical for verifying neuroprotective activity of compound prior to in vivo experiment in SOD1-ALS transgenic mice. All compounds were dosed to N2a cells at the concentrations of 0.1, 1 and 10 µM and then incubated for two days before measuring cell viability by MTS assay. Relative cell viability of each compound at different concentrations are presented in bar charts in Fig. 3. We used N2a cells expressing wild-type SOD1 as a control, because wild-type SOD1 does not induce any obvious damage to cells in this experimental condition. We found that transgenic N2a cells expressing G93A SOD1 used as negative control have significantly lower cell viability, about 62% of wild-type SOD1. Fortunately, edaravone which is an approved ALS drug, was used as positive control in this screen. With the aim of producing an ALS drug that can stabilise SOD1, a good compound should redeem G93A SOD1-transfected cell using a lower dose than edaravone. That can stabilise SOD1, a good compound should redeem G93A SOD1, but also enhanced cell survival in other ALS SOD1 transfected cell, but also enhanced cell survival in other ALS SOD1 mutants, including A4V. This result suggests that ebselen-based compounds are also applicable and beneficial for neuronal cells expressing other ALS-linked SOD1 mutants.

### Table 2

| Structure | ATₘ (°C) | ClogP | ClogD | tPSA (Å²) | MW  | IBD  | pKₘ | MPO |
|-----------|----------|-------|-------|-----------|-----|------|------|-----|
| Ebselen   | +8.77    | 3.70  | 2.62  | 20.31     | 274.19 | 0   | -0.4 | 4.4 |
| 11        | +4.03    | 3.45  | 1.91  | 32.78     | 374.33 | 0   | 6.6  | 5.3 |
| 12        | +9.90    | 2.33  | 1.25  | 49.85     | 387.30 | 0   | -1.0 | 5.8 |
| 13        | +10.52   | 2.48  | 1.26  | 49.85     | 405.30 | 0   | -1.5 | 5.7 |
| 14        | +8.67    | 2.36  | 1.84  | 49.85     | 401.32 | 0   | -2.0 | 5.7 |

instead of impairing dimer interface like A4V. Native mass spectrometry demonstrated that G93A SOD1 has wild-type-like dimer interaction. Ebselen increased dimer affinity of wild-type by only 2 fold and G93A SOD1 by just over 5 fold in contrast to the 60 fold increase for A4V. GCMS of compound 8, which possesses excellent activity in N2a and edaravone which is an approved ALS drug, was used as positive control in this screen. With the aim of producing an ALS drug that can stabilise SOD1, a good compound should redeem G93A SOD1 transfected cell using a lower dose than edaravone. Most of the ebselen-based compounds seem to be toxic to N2a cell at 10 µM resulting in lower viability than the negative control. Fortunately, many compounds performed good neuroprotective effect at 0.1 and 1 µM. Compounds 9 and 10 appeared to be the most potent compounds recovering G93A SOD1-transfected cell to the same level of wild-type cell even at 10µM (indicating better overall safety index) and could be selected for in vivo experiments. In contrast, compounds 12 and 13, which possess excellent ΔTₘ values, gave poor neuroprotection at 1 and 10 µM. Incorporation of the carbonyl morpholine group in compounds 12 and 13 might not only enhance stabilisation of SOD1 dimer, but also stimulate unexpected cytotoxicity. To verify this concept, cell viability of N2a cells in ebselen, compounds 6 and 12 at seven concentrations were measured and median lethal doses (LC₅₀) were calculated. The results in Fig. S5a show that LC₅₀ of compound 12 (17.6 µM) was about half of ebselen (32.8 µM), indicating that compound 12 is more cytotoxic than ebselen. On the other hand, compound 6 had LC₅₀ at 71.0 µM that is considered as a safer compound than ebselen. To gain further insight, we assayed compound 6 at 0.1 µM in N2a cells expressing other SOD1 mutants- A4V, G37R, G85R (Fig. S5b). This compound was found to be effective not only in G93A SOD1-transfected cell, but also enhanced cell survival in other ALS SOD1 mutants, including A4V. This result suggests that ebselen-based compounds are also applicable and beneficial for neuronal cells expressing other ALS-linked SOD1 mutants.

#### 3.4. In-vivo experiments: impact of Ebselen in G93A SOD1 transgenic mice

In preparation for selecting potential candidates to progress into the clinical phase, we have set up in vivo trial of compounds using G93A SOD1 transgenic mice. The first set of experiments have been completed using ebselen to give an indication of the potential in-vivo efficacy of this scaffold and identify potential lead compounds. The mice were fed by powder food mixed with/without 0.016% w/w ebselen (estimated dose: 24 mg/kg) from day 70 till end-stage. Survival experiments were performed to determine disease onset and mouse survival time, shown in Fig. 4. We found that ebselen significantly delayed the disease onset of ALS in mice (control: 118.9±5.8 days and ebselen: 128.0±5.2 days shown as mean±SD). However, mean survival time of transgenic mice was extended only marginally (control: 153.1±11.4 days and ebselen: 156.6±11.4 days, mean±SD). The body weights, the rotarod performance, and the clasping signs, which are the markers used to evaluate disease progression, were not affected by ebselen (Fig. S6).

#### 4. Discussion

SOD1 is a ubiquitously expressed protein enriched in nervous system. Genetic mutations throughout the SOD1 structure, including A4V substitution, result in protein destabilisation and cystolic...
aggregation leading to neuron death [5]. A4V mutation causes steric clashes with neighbouring residues, altering dimer interface and disturbing protein dimer stability and affinity [5,39,40]. Thus, preventing monomerisation of SOD1 is considered an important feature of novel therapeutic agent that delays ALS progression. Cys111 in SOD1 is amenable to be oxidised that is considered to promote monomerisation and inclusion formation [18,41]. As such it has become a focus for small molecule intervention for stabilising SOD1 monomerisation and inclusion formation [18,41]. As such it has become a focus for small molecule intervention for stabilising SOD1 monomerisation and inclusion formation [18,41].

Ebselen has been proven as a potential template with good ability to increase the affinity of A4V SOD1 dimer formation [17] and thermal stability of A4VC6S SOD1 [26] when bound with cystein. In this study, we generated a group of organoselenium compounds using benzoisoleselenazolone ring of ebselen incorporated with functionalised methylene linker and N-aryl moieties. The new molecules were designed as CNS-accessible compounds with drug-like-ness profiles on the basis of high scores of MPO from predicted physicochemical properties (Tables 1&2) before evaluating their performance against A4VC6S SOD1 in DSF assay. Some compounds in methylene linker series (compounds 1, 3, 6, 9, 10, 12, 13), which exhibited SOD1 stabilisation with large positive Dₜm values at the same degree or better than ebselen, were listed as potent candidates. Crystallographic investigation of A4V SOD1 bound with those candidates revealed binding poses at cystein and rationalised how each chemical moiety interacts with SOD1 and also reducing the chance of oxidative modification [15–17].

Table 3 Crystallographic data collection and refinement statistics of A4V SOD1 with bound compound crystals in P2₁, space group.

| Parameter/ligand | Compound 1 | Compound 2 | Compound 3 | Compound 5 |
|------------------|------------|------------|------------|------------|
| Data collection  | P2₁        | P2₁        | P2₁        | P2₁        | P2₁        |
| Space group      |            |            |            |            |            |
| Cell dimensions  | a, b, c (Å) | 38.95, 67.72, 50.65 | 38.34, 67.95, 50.46 | 38.61, 68.29, 50.90 | 37.33, 68.26, 50.71 | 38.50, 68.32, 51.32 |
| α, β, γ (°)      | 90.00, 106.40, 90.00 | 90.00, 105.54, 90.00 | 90.00, 105.76, 90.00 | 90.00, 105.07, 90.00 | 90.00, 106.36, 90.00 |
| Resolution (Å)* | 39.48–1.45 (1.47–1.45) | 48.61–1.35 (1.37–1.35) | 39.80–1.95 (2.00–1.95) | 32.46–1.60 (1.63–1.60) | 36.94–1.55 (1.58–1.55) |
| Rmerge            | 0.0181 | 0.0151 | 0.0117 | 0.0118 | 0.0131 |
| Rfree             | 0.0147 | 0.0115 | 0.0103 | 0.0155 | 0.0131 |
| CC1/2 (%)*        | 99.3 (99.8) | 99.8 (99.4) | 99.7 (98.9) | 100.0 (99.8) | 99.8 (99.6) |
| Compleness (%)*  | 1.74 (1.75) | 1.68 (1.69) | 1.60 (1.61) | 1.55 (1.56) | 1.50 (1.51) |
| Redundancy*       | 3.3 (3.4) | 3.2 (3.6) | 3.1 (3.2) | 3.4 (3.4) | 3.2 (3.3) |
| No. reflections   | 44,730 | 54,619 | 48,610 | 54,200 | 33,188 |
| Rwork/Rfree       | 18.1610 | 18.1610 | 18.1610 | 18.1610 | 18.1610 |
| No. atoms         | 2237 | 2291 | 2200 | 2289 | 2291 |
| Protein           | 32/34 195 | 34/19 229 | 36/14 95 | 57/22 223 | 36/166 |
| Ligand/ion Water  | 31.28 | 31.43 | 31.43 | 31.43 | 31.43 |
| B-factors         | 22.06 | 20.27 | 31.69 | 26.15 | 22.24 |
| Redundancy*       | 3.3 (3.0) | 3.5 (3.1) | 3.1 (3.1) | 3.4 (3.2) | 29.98 |
| Bond angles (°)   | 0.0181 | 0.0117 | 0.0118 | 0.0131 | 0.0131 |
| PDB code          | 6Z4G | 6Z4O | 6Z4H | 6Z4I | 6Z4J |

* Values in parenthesis denote the highest resolution shell.
can establish water bridging with SOD1 residues at dimer interface through hydrogen bonds that results in fantastic thermal stability. In contrast to N-aryl series, the compounds with methylene linker have binding mode almost perpendicular to N-aryl compounds due to their higher flexibility. They reinforce dimer stability by picking up interactions with Thr2 and Ile151 of N- and C-termini of SOD1, respectively. This point could be utilised by growing small functional groups at benzyl tail to develop additional contacts. Incorporation of para-methoxy group in compound 3 seems to be more beneficial than halogen atoms according to significant greater $\Delta T_m$ because its size and position may perfectly fit the space between Thr2 and Ile151. Nevertheless, meta-methoxy group results in distinct poses of benzyl tail in A4V SOD1 dimer. Interestingly, crystal structures of compounds 6 and 9 shows that benzyl tail of one ligand establish cross-monomer linking with Asp109 of opposite monomer. This seems to result in better stabilisation especially in compound 8 promotes thermal stability above other compounds in the same series. Moreover, chemical modification using a pyridyl ring as a more polar aromatic tail in compound 9 confers exceptional values of MPO scores and physicochemical parameters that opens opportunity to explore more potential compounds based on this moiety. We modified compound 9 by incorporating with a para-methoxy group (compound 10) or carbonyl morpholine (compound 14). Both compounds certainly gave the highest rank in MPO scores and other parameters and effective A4V SOD1 stabilisation which may be potential neuroprotective agents with expected good drug-likeness.

We tested the neuroprotective activity against G93A SOD1 expressing mouse neuronal cells of potent candidates which were selected by exceptional $\Delta T_m$, MPO and physicochemical parameters scores. The parent ebselen can redeem G93A SOD1 neuron cell viability almost up to wild-type level in much lower dose than the licensed drug edaravone, which implies better potency for this class of compounds. We can see potentially high survival level of neuronal cells after treatment with low concentrations at 0.1 and 1 $\mu$M of all methylene linker compounds. However, most compounds in this series and ebselen show lethality to neuron cells at concentration of 10 $\mu$M. Compounds 9 and 10 containing a pyridyl ring seems to be the most promising candidates among tested compounds with the strongest in vitro neuroprotective activity and much smaller loss of neuron cells at high dose. Thus, the methylene linker compounds with pyridyl group should be the focus for further development and optimisation of safe neuroprotective compounds, but para-methoxy group appears not essential to improve healing effect in mouse neuronal cells. Carbonyl morpholine derivatives (compounds 12, 13), which are the most promising candidates due to their excellent $\Delta T_m$ and predicted MPO scores, exhibit the poorest level of neuroprotection among other candidates. This observation suggests carbonyl morpholine group displays non-selective binding to other endogenous targets leading to neuron cell death.

In addition to in vitro neuroprotective performance of our compounds, transgenic mice expressing G93A SOD1, a frequently used SOD1-ALS mouse model, were treated orally with ebselen and monitored for their motor activities and survival times. Even though ebselen does not extend total lifetime of experimental mice appreciably compared to the controls, we observed a delayed disease onset in ebselen-treated G93A mice (>10 days). This confirms that ebselen can be absorbed in digestive tract and replicate its neuroprotective effect in animal model. Ebselen has a reducing activity to suppress oxidative stress that may contribute to neuroprotective effect [25]. However, our compounds, apart from their reducing potential,
showed significantly improved neuroprotective activity measured by our MTS assay. This suggests that the direct interaction with SOD1 is a key determinant of neuroprotective activity of our new compounds rather than their reducing activity per se. We propose that compounds 9 and 10 may prove better in the in vivo activities than ebselen and result in an increased survival rate of transgenic mice and thus may have better prospects for clinical trials.

Thus, we have generated two promising series of ebselen-based compounds for stabilising mutant SOD1 including A4V. We used DSF assays, MPO scores and predicted physicochemical parameters to rank lead compounds before measuring neuroprotective effects in cell-based assays and animal models. The methylene linked series showed outstanding neuroprotective activities in mouse neuronal cells, particularly compounds 9 and 10 with a pyridyl tail, which can be considered an excellent scaffold for further optimisation in drug development programme. However, our N-aryl compounds with carbonyl morpholine are less selective in spite of excellent outcomes in DSF assays. These results demonstrate that DSF assay is a good tool for hit to lead optimisation in this disease area, but phenotypic cell-based experiments are still required to evaluate in vitro effectiveness and cytotoxic effects. Apart from neuroprotection and safety profiles, in vitro and in vivo pharmacokinetic and pharmacodynamics studies of lead compounds should be addressed to select the most powerful candidates. Here, we also verified neuroprotective action of ebselen in mouse model showing effective oral availability and delay in disease onset. Further optimisation of this template can be envisaged to improve drug potency and increase the overall survival time.

Fig. 2. Superimposed and individual crystal structures of A4V SOD1 bound with compounds. a Ebselen (pink) and compound 13 (yellow). b Compounds 2 (cyan) and 5 (pink). c Compounds 3 (cyan) and 6 (pink). d Compounds 9 (cyan) and 10 (pink). Surrounding amino acid residues of each monomers are coloured in purple and blue. Waters are represented as red dots. Hydrogen bonds are illustrated as green dashed lines. The distances between atoms and between aromatic rings are illustrated as yellow and black dashed lines, respectively. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)
Fig. 3. Ebselen-based compounds confer neuroprotection in mouse N2a cells expressing G93A mutant form of SOD1. MTS assays of mouse N2a cells transfected with wild-type (WT) or G93A SOD1 in the presence of ebselen, ebselen-based compounds, or edaravone at the indicated concentrations. The results with each the ebselen-based compounds are displayed with colors based on its physicochemical parameters shown in Table 1. Green = good/desirable; Light green = almost good/desirable; Amber = ok; Red = poor/undesirable. Relative cell viability levels of WT and G93A SOD1 are shown as blue and black dashed lines, respectively. Asterisks indicate statistically significant protection ($p < 0.05$) against G93A SOD1. Averaged cell viability ±SEM relative to the one expressing WT SOD1 was plotted. (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)

Fig. 4. Onset and survival curves of G93A SOD1 transgenic mice treated with ebselen. a Disease onset curves for ebselen-treated G93A SOD1 mice and control G93A SOD1 mice (Ebselen: 128.0±5.2 days, Control: 118.9±5.8 days; mean±SD, $p = 0.0006$ in log-rank test). b Survival curves for ebselen-treated G93A SOD1 mice and control (Ebselen: 156.6±11.4 days, Control: 153.1±11.4 days; shown as mean±SD, $p = 0.3530$ in log-rank test). The same numbers of mice were used for onset and survival analysis (control, n=10; ebselen, n=12). All the mice are female.
We have established that the new generation of ebselen-based compounds are potent, multimodal SOD1 pharmacological chaperones targeting cysteine 111, restoring the dimer affinity of, say A4V SOD1 to that of uncompromised wild-type SOD1, prevent the formation of oligomers/aggregates. This is an important significant milestone in the chemical-assisted manipulation of SOD1 for the treatment of ALS endorsed by the neuroprotection offered in mouse N2a neuronal cells transfected with human wild-type and G93A SOD1. Several of the compounds including ebselen redeemed G93A SOD1 transfected cells better than the drug Edaravone (approved in USA in 2017), which has had limited efficacy and acceptance as a drug worldwide, requiring daily intravenous infusion. In addition to the in vitro neuroprotective performance, oral treatment with ebselen did not extend total lifespan of G93A SOD1 mice, but significantly delayed a disease onset by ~10 days, confirming that ebselen (and by extension ebselen-based compounds) can be absorbed in digestive tract and replicate its neuroprotective effect in animal model. In summary, we have demonstrated outperformance of ebselen and its new generation of leads over edaravone where the formulations are suitable for effective oral administration for a sustained treatment for ALS.

The ability of our compounds to target a cysteine may extend to other protein targets such as the main protease (Mpro) of COVID-19 virus whose recent crystallographic studies with inhibitor and the docking study with ebselen indicates covalent interaction with Cys145 is the mechanism of its inhibition [42]. This is further supported by the docking of new ebselen-based compound developed here, Fig. 57.

Author contributions
Conceptualization, S.S.H. and P.M.O.; Methodology, K.A., M.C.R., and S.W.; Investigation, K.A., M.C.R., S.W., S.S.H., P.M.O. and K. Y.; Supervision, S.S.H., P.M.O. and K. Y. All authors read and approved the final version of the manuscript.

Declaration of Interests
The authors declare no competing interests.

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Supplementary materials
Supplementary material associated with this article can be found, in the online version, at doi:10.1016/j.ebiom.2020.102980.
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