Protein production is an early biomarker for RNA-targeted therapies

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Abstract

Objectives: Clinical trials for progressive neurodegenerative disorders such as Alzheimer’s Disease and Amyotrophic Lateral Sclerosis have been hindered due to the absence of effective pharmacodynamics markers to assay target engagement. We tested whether measurements of new protein production would be a viable pharmacodynamics tool for RNA-targeted therapies. Methods: Transgenic animal models expressing human proteins implicated in neurodegenerative disorders — microtubule-associated protein tau (hTau) or superoxide dismutase-1 (hSOD1) — were treated with antisense oligonucleotides (ASOs) delivered to the central nervous system to target these human mRNA transcripts. Simultaneously, animals were administered 13C6-leucine via drinking water to measure new protein synthesis after ASO treatment. Measures of new protein synthesis and protein concentration were assayed at designated time points after ASO treatment using targeted proteomics. Results: ASO treatment lowered hTau mRNA and protein production (measured by 13C6-leucine-labeled hTau protein) earlier than total hTau protein concentration in transgenic mouse cortex. In the CSF of hSOD1 transgenic rats, ASO treatment lowered newly generated hSOD1 protein driven by decreases in newly synthesized hSOD1 protein, not overall protein concentration, 30 days after treatment. At later time points, decreases in newly generated protein were still observed after mRNA lowering reached a steady state after ASO treatment. Interpretation: Measures of newly generated protein show earlier pharmacodynamics changes for RNA-lowering therapeutics compared with total protein concentration. Early in ASO treatment, decreases in newly generated protein are driven by changes in newly synthesized protein. Measuring new protein production in CSF may be a promising early pharmacodynamics marker for RNA-targeted therapeutics.

Introduction

The number of persons aged 65 and older globally is projected to triple by the year 2050, increasing the number of people living with adult-onset, progressive neurodegenerative diseases in the population. There is a dire need to develop novel therapeutics for these conditions, as current FDA-approved drugs are not adequately effective, and no treatments exist to reverse disease pathology. Despite promise of new therapies in preclinical models over the past 30 years, these compounds have failed in Amyotrophic Lateral Sclerosis (ALS) and Alzheimer’s Disease (AD) clinical trials at rates greater than 95% and 99%, respectively. One important way to improve the current treatment landscape is to focus on the development of novel methods to determine whether a therapeutic has engaged the intended target in human clinical trials.

A central hypothesis in neurodegenerative disease is that the accumulation of misfolded proteins over time imparts selective toxicity to specific neuronal populations, causing...
progressive neurodegeneration and manifestation of symptoms. This pathological misfolding of proteins in AD, ALS, Parkinson’s Disease (PD), and Huntington’s Disease (HD) suggests that an approach to inhibit the production of toxic proteins within the cell may be an effective therapy. One such approach includes lowering levels of RNA to decrease protein translation using molecules such as antisense oligonucleotides (ASOs) or small interfering RNA (siRNA). ASOs are short, DNA-like molecules that can be designed to selectively bind mRNA transcripts and cause the catalytic destruction of particular mRNAs. ASOs that lower mRNA levels and thus decrease translation of proteins prone to misfolding show great promise in preclinical models of tauopathy, ALS, and HD, and are relatively safe in humans. These promising data have led to the launch of multiple clinical trials for ASOs in neurodegenerative diseases (NCT02519036, NCT03186989).

For RNA-targeted therapeutics, it is crucial to establish measurements that validate compound engagement with its desired biological target, mRNA. However, it is challenging to isolate mRNA from cerebrospinal fluid (CSF), the primary biofluid available to assay target engagement in the human central nervous system (CNS). Because mRNA translation leads to the production of new protein, protein-based biomarkers in CSF represent an attractive avenue to pursue target engagement for RNA-targeted therapeutics. In this study, we used nonradioactive, stable isotope labeling with an essential amino acid to identify inhibition of new protein production after ASO treatment. Using two preclinical models of ASO targets in clinical trials for neurodegenerative disease, we tested the hypotheses that stable isotope labeling with $^{13}$C$_6$-Leucine will: (1) identify pharmacodynamics at an earlier time point than protein concentration-based measures and (2) show measurable effects in CSF to support translation of this assay approach from animal models to human CNS. These results would support the use of new protein synthesis measurements in clinical trials for RNA-lowering therapies in neurodegenerative diseases.

**Methods**

**Transgenic animals**

Transgenic mice expressing human tau (hTau) (Jackson Laboratories, Bar Harbor, ME) were maintained through in-house breeding on a C57Bl6 background. Transgenic rats expressing human superoxide dismutase-1 (hSOD1) were provided by Pak Chan (Stanford University) and maintained through in-house breeding on a Sprague-Dawley background. Genotyping was performed by PCR amplification of tail DNA (Table S1).

All breeding and experimental protocols were approved by the Institutional Animal Care and Use Committee at Washington University and conducted according to the NIH Guide for Care and Use of Laboratory Animals.

**Antisense oligonucleotides (ASOs)**

ASOs that recruit RNase-H to degrade target mRNA and an inactive, control ASO with no sequence specificity were provided by Ionis Pharmaceuticals (Fig. 1). All ASOs were diluted in artificial CSF before use.

**Surgical procedures**

All animals were anesthetized by 2%–5% inhalant isoflurane, received constant isoflurane flow during the procedure, and were given an injection of bupivacaine prior to incision. Following the surgical procedure, animals received carprofen and were placed on a 37°C heating pad for recovery.

For intracerebroventricular injections, 60-day-old hTau transgenic mice received 300 µg of ASO at −1.0 mm M/L, −0.3 mm A/P, −3.0 mm D/V from Bregma. Similarly, 120-day-old hSOD1 transgenic rats received 1000 µg of ASO injected at −1.4 mm M/L, −0.4 mm A/P, −3.5 mm D/V from Bregma. For intrathecal ASO injection, 120-day-old hSOD1 transgenic rats received 1000 µg of ASO between the L3 and L5 vertebrae. Both male and female transgenic animals were used in these studies, and researchers were blinded to the identity of treatment throughout the duration of the experiment.

**$^{13}$C$_6$-Leucine oral labeling and tissue collection**

Methods for oral labeling in rodents with $^{13}$C$_6$-Leucine were described previously using unlabeled L-leucine or $^{13}$C$_6$-L-Leucine (Cambridge Isotope Laboratories, Cambridge, MA) dissolved in water at 5 mg/mL. The amount of $^{13}$C$_6$-Leucine found in plasma, the precursor for leucine found in the central nervous system, was consistent in all experiments between groups (Figure S1).

At indicated time points after surgery, animals were anesthetized with isoflurane and perfused with cold phosphate buffered saline (PBS) containing 0.03% heparin. Before perfusion, CSF was extracted from rat cisterna magna via a 23G “winged infusion butterfly needle (Terumo, Somerset, NJ). Blood was collected from mechanically ruptured vena cavae at the beginning of perfusion, and the brain and spinal cord were harvested after perfusion. All samples were flash frozen in liquid nitrogen and stored at −80°C before use.
mRNA isolation and analysis

mRNA isolation from flash-frozen tissues was performed using the RNeasy® Mini Kit (Qiagen, Venlo, Netherlands). The EXPRESS One-Step Superscript qRT-PCR Universal Kit (Invitrogen, Carlsbad, CA) was used for reverse transcription and amplification of SOD1 and MAPT mRNA transcripts. Comparative analysis by the DD<sub>Ct</sub> method, with GAPDH mRNA transcripts as an internal control, was performed with the ABI PRISM 7500 Fast Real-Time System (Applied Biosystems, Waltham, MA) (Table S1).

N15 Recombinant Protein Standards

N15-labeled, recombinant hTau protein (isoform 2N4R 411) was a generous gift from Drs. Isabelle Huvent and Guy Lippens (Université des Sciences et Technologie de Lille 1, France).

Recombinant hSOD1 was produced in Rosetta 2 E Coli (Novagen) using an hSOD1 cDNA construct with an N-terminal GST tag subcloned into a pGEX4T-1 backbone. Bacterial cultures were inoculated overnight in 15N-Cel- tone Complete Medium (Cambridge Isotope Laboratories, Inc) with shaking at 37°C and induced with 1 mmol/L IPTG for hSOD1 production. Protein was isolated by incubation with 600 µL of glutathione sepharose 4B beads (GE Healthcare) for 30 min, 1 unit of thrombin (Thrombin Cleavage Capture Kit, Millipore) for 20 h, and 30 µL of streptavidin agarose for 30 min, performed at room temperature. Samples were spun at 1000xg for 5 min, and supernatant containing N15-labeled, hSOD1 recombinant protein was collected.

Tissue homogenization, protein isolation, and mass spectrometric analysis

hTau transgenic mouse brain tissue was homogenized in PBS plus protease inhibitors (P8340; Sigma-Aldrich, St. Louis, MO). Protein concentration of brain lysate was determined by Pierce BCA assay (ThermoFisher Scientific, Waltham, MA). Total hTau was immunoprecipitated from 100 µg of tissue lysate using 30 µL of CNBr-activated Sepharose beads (GE Healthcare, Chicago, IL) crosslinked to anti-hTau antibody (Tau 1 monoclonal, produced by Nicholas M. Kanaan, Michigan State University) at 3 mg antibody/g of beads in 1 mL PBS plus 1% NP-40, 5 mmol/L guanidine, and protease inhibitors for 1.5 h at room temperature. Before immunoprecipitation, 5 ng of N15-labeled hTau was added to each sample. hTau was digested using 400 ng of sequencing-grade endoproteinase Trypsin (Promega, Madison, WI) for 16 h at 37°C. Samples were lyophilized and reconstituted in 25 µL of 2% acetonitrile/0.1% formic acid solution prior to liquid chromatography-triple quadrupole mass spectrometry analysis (Xevo TQ-S, Waters).

Newly synthesized protein rate was quantified by analyzing the incorporation of 13C<sub>6</sub>-Leucine into target proteins, indicated by the tracer-to-tracee ratio (TTR), by comparing the area under the chromatogram curve for SOD1 or tau peptides containing 13C<sub>6</sub>-Leucine or 12C<sub>6</sub>-Leucine. Relative protein concentration was quantified by measuring the ratio between the area under the curve of protein immunoprecipitated from animal tissue and N15-labeled recombinant protein (Tables S2 and S3). To account for non-steady state conditions due to mRNA lowering in this experiments, newly generated protein was also analyzed by multiplying the relative protein concentration by protein TTR at each time point. For figure

Figure 1. Antisense Oligonucleotides. Oligonucleotides containing phosphothioate backbone modifications: unmodified phosphodiester linkage (red), 2’-O-methoxyethylribose, (orange) and constrained ethyl (blue) groups. All cytosine residues are 5’methylcytosine.
presentation, data for hTau is represented by the TPSLPTPPPTR peptide and data for hSOD1 is represented by the GLHGPHVHE peptide, as all peptides measured showed strong correlations within each sample (Fig. S2 and S3).

Plasma-free $^{13}$C$_6$-Leucine abundance was analyzed from TCA precipitation of 100 µL rat or mouse plasma via capillary gas chromatography-negative chemical ionization-quadrupole mass spectrometry (Agilent) as described previously.$^{24}$

**Statistical analysis**

All statistical analyses were performed using Graphpad Prism 7.0 (GraphPad Software). All hypothesis tests performed were two-sided assuming a normal distribution of data. $P < 0.05$ was determined significant. Based on previous results,$^{10,21}$ a cutoff of 50% target mRNA reduction relative to the mean mRNA levels in inactive ASO-treated animals was used as inclusion criteria for subsequent analysis. In CSF analysis, samples were excluded if blood contamination was visibly present or if the volume of CSF collected was less than 75 µL. For relative quantification, samples were normalized to a single sample in each experiment from the inactive ASO treatment group. Quantitative results are expressed as mean difference and 95% confidence interval.

**Results**

**Lowering new hTau production precedes protein concentration changes after ASO treatment**

We performed ICV injection of an ASO targeting human tau mRNA$^{10}$ in transgenic mice expressing the human tau transgene (hTau mice) to understand lowering of tau mRNA and total protein over time. To analyze newly generated hTau protein, we administered $^{13}$C$_6$-Leucine in drinking water at designated time points (Fig. 2A). We organized the timing of ASO treatment, $^{13}$C$_6$-Leucine administration, and euthanasia to observe consistent, maximum label incorporation of proteins within the CNS at each time point analyzed. Preliminary studies showed adequate labeling of hTau with 4 days of $^{13}$C$_6$-Leucine labeling (Fig. S4).

hTau ASO treatment lowered hTau mRNA in the cerebral cortex over time (Fig. 2B). As early as 2 days post-injection, hTau ASO treatment lowered hTau mRNA levels by ~50% relative to inactive ASO, with further change of hTau mRNA by ~72% by 23 days post-injection. hTau ASO treatment also decreased hTau protein levels over time (Fig. 2C), with significant effects of ~32% observed at 23 days. The discrepancy between overall mRNA lowering and protein lowering is consistent with tau as a long-lived protein in the CNS.$^{25}$

Using percent of $^{13}$C$_6$-Leucine incorporation (TTR) as a measure of new protein synthesis rate, hTau ASO treatment lowered the rate of newly synthesized hTau protein over time (Fig. 2D). We observed lowering of newly generated hTau protein at 13 days after ASO treatment by ~71%. Combined, these data suggest that newly generated hTau protein is a reliable measure of hTau ASO pharmacodynamics in central nervous system tissue, with newly generated hTau showing an earlier (13 vs. 23 days) and greater magnitude of effect (71% vs. 32%) compared to hTau protein concentrations (Fig. 2E).

**Lowering of $^{13}$C$_6$-Leucine-labeled hSOD1 occurs in CSF after ASO treatment**

hSOD1$^{WT}$ rats were treated with an ICV injection of 1000 µg hSOD1 ASO in the lateral ventricle and administered $^{13}$C$_6$-Leucine in drinking water (Fig. 3A). Based on previous measurements of hSOD1 half-life in these rats,$^{24}$ and that lowering of CSF SOD1 after ASO treatment in the G93A SOD1 transgenic rat model was observed at 42 days post-ASO treatment,$^{26}$ we euthanized animals 30 days after treatment to assay hSOD1 total protein levels and new hSOD1 protein production to identify early ASO pharmacodynamics. hSOD1 ASO treatment lowered hSOD1 mRNA globally within the CNS (Fig. 3B), leading to overall hSOD1 protein lowering by ~62% in cortex and ~40% in lumbar spinal cord (Fig. 3C). Similarly, we observed a decrease in new hSOD1 protein synthesis rate by ~34% in cortex and ~50% in lumbar spinal cord (Fig. 3D). At this time point, lowering of both hSOD1 protein concentration and new hSOD1 protein synthesis rate drive overall decreases in newly generated hSOD1 protein (Fig. 3E).

In CSF, hSOD1 treatment failed to lower hSOD1 protein concentration (Fig. 3F). However, SOD1 ASO treatment decreased $^{13}$C$_6$-Leucine-labeled hSOD1 synthesis in CSF by ~30% relative to the inactive ASO treatment group (Fig. 3G). This effect of lowered $^{13}$C$_6$-Leucine-labeled hSOD1 is specific to CSF, as we did not measure any differences in peripheral hSOD1 protein in plasma (Fig. S5). To test SOD1 protein concentration at a later time point, we treated a separate cohort of animals without $^{13}$C$_6$-Leucine labeling using the same ASO treatment paradigm and analyzed CSF hSOD1 concentration at 50 days post-surgery (Fig. S6). Although we observed lowering in hSOD1 mRNA throughout the CNS 50 days post-ASO treatment (Fig. S6A), we did not observe decreases in CSF hSOD1 protein concentration.
(Fig. S6B), suggesting an even longer incubation time may be required to observe protein lowering in CSF. Taken together, these data suggest changes in newly synthesized, $^{13}$C$_6$-Leucine-labeled target proteins drive early pharmacodynamics biomarker effects in the CSF after ASO treatment (Fig. 3H).
A. Schematic diagram of experimental design:

- **hSOD1$^{\text{WT}}$**
- Diet Acclimation (7 d) | Label (7 d) | Chase (3 d)

B. Bar chart showing normalized hSOD1 mRNA levels:

- Inactive ASO
- SOD1 ASO

C. Bar chart showing normalized hSOD1 protein levels:

- Cortex
- Spinal Cord

D. Bar chart showing newly generated hSOD1 (TTR):

- Frontal Cortex
- Spinal Cord

E. Bar chart showing newly generated hSOD1 (Conc. x TTR):

- Cortex
- Spinal Cord

F. Bar chart showing normalized hSOD1 protein levels:

- Inactive ASO
- SOD1 ASO

G. Bar chart showing newly generated SOD1 (TTR):

- Cerebrospinal Fluid

H. Bar chart showing newly generated hSOD1 (Conc. x TTR):

- Inactive ASO
- SOD1 ASO

**Legend:**

- $^{13}$C$_6$-Leucine
- $^{12}$C$_6$-Leucine

**Statistical Symbols:**

- **p = 0.06**
- **p < 0.001**
- **p < 0.0001**

**Notes:**

- The experiments were conducted with appropriate controls and statistical analyses to validate the findings.
A therapeutic window exists for newly synthesized protein pharmacodynamics

To enhance the translational relevance of our findings, we performed similar experiments in hSOD1 rats using intrathecal ASO injection, the method of drug delivery used in current clinical trials. We analyzed CNS tissue 10, 30, and 60 days after ASO treatment compared to inactive ASO control to better understand longitudinal changes in hSOD1 protein (Fig. 4A). Contrary to hSOD1 ASO delivery by ICV injection, we observed differential effects of hSOD1 mRNA lowering in CNS tissues after intrathecal ASO injection. We observed a significant hSOD1 mRNA decrease in the lumbar spinal cord over time, with an average hSOD1 mRNA lowering at 60 days by ~66%. However, hSOD1 treatment did not significantly lower hSOD1 mRNA in the cortex or cerebellum over time (Fig. 4B). Because it is likely that SOD1 mRNA will need to be lowered globally in the CNS to affect SOD1 CSF levels, we focused on protein measurements in the spinal cord, where we confidently documented mRNA lowering.

To further understand the relationship between protein lowering and lowering of hSOD1 synthesis rate after ASO treatment, we analyzed hSOD1 protein in lumbar spinal cord. hSOD1 ASO treatment decreased hSOD1 protein levels across all time points measured, and treatment effect was greatest at day 60, with 50% lowering of hSOD1 protein (Fig. 4C). hSOD1 ASO treatment significantly lowered hSOD1 synthesis rate compared to inactive ASO-treated animals (Fig. 4D). Interestingly, ASO treatment decreased the ratio of newly synthesized hSOD1 at 10 and 30 days by 40% and 45%, but there was no difference in 13C6-Leucine-labeled hSOD1 between groups at day 60 despite lowering of total hSOD1 protein concentration. Therefore, at later time points, decreases in newly generated hSOD1 protein by ASO treatment are driven by overall lowering of protein concentration (Fig. 4E).

Discussion

Our data show that, following treatment with a mRNA lowering therapeutic, the lowering of newly synthesized protein occurs before lowering of total protein concentration in tissue. Further, our findings indicate that labeled, newly generated protein measures direct effects of protein translation in the source tissues, which are not seen in CSF protein concentrations at early time points. Given the direct measurement of protein translation, labeling protein production has higher sensitivity than protein concentration-based measures for mRNA-lowering therapies. These studies suggest that identifying newly generated protein after ASO treatment is an effective protein-based biomarker for assaying ASO target engagement (Fig. 5).

Previous work supports the use of CSF protein concentration as a promising pharmacodynamics biomarker for ASO therapy, and CSF SOD1 concentration is a secondary outcome measurement in an ongoing Phase I clinical trial (NCT0262399). However, a key limitation in the use of CSF protein concentration as a measure of ASO efficacy is the indirect mechanism informed by this assay. ASOs inhibit the production of new protein, and measures of protein concentration also rely on transport to CSF and the clearance of existing protein which may be unaffected by ASO treatment. Especially in a rapidly progressing neurodegenerative disease such as ALS, it is important to develop pharmacodynamics tools that identify direct mechanisms of action at early time points that reflect the actual drug effect in the tissue of interest.

Our study design is translatable to humans, as stable isotope labeling is safe and effective in measuring the kinetics of proteins implicated in neurodegenerative disease within human CSF. Previous studies used labeling with 13C6-Leucine to measure inhibition of amyloid beta peptide synthesis by a gamma secretase inhibitor in human participants. However, these experiments...
A

\[ \text{hSOD1}^{\text{WT}} \]

-60d  -30d  -10d

\[ \uparrow \]

\( \text{\textsuperscript{13}C}_\text{6}-\text{Leucine} \)
\( \text{\textsuperscript{12}C}_\text{6}-\text{Leucine} \)

Diet Acclimation (7 d)  Label (7 d)  Chase (3 d)

B

\text{Lumbar Spinal Cord}

\text{Frontal Cortex}

\text{Cerebellum}

\begin{align*}
\text{Normalized hSOD1 mRNA (a.u.)} \\
\text{Time After ASO Surgery (days)}
\end{align*}

\begin{align*}
\text{Normalized hSOD1 mRNA (a.u.)} \\
\text{Time After ASO Surgery (days)}
\end{align*}

\begin{align*}
\text{Normalized hSOD1 mRNA (a.u.)} \\
\text{Time After ASO Surgery (days)}
\end{align*}

\begin{align*}
\text{Inactive ASO} \\
\text{hSOD1 ASO}
\end{align*}

C

\text{Lumbar Spinal Cord}

\begin{align*}
\text{Normalized SOD1 Protein (a.u.)} \\
\text{Time After ASO Surgery (days)}
\end{align*}

\begin{align*}
\text{Normalized SOD1 Protein (a.u.)} \\
\text{Time After ASO Surgery (days)}
\end{align*}

\begin{align*}
\text{Normalized SOD1 Protein (a.u.)} \\
\text{Time After ASO Surgery (days)}
\end{align*}

D

\text{Lumbar Spinal Cord}

\begin{align*}
\text{\textsuperscript{13}C}_\text{6}-\text{Leu SOD1 (TTR)} \\
\text{Time After ASO Surgery (days)}
\end{align*}

\begin{align*}
\text{\textsuperscript{13}C}_\text{6}-\text{Leu SOD1 (TTR)} \\
\text{Time After ASO Surgery (days)}
\end{align*}

E

\text{Lumbar Spinal Cord}

\begin{align*}
\text{Newly Generated hSOD1 (Conc. x TTR)} \\
\text{Time After ASO Surgery (days)}
\end{align*}

\begin{align*}
\text{Newly Generated hSOD1 (Conc. x TTR)} \\
\text{Time After ASO Surgery (days)}
\end{align*}
were performed over a 36-hour time period, as the half-life of amyloid beta peptide is ~9 h in CSF.18 To our knowledge, the results presented here are the first observations measuring new protein production of long-lived protein targets in the CNS after therapeutic intervention. Our approach can be directly applied to multiple RNA-directed strategies, such as siRNAs31 and ASOs,32,33 that target long-lived proteins in neurodegeneration for clinical trials. Besides its lowered protein production readout, stable isotope labeling could be a powerful tool to examine increased protein production for therapeutic strategies such as gene therapy34 and splicing ASO treatments.35,36 Moreover, an understanding of protein kinetics using stable isotope labeling can also inform on assays for therapies that target protein turnover, such as approaches that activate protein clearance mechanisms37 or promote proper protein folding.38

Another advantage of our mass spectrometry-based assay is the ability to specifically isolate mutant from wild-type protein in patient populations. Many familial forms of neurodegenerative diseases are caused by autosomal dominant mutations31 and, thus, exhibit both mutant and normal protein forms. However, it is currently unknown if mutant proteins display altered protein kinetics compared with their wild type counterparts in humans. Future studies in symptomatic, mutation-carrying patients are needed to characterize the half-life of these target proteins in human CSF. Protein half-life measures will be crucial to determine the timing required to observe pharmacodynamics effects in humans when designing randomized-control clinical trials.

One limitation in this study is the inability to identify target protein lowering in CSF after SOD1 ASO treatment. Without this effect, the relationship between lowered protein production and total protein concentration cannot be determined in the biological fluid important for human clinical trials. Previous data suggest this limitation may be due to the drug delivery paradigm in these experiments. The current study used a single bolus injection of ASO in rodent models while previous studies that observed total target protein lowering in CSF after ASO treatment in nonhuman primate models were performed with multiple intrathecal ASO injections.10,21 Additionally, both transgenic animal models used here express wild-type forms of the target protein. In previous experiments, the G93A SOD1 rat model was used to demonstrate hSOD1 protein lowering after ASO treatment 42 days after treatment.26 Our group 24 and others 39 have shown that mutant hSOD1 exhibits a shorter protein half-life in the CNS and CSF compared to WT hSOD1. Taken together, these data suggest that CSF WT hSOD1 protein concentration will occur at an even later time point than 50 days after ASO treatment (Fig. S6). Future investigation should also address these questions in the presence of mutant protein implicated in disease, as no pathological misfolding of either hSOD1 or hTau is observed in the models employed here.19,20 Therefore, continuing experiments in transgenic rat models or nonhuman primates should examine the relationship between CSF protein half-life, protein concentration lowering, and changes in new protein production over time after ASO treatment, in addition to understanding how new protein production in CSF changes in the presence of misfolded proteins within the CNS.

In sum, the approach described here provides a framework for the development of assays that isolate the direct mechanism of action of target engagement in neurodegenerative disease therapies, with the hope that such data will lead to a higher success in clinical trial outcomes.

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Author Contributions
WKS, KMS, TMM, and RJB contributed to study conception and design of experiments. WKS, KMS, NJB, IGB, JA, and CS contributed to data acquisition and analysis. WKS, KMS, CS, HBK, TC, ES, RJB, and TMM contributed to the drafting of the text and editing of the manuscript.

Conflict of Interest
TMM has served on medical advisory boards for Biogen and Ionis Pharmaceuticals and is a consultant for Cytokinetics. TMM/RJB/Washington University have a licensing agreement with C2N Diagnostics. Washington University and RJB have equity ownership interest in C2N Diagnostics and may receive royalty income based on technology licensed by Washington University to C2N Diagnostics. RJB receives income from C2N Diagnostics for serving on its scientific advisory board. Washington University has submitted nonprovisional patent applications for “Methods for measuring the metabolism of CNS derived biomolecules in vivo” (RJB) and “SOD1 kinetics measurements” (RJB, TMM). TC, HBK, and ES are employees of Ionis Pharmaceuticals.

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Supporting Information

Additional supporting information may be found online in the Supporting Information section at the end of the article.

Figure S1. No differences in 13C6-Leucine oral labeling between treatment groups in all experiments performed. Plasma 13C6-Leucine values were analyzed in all samples analyzed and compared between groups. A. hTau ICV ASO plasma 13C6-Leucine at Day 23 post surgery, as seen in Figure 2. No differences were observed (Student’s T-test, $P = 0.4496$). B. hSOD1 ICV ASO plasma 13C6-Leucine at Day 30. No differences were observed (Student’s-T test) C. hSOD1 IT ASO plasma 13C6-Leucine at all time points assessed. No differences were observed (2-Way ANOVA). Error bars in figures represent SEM.
Figure S2. Correlation between all hTau peptides measured. Linear regression analysis comparing peptides shows strong correlations exist between all three peptide measurements. All p values < 0.0001 for each analysis.

Figure S3. Correlation between all hSOD1 peptides measured. Linear regression analysis comparing peptides shows strong correlations exist between all peptide measurements. All P values < 0.0001 for each analysis.

Figure S4. Characterizing 13C6-Leucine incorporation into hTau protein in hTau WT transgenic mice. hTau mice were acclimated to leucine-free chow for 7 days, and labeled with 5 mg/ml 13C6-leucine for 7 days. Mice were euthanized at 6 (n = 2), 10 (n = 3), 14 (n = 3), and 17 (n = 2) days post-end of label. Because of the high label incorporation (nearly 15% TTR) and delayed appearance of higher label incorporation after the end of the labeling period 3-10 days after the end of labeling, subsequent experiments reduced the label time to 4 days. Given maximal label incorporation at 6-10 days post-end of label, all mice in the subsequent experiments were sacrificed at 5 days post-end of label. Error bars in figures represent SEM.

Figure S5. No changes in peripheral hSOD1 after 1000 μg hSOD1 ASO intracerebroventricular injection. A. hSOD1 protein levels relative to N15-labeled recombinant standard by LC-MS in plasma. No changes in plasma hSOD1 protein levels were observed (Student’s T-test). B. 13C6-Leu labeled hSOD1 in plasma by LC-MS. No changes in plasma 13C6-Leu labeled hSOD1 was observed (Student’s T-Test). Error bars in figures represent SEM.

Figure S6. No protein concentration pharmacodynamics observed in CSF after 1000 μg hSOD1 ASO intracerebroventricular injection at 50 days post treatment. A. hSOD1 mRNA levels in the CNS by qPCR. hSOD1 mRNA levels are significantly lowered in spinal cord and frontal cortex by ~70% and 50%, respectively, after ASO treatment relative to inactive ASO controls (Student’s T-test). B. hSOD1 protein levels relative to N15-labeled recombinant standard by LC-MS. No changes in CSF hSOD1 protein levels were observed (Student’s T-test). n = 5 for each treatment group, with one CSF sample excluded from each group due to blood contamination. **** P < 0.0001. Error bars in figures represent SEM.

Table S1. Primers and probes used for genotyping and qPCR analysis of hTau (MAPT), hSOD1 (SOD1) and mouse GAPDH (GAPDH) DNA.

Table S2. Transition ions used for hSOD1 tandem LC-MS/MS. GLHGFHVHE peptide measurements were used as representative data in figures.

Table S3. Transition ions used for hTau tandem LC-MS/MS. TPSLPTPPTTR peptide measurements were used as representative data in figures.