Development of novel small molecules targeting neurotrophic HGF signaling for the treatment of Alzheimer's, Parkinson's, and ALS

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CONCLUSIONS

- Positive modulation of neurotrophic HGF signaling is a promising therapeutic strategy that may be beneficial in a range of neurodegenerative disorders
- ATH small molecules promote neurotrophic signaling, protect neurons, reduce protein pathology, and improve function in several models of neurodegenerative disorders
- Preclinical evidence supports continued development of ATH small molecule candidates fosgonimeton, ATH-1020, and ATH-1105 for the treatment of AD, PD, and ALS

KEY TAKEAWAY

The neurotrophic and neuroprotective effects of HGF positive modulators across diverse neurodegenerative models supports the broad therapeutic potential of this emerging class of compounds





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Abbreviations: 6-OHDA, 6-hyrdoxy dopamine; AD, Alzheimer's disease; α-syn, alpha synuclein; ALS, amyotrophic lateral sclerosis; AM, active metabolite; ATF6, activating transcription factor 6; Aβ, amyloid beta; CBE, conduritol β-epoxide; CMAP, compound muscle drug; LAMP2, lysosomal associated membrane protein 2; MAP2, microtubule associate protein 2; MPP+, 1-methyl-4 um; NeuN, neuronal nuclei; PD, Parkinson's disease; PFF, pre-formed fibril; PM, positive modulator; SC, subcutaneous injection; **SOD1**, superoxide dismutase 1; **TDP43**, transactive response DNA binding protein; **TH**, tyrosine hydroxylase

References: .1. Dugger and Dickson. Cold Spring Harbor Perspect Biol. 2017;9(7).2 Gontijo et al. Current Neuropharm. 2020;18(5):348-407 3. Desole et al, Font Cell Dev Biol. 2021;9, 683609. 4. Johnston JL et al. Neurotherapeutics. 2023;20(2):431-451.

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Disclosures

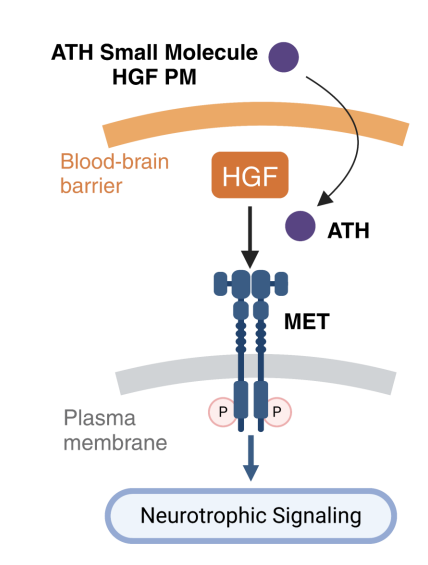
Kevin Church, Sherif Reda, Andrée-Anne Berthiaume, Sharay Setti, Kayla Kleist, Wei Wu, Robert Taylor, Jewel Johnston are employees and stockholders of Athira Pharma, Inc.

Fosgonimeton, ATH-1020, and ATH-1105 are investigational therapies and have not received FDA approval nor been demonstrated to be safe or effective for any use.

INTRODUCTION

- Neurodegenerative diseases such as AD, PD, and ALS present diverse symptomology, yet they share common pathological themes, including increased neuroinflammation, protein accumulation, and oxidative, lysosomal, and mitochondrial stress ultimately leading to the death of neurons1.
- The complex pathology involved in neurodegenerative disorders suggests that multifactorial treatment strategies may be required for disease modification^{2,3}.
- We have developed a series of small molecule positive modulators (PM) of the neurotrophic HGF system (ATH small molecules) capable of promoting neuroprotective and neurotrophic signaling that may counteract neurodegeneration in a range of diseases⁴.

Figure 1: ATH small molecules positively modulate the neurotrophic HGF signaling system to promote neuroprotective signaling pathways to counteract neurodegenerative processes



ATH small molecule HGF PMs promote the phospho-activation of the MET receptor in the presence of endogenous HGF ligand.

ATH-1020

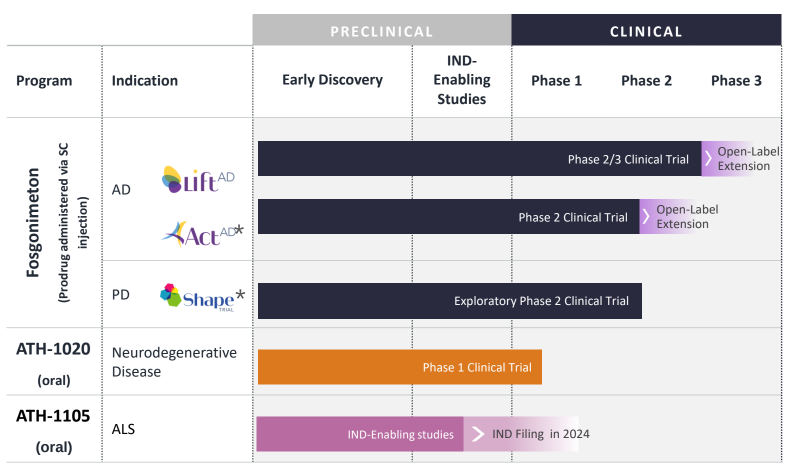
ATH-1105

ATH small molecules presented here distribute to the CNS.

ATH HGF PMs in the presented studies:

- Prodrug delivered via SC injection rapidly converts to active metabolite fosgo-AM
- ATH-1020 Orally bioavailable
- ATH-1105
- Orally bioavailable

Figure 2: ATH small molecules are under clinical investigation or development for the treatment of multiple neurodegenerative diseases



*These trials are complete with topline data reported in June 2022 and December 2023 respectively

METHODS



- MET activation: HEK293 cells were treated with the indicated ATH molecule in vehicle containing a low dose of HGF (1 ng/ml). MET phosphorylation (Y1234/1235) was quantified via ELISA
- Neurotrophic activity: Primary hippocampal neurons isolated from newborn rats were treated with the indicated ATH compound, and in some cases in the presence of exogenous HGF protein for 3-4 days (neurite assay) or 8-9 days (synapse assay). Cultures were immunostained against βII-tubulin to measure neurite length and synaptobrevin II to count
- AD models: Primary cortical neurons isolated from embryonic rats were incubated with Aβ₁₋₄₂ oligomers for 24 hours and assessed for survival, accumulation of pTau, and mitochondrial stress by automated image acquisition of cultures immunostained for MAP2, AT100, and MitoSox respectively.
- PD models: Primary cultures of rat dopaminergic neurons were treated with α-synuclein PFF. Neuron survival and lysosomal stress were assessed by automated image acquisition of cultures immunostained for MAP2 and LAMP2 respectively. Neuron survival was assessed by cell titer glow.
- ALS models: Primary motor neurons with a SOD1^{G93A} genotype in culture were treated with glutamate to induce excitotoxicity. Neuron survival, ER stress, and mitochondrial stress were assessed by immunostaining for MAP2, ATF6, and MitoTracker respectively.



used to produce PD-related deficits. Grip strength was measured after 6 weeks of treatment.

In Vivo

- AD model: Aβ_{1.42} oligomers were administered via intrahippocampal injection to induce neuron death and cognitive impairment in mice. After 7 days of treatment with fosgonimeton, cognitive performance was evaluated in the Y-maze. After
- 28 days of treatment, neuron survival was assessed by quantifying NeuN+ cells in the CA1 layer of the hippocampus. PD models: In one model, Parkinson's disease-related phenotypes were elicited by single intranigral injection of αsynuclein and CBE (lysosomal disruptor) administration 3 times per week in mice. Motor function was assessed by the ladder test after 6 weeks of treatment. α-synuclein area in dopaminergic neurons was assessed in the substantia nigra via
- ALS model: Transgenic Prp-TDP43A315T mice were used as a model of ALS. Motor function was assessed by latency to fall in rotarod, and nerve function was assessed by CMAP-amplitude after 2 months of treatment. In a separate study, survival was assessed in animals treated with ATH-1105 from 1-5 months of age.

immunostaining for tyrosine hydroxylase (TH) and α-synuclein. In another model in rats, unilateral injection of 6-OHDA was

RESULTS

ATH small molecules enhance neurotrophic HGF signaling

ATH small molecule

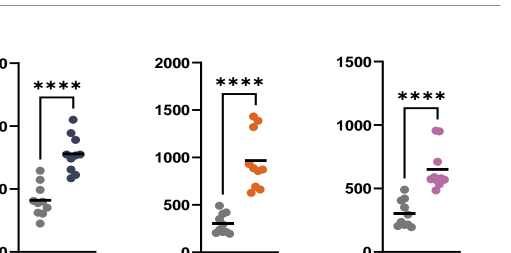
Fosgonimeton

ATH-1105

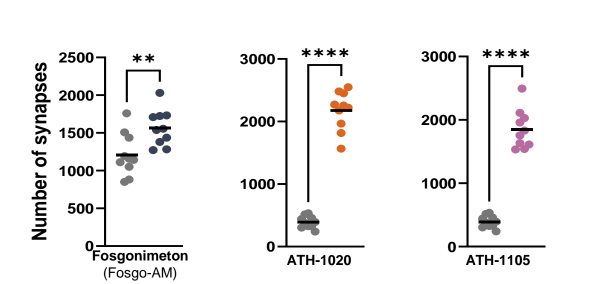
Figure 3: ATH small molecules promote MET phosphorylation and stimulate neurotrophic effects

Neurite Outgrowth MET Activation

ATH small molecules enhance MET activation (phosphorylation) in HEK cells after 15 minutes of treatment. Fosgo-AM: 0.01 nM, ATH-1020: 1 nM, ATH-1105: 0.1 nM. Statistics: Student's t-test, * p< 0.05, * p<0.01, *** p< 0.001.



ATH small molecules promote extension of neurites in cultured primary hippocampal neurons after 3-4 days treatment. Fosgo-AM: 10 nM, ATH-1020: 1 nM, ATH-1105: 1 nM. Statistics: Student's t-test, * p< 0.05, * p<0.01

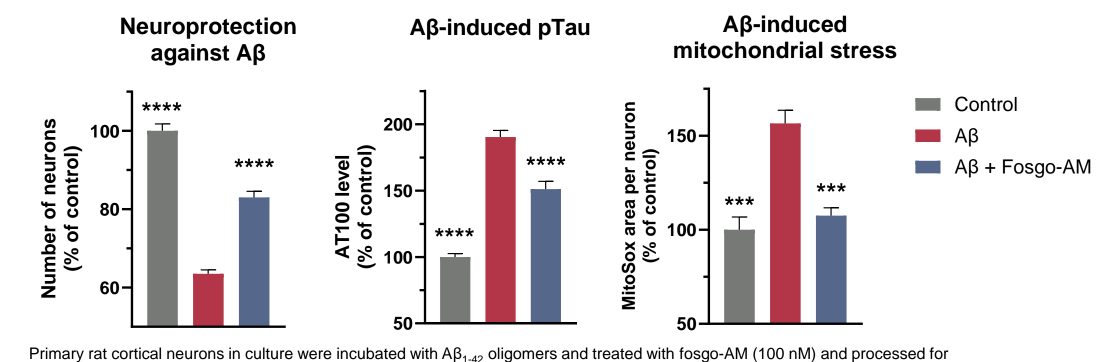


Synapse Count

ATH small molecules promote the formation of synapses in cultured primary hippocampal neurons after 8-9 days of treatment. Fosgo-AM: 1 nM, ATH-1020: 1 nM. ATH-1105: 1 nM. Statistics: Student's t-test, * p< 0.05, * p<0.01,

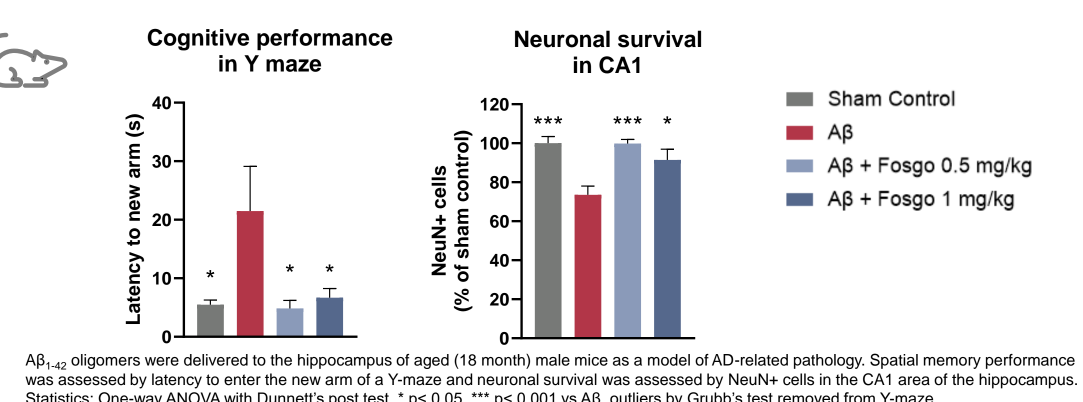
Fosgonimeton in preclinical models of AD

Figure 4: Fosgo-AM is neuroprotective and reduces pTau and mitochondrial stress in AD models in vitro



imaging with the following assessments: Neuron survival by number of MAP2+ cells, pTau accumulation by AT100 staining, and mitochondrial stress by MitoSox. Statistics: One-way ANOVA with Fisher's LSD, *** p< 0.001, **** p < 0.0001 vs Aβ

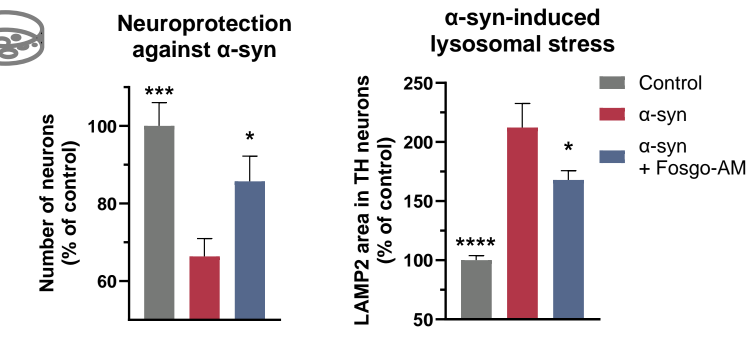
Figure 5: Fosgonimeton improves cognitive performance and protects hippocampal neurons in animals challenged with Aβ1-42



Statistics: One-way ANOVA with Dunnett's post test, * p< 0.05, *** p< 0.001 vs Aβ, outliers by Grubb's test removed from Y-maze.

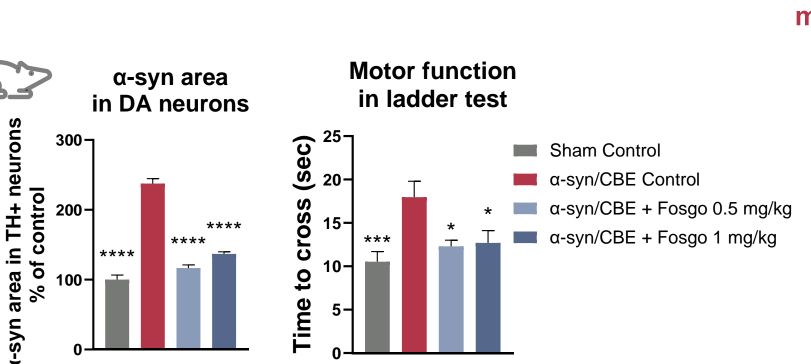
ATH small molecules in preclinical models of PD

Figure 6: Fosgonimeton promotes neuron survival and reduces lysosomal stress in vitro and improves DA neuron survival and motor function in vivo



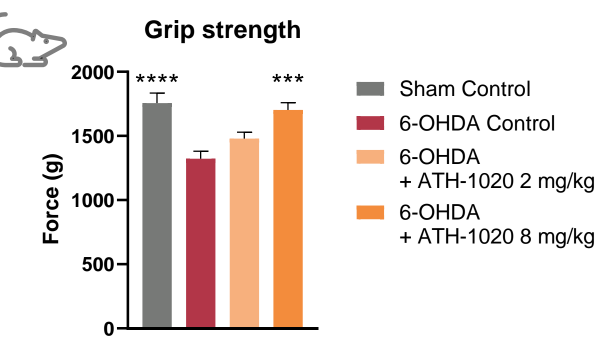
α-synuclein PFF were used to model PD-related phenotypes in primary dopaminergic neuron culture. Treatment effects of fosgo-AM (100 nM) were assessed in imaging experiments using TH+ cell counts and staining for LAMP2. Statistics: One-way ANOVA with Fisher's LSD, * p< 0.05 *** p< 0.001, **** p< 0.0001 vs α-syn

* p< 0.05, *** p< 0.001, **** p < 0.0001 vs glutamate



Aged (18 month) mice received a bilateral intranigral injection of α-synuclein PFF and injections of CBE 3 times a week to model PD-related phenotypes. α-synuclein area was determined in TH+ neurons in the substantia nigra. Motor behavior was assessed in the ladder test. Statistics: One-way ANOVA with Dunnett's post test, * p< 0.05, *** p< 0.001. **** p< 0.0001 vs α-syn

Figure 7: ATH-1020 improves grip strength in the 6-OHDA model of PD



6-OHDA was delivered to the striatum of male rats to produce PD-like pathology. Preclinical efficacy of ATH-1020 via oral gavage was assessed by measurement of grip strength. Statistics: One-way ANOVA with Dunnett's post test, *** p< 0.001, **** p< 0.0001 vs 6-OHDA

ATH-1105 in preclinical models of ALS

Figure 8: ATH-1105 promotes neuronal survival, reduces ER stress, and improves mitochondrial function following excitotoxic glutamate injury

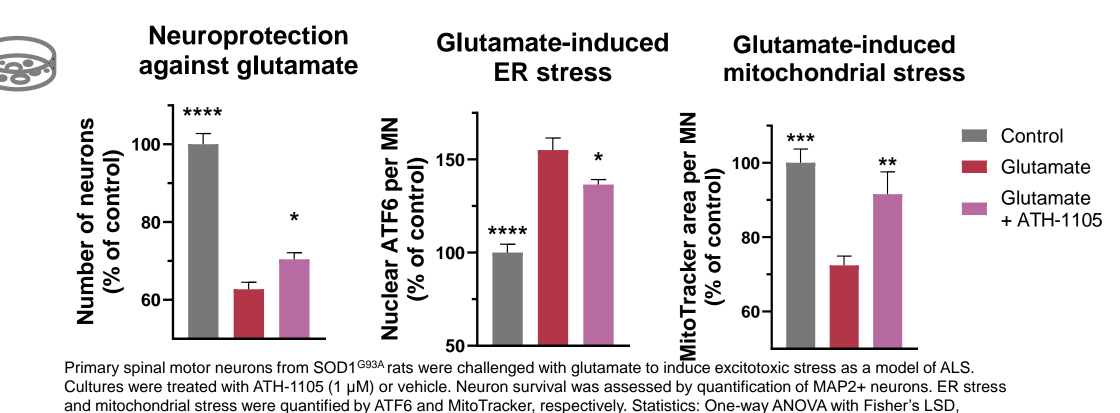
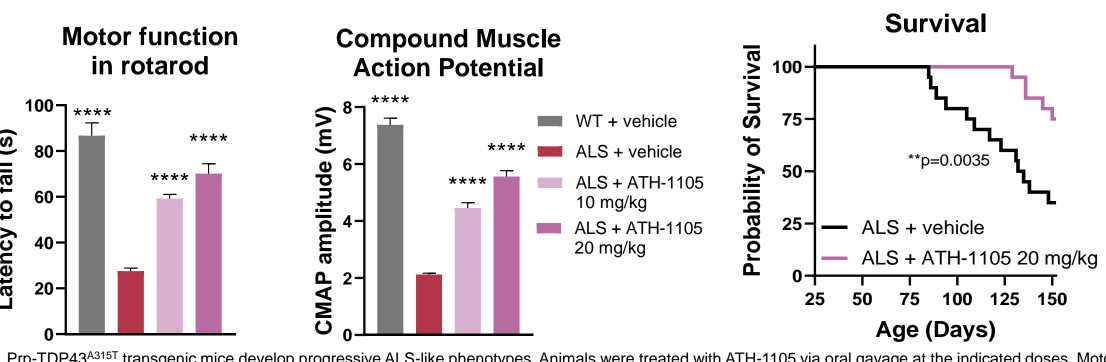


Figure 9: ATH-1105 improves motor function, nerve function, and survival in the Prp-TDP43^{A315T} transgenic mouse model of ALS.



Prp-TDP43^{A315T} transgenic mice develop progressive ALS-like phenotypes. Animals were treated with ATH-1105 via oral gavage at the indicated doses. Motor behavior was assessed by latency to fall in rotarod, and nerve function was assessed by measuring the CMAP amplitude between the sciatic nerve and the muscles of the foot. Statistics: One-way ANOVA with Dunnett's post test, **** p < 0.0001 vs ALS + vehicle. In a separate experiment, survival was assessed in animals treated with ATH-1105 from 1 to 5 months of age. Statistics: Log-rank (Mantel-Cox) test.