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3 **Title:** Novel HDAC6 inhibitors increase tubulin acetylation and rescue axonal transport of
4 mitochondria in a model of Charcot-Marie-Tooth Type 2F.
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ABSTRACT

Disruption of axonal transport causes a number of rare, inherited axonopathies and is heavily implicated in a wide range of more common neurodegenerative disorders, many of them age-related. Acetylation of α -tubulin is one important regulatory mechanism, influencing microtubule stability and motor protein attachment. Of several strategies so far used to enhance axonal transport, increasing microtubule acetylation through inhibition of the deacetylase enzyme HDAC6 has been one of the most effective. Several inhibitors have been developed and tested in animal and cellular models but better drug candidates are still needed. Here we report the development and characterisation of two highly potent HDAC6 inhibitors, which show low toxicity, promising pharmacokinetic properties, and enhance microtubule acetylation in the nanomolar range. We demonstrate their capacity to rescue axonal transport of mitochondria in a primary neuronal culture model of the inherited axonopathy Charcot-Marie-Tooth Type 2F, caused by a dominantly acting mutation in heat shock protein beta 1.

Keywords: HDAC6; CMT; Axonal transport; Mitochondria, Axonopathy, α -tubulin.

INTRODUCTION

The bidirectional movement of macromolecules and organelles along axons is essential for axon survival and function, and requires a complex machinery involving motor proteins, adapters coupling to specific cargoes, microtubule tracks and regulators of all the above. Not surprisingly, such an essential process involving many components can malfunction in a number of ways and when it does the consequences can be profound. Mutation of genes encoding axonal transport machinery and regulators cause a number of axonopathies, and especially diseases of long axons. For example, mutations in KIF5A encoding a major anterograde motor protein are an established cause of hereditary spastic paraplegia SPG10

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3 ¹, and have also been linked to Charcot-Marie-Tooth Disease type 2 (CMT2) ², amyotrophic
4 lateral sclerosis (ALS) ^{3,4} and neonatal intractable myoclonus ⁵. Mutant dynactin causes motor
5 neuron disease and distal spinal and bulbar muscular atrophy ^{6,7} and in mice the mutation of
6 tubulin chaperone protein Tbc1e causes a severe, early onset loss of motor axons with a major
7 deficiency of microtubules ⁸.

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14 Charcot-Marie-Tooth Disease type 2 (CMT2) is an axonal, non-demyelinating peripheral
15 neuropathy characterized by distal muscle weakness and atrophy, mild sensory loss, and
16 normal or near-normal nerve conduction velocities ⁹. The Charcot-Marie-Tooth disease
17 subtype 2F (CMT2F) and distal hereditary motor neuropathy subtype 2B (dHMN2B) are
18 caused by autosomal dominantly inherited mutations in the small heat shock protein B1
19 (*HSPB1*) gene ^{10,11}. The gene codes for heat shock protein beta-1 (HSPB1, also known as
20 HSP27), which is a member of the small heat shock protein family comprising a highly
21 conserved α -crystalline domain. HSPB1 acts as a chaperone binding with partially denatured
22 proteins to prevent aggregation ^{12,13}. Up to now, 18 mutations in *HSPB1* have been linked to
23 CMT2F and 27 mutations to dHMN 2 ¹⁴. The S135F and P182L mutations are among the best
24 characterized mutations so far ^{11,15,16}. The S135F mutation is the only one that causes both
25 CMT2 and dHMN2B. P182L mutation is associated only with dHMN2B ¹⁵. The S135F mutation
26 is located in the α -crystallin domain while P182L mutation lies in the short C-terminal tail of
27 the protein ¹⁵. Interestingly the localization of the mutation was shown to have different effects
28 on the protein function. While the S135F mutation caused the protein to increase its chaperone
29 activity accompanied with an increased in its monomeric state the chaperone activity of
30 HSPB1 was not affected by the P182L mutation ¹⁷.

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51 Four mutant *HSPB1* transgenic mouse models of CMT2F and/or dHMN2B have been
52 developed so far, which partially recapitulate the hallmarks of peripheral neuropathy ^{11,18-20}.

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S135F and P182L transgenic mice generated by d'Ydewalle et al. demonstrated noticeable
phenotypes, the latter presents more like dHMN2B than CMT2F with a lack of sensory

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3 symptoms ¹¹ which recapitulates all key features of CMT2F or distal HMN2B, dependent on
4 the mutation. However, CMT2F mouse models generated by other groups had notable
5 differences. S135F transgenic mice reported by Lee et al. had no sensory phenotype and
6 presented only a strict motor loss, similar to the P182L, but not the S135F mice of d'Ydewalle
7 et al ¹⁸. In further contrast, the R136W mouse model did not demonstrate any functional or
8 behavioral deficits ²⁰. When R127W and P182L mutant proteins were expressed at
9 physiological levels to alleviate concerns of artifacts due to overexpression, no pathology and
10 behavioural deficits were found in mice ¹⁹. This could be due to insufficient expression of
11 HSPB1 under the ROSA26 locus.
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15 In addition to rare disorders and animal models, axonal transport deficiency is heavily
16 implicated in many more common neurodegenerative and axonal disorders. Several cancer
17 chemotherapeutics that cause peripheral neuropathy as a dose-limiting complication target
18 microtubules ²¹ and disrupt axonal transport ²². In Alzheimer's disease, aggregation of
19 microtubule associated protein tau, whose normal functions include regulation of microtubule
20 stability and motor protein attachment ^{23, 24} plays a prominent role, exogenously applied A β ₁₋₄₂
21 is able to disrupt axonal transport in a tau-dependent manner ²⁵. The two may also interact
22 ²⁶ and impairment of axonal transport exacerbates animal models ²⁷. There are many
23 indications of a wider role also in ALS ²⁸, Huntington's disease ²⁹, Parkinsonism and
24 frontotemporal dementia ^{30, 31}, and normal ageing, the biggest risk factor in each of these ³²,
25 is accompanied by a twofold decline in axonal transport ³³. Thus, rare but often better-
26 understood inherited disorders involving an axonal transport mechanism are an important
27 starting point to develop therapies that could have far wider application in neurodegenerative
28 disease.
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32 Axonal microtubules exist in a state of dynamic instability ^{34, 35}, constantly both growing and
33 severing to maintain them typically between 0.15-20 μ m in length ³⁶. Acetylation of α -tubulin
34 at Lys40 is reported to increase microtubule stability under mechanical stress ³⁷ and to
35 influence severing by katanin ³⁸. It also enhances the binding of kinesin-1 and axonal transport
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3 ^{39, 40}. Beside SIRT2 ⁴¹ HDAC6 is the other major deacetylase for α -tubulin and its inhibition
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5 increases axonal transport of some cargoes in models of Charcot-Marie-Tooth disease types
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7 2F ¹¹ and 2D ⁴², ALS ⁴³, and Vincristine neuropathy ⁴⁴, also alleviating some symptoms.
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9 Beneficial outcomes have also been reported in models of Alzheimer's disease ⁴⁵ and stroke
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11 ⁴⁶. Early studies of HDAC6 inhibition used tubacin, whose high lipophilicity and short *in vivo*
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13 half-life limited its usefulness. This was largely superseded by the development of Tubastatin
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15 A ⁴⁷. However, further improvement of potency is possible ^{48, 49} so it is important to develop
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17 new compounds targeting HDAC6 with greater potential for clinical application.
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21 The HDAC inhibitors share a well-recognized pharmacophore that consists of three parts: a
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23 zinc binding group (ZBG), a linker, and a cap moiety. Classical HDAC inhibitors typically have
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25 the hydroxamic acid moiety as ZBG but the hydroxamic acid causes poor pharmacokinetics,
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27 low selectivity profiles, and production of active metabolites ⁵⁰. These features of hydroxamic
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29 acid are red flags for drug discovery in chronic diseases that are not life threatening. Therefore,
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31 we focussed here on the discovery of non-hydroxamic acid derivatives. High throughput
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33 screening (HTS) with a Takeda internal library provided several non-hydroxamic acid
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35 derivatives as hit compounds against HDAC6. By our medicinal chemistry efforts, we
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37 developed two compounds T-3796106 and T-3793168 that are highly selective for HDAC6,
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39 show CNS penetration and low toxicity both *in vivo* and *in vitro*. We report their dose-response
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41 effects for α -tubulin acetylation in primary neuronal cultures and their influence on axonal
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43 transport of mitochondria in a primary culture model of CMT2F.
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49 RESULTS

50 Evaluation of inhibitory potencies (IC₅₀) of T-3796106 and T-3793168

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52 The inhibitory potencies (IC₅₀) of T-3796106 and T-3793168, which were developed through
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54 medicinal chemistry campaign from HTS hit compounds, were evaluated in HDAC panel assay
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56 (Table 1). T-3796106 showed potent inhibitory activity against HDAC6 with the IC₅₀ value of
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12 nM. IC₅₀ values for HDAC3, HDAC8, HDAC5, HDAC7, and HDAC9 were in the range of 1,000-3,000 nM. IC₅₀ values for HDAC1 and HDAC4 were over 6,000 nM. T-3796106 did not show inhibitory activity against HDAC2, HDAC10, and HDAC11 up to 10,000 nM. IC₅₀ values of T-3793168 were 86 nM for HDAC6 and over 2,000 nM against other HDACs.

Target	Compounds IC ₅₀ (nM) ^a		
	T-3796106	T-3793168	Tubastatin A
HDAC1	6200 (5820-6660)	b	>10000
HDAC2	>10000	b	>10000
HDAC3	4000 (3480-4470)	b	>10000
HDAC8	>1000	5600 (4170-7080)	>1000
HDAC4	6200 (5970-6380)	>10000	6200 (6030-6330)
HDAC5	1700 (1610-1800)	2300 (1550-3060)	1900 (1580-2310)
HDAC7	1100 (1050-1130)	5800 (4000-7660)	590 (530-641)
HDAC9	2700 (2540-2840)	5000 (4950-4960)	1100 (913-1380)
HDAC6	12 (12.3-12.4)	86 (67.4-104)	15 (14.3-15.2)
HDAC10	>10000	b	>10000
HDAC11	>10000	b	>10000

Table 1 HDAC panel assay. The selectivity of T-3796106 and T-3793168 was analyzed based on HDAC enzyme inhibition. [a] The compound activity against 11 HDACs represented with the IC₅₀ value. The IC₅₀ values shown are the mean values of duplicate measurements; the numbers in parentheses represent each data. [b] No inhibition or compound activity that could not be fit to an IC₅₀ curve.

T-3796106 and T-3793168 do not cause neuronal toxicity even at high concentrations

First, we tested whether T-3796106 or T-3793168 induces any cytotoxicity in murine neuronal explant cultures, using concentrations substantially higher than those we subsequently used

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3 for axonal transport studies to indicate a large therapeutic window. We used superior cervical
4 ganglion (SCG) explants because this neuron type is well-suited for genetic manipulation by
5 microinjection and for axonal transport studies^{51, 52}, and incubated with concentrations from 1
6 μM to 100 μM for 24 h. No toxicity was observed at any concentration. Neurites remained
7 morphologically similar to vehicle-treated or untreated cultures, even in their distal terminals
8 which are typically the most vulnerable site (Fig 1A, B). Thus, both compounds are safe for
9 neurons up to 100 μM for at least 24 h.

18 **Increased acetylation levels of α -tubulin after T-3796106 and T-3793168 treatment in** 19 **neurons**

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23 We next confirmed that steady state α -tubulin acetylation increases with either T-3796106 or
24 T-3793168 within the above concentration range (data not shown), before titrating down to
25 determine the dose-response curve for α -tubulin acetylation at sub-saturating levels, thereby
26 minimizing the risk of off-target effects. Both compounds showed a clear dose-response effect
27 between 1 nM and 250 nM in a 24 h incubation (Fig 2A), reaching significance at 50 nM for T-
28 3796106 and 250 nM for T-3793168 (Fig 2B). Based on this characterization we used
29 concentrations of 100 nM and 250 nM respectively in our subsequent axonal transport
30 experiments. At these concentrations, there was no effect on histone acetylation which
31 indicates a high selectivity of these compounds towards HDAC6 (Supplementary Figure 1).

43 **Axonal transport of mitochondria in wild type SCG cultures is not altered by T-3796106** 44 **or T-3793168**

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48 In the absence of a pathogenic mutation, we found no significant change in either the number
49 or the average and maximum velocity of axonally transported mitochondria in dissociated wild-
50 type SCG neurons treated with T-3796106, T-3793168 or Tubastatin A (Fig 3B, C, D, E). Thus,
51 there is no change in basal axonal transport parameters for this cargo.

57 **Mitochondrial transport impairment induced by S135F mutation is rescued by T-** 58 **3793168**

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3 In the presence of the HSPB1^{S135F} mutation, which causes CMT2 and distal HMN in patients
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5 ¹⁵, both the numbers of anterogradely and retrogradely moving mitochondria were significantly
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7 decreased relative to wild type neurons 12 h after microinjection (Fig 4A, B), mirroring similar
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9 changes reported in sensory neurons ¹¹. The transport deficits in both directions were
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11 significantly rescued in neurons treated for 24 h with 250 nM T-3793168, while those treated
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13 with 100 nM compound T-3796106 showed a trend towards increased mitochondrial transport
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15 but the difference was not statistically significant (Fig 4A, B). As previously reported ¹¹, we
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17 also found a rescue of anterograde axonal transport with 1 μ M Tubastatin A, but in the
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19 retrograde direction the trend towards a rescue with Tubastatin A was not significant. The
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21 average and maximum speed of mitochondria movement was not significantly altered in the
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23 neurons with the HSPB1^{S135F} mutation and was unaffected by any of these treatments (Fig 4C,
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25 D).

26 27 28 **P182L mutation does not alter mitochondrial transport in SCG neurons**

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30 Consistent with previous findings in sensory neurons ¹¹, SCG neurons expressing
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32 HSPB1^{P182L} showed no significant changes in mitochondrial transport compared to wild type
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34 neurons (Fig 5). Treatment of these mutant-expressing neurons with T-3796106, T-3793168
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36 or Tubastatin A also had no effect on mitochondrial movement (Fig 5).

37 38 39 **T-3796106 and T-3793168 increase α -tubulin acetylation in human whole blood**

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41 We finally investigated the effects of T-3796106 and T-3793168 on acetylation of α -tubulin in
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43 human cells, using whole blood. For both compounds, clear dose-response effects were
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45 observed between 10 nM and 30 μ M in a 4 h incubation (Fig 6). Over 100 μ M of our
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47 compounds and 300 μ M of hydroxamic acid-based HDAC6 inhibitor ACY-1215 showed
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49 some precipitate when the compounds were added in the culture medium. We also observed
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51 the effect of Tubastatin A on acetylated α -tubulin in the human whole blood assay. It showed
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53 a similar trend. In brief, the levels of acetylated α -tubulin were almost the same at 10 and 30
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55 μ M of Tubastatin A (data not shown) as well as that of ACY-1215.
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DISCUSSION

We report the development and characterization of two novel non-hydroxamic acid-based inhibitors with high potency and specificity for HDAC6, low toxicity in murine primary neuronal cultures and a dose-dependent effect on neuronal α -tubulin acetylation between 1-250 nM. T-3793168 significantly increases both the anterograde and retrograde flux of mitochondria in axonal transport within 24 h of application to neurons expressing the CMT-2F HSPB1 mutation S135F, and T-3796106 shows a similar, albeit non-significant trend. Neither alters axonal transport in wild-type cells.

For both compounds the changes in acetylated tubulin in whole blood was several orders of magnitude greater than those in mouse primary neuronal cultures. This suggests a tissue-, tubulin isoform-, or species-specific effect on the efficacy of these HDAC6 inhibitors indicating significant scope of lead compound optimization. Further studies to understand the basis of this specificity should help to optimize their efficacies to achieve substantial enhancement of axonal transport in human axonal disorders.

A major advantage over hydroxamic acid-based inhibitors is greater selectivity over other HDAC family members and this is the case for these compounds. For example, hydroxamic acid-based HDAC6 inhibitor ACY-1215 has a high HDAC6 enzyme potency with IC_{50} value of 4.7 nM but much lower selectivity (12-fold selectivity for HDAC6 and HDAC1 at IC_{50})⁵³. In contrast, non-hydroxamic acid-based HDAC6 inhibitors T-3796106 and T-3793168 showed excellent selectivity (>25-fold over other HDAC family members; >100-fold selectivity over HDAC1 at IC_{50}).

It will be important now to test the effect of these compounds on axonal transport in other disease models where transport is impaired and the effect on other axonal transport cargoes. For example, axonal transport defects underlie vincristine neuropathy and some forms of hereditary spastic paraplegia and ALS, and have also been implicated in glaucoma,

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3 Alzheimer's disease and multiple sclerosis. Many axonal transport cargoes need to be
4 continuously shuttled back and forth but among some of the most important ones are
5 NMNAT2, whose absence limits the survival of transected axons⁵¹ and the retrograde
6 transport of lysosomes to maintain efficient autophagy and mitochondrial quality control⁵⁴,
7 and neurotrophins.
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14 Finally, it will be important to test the efficacy of HDAC6 inhibition and rescue of axonal
15 transport *in vivo* using methods for live imaging of transport cargoes in live nerves and CNS
16 tissue^{33, 55, 56}, to assess how HDAC6 inhibition compares to other methods of boosting
17 axonal transport in experimental models^{57, 58} and to further develop these lead compounds.
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26 **METHODS**

27 **Chemicals**

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31 T-3796106 and T-3793168 are novel HDAC6 inhibitors developed by Takeda Pharmaceutical
32 Company Limited (Patent WO2017014321)⁵⁹. The purity of T-3796106 and T-3793168 was
33 determined to be $\geq 95\%$ by elemental analysis which was performed by Sumika Chemical
34 Analysis Service, Ltd. experimentally determined hydrogen, carbon, and nitrogen composition
35 by elemental analysis was within $\pm 0.4\%$ of the expected value, implying a purity of $\geq 95\%$.
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46 **Enzyme assay**

49 HDAC panel assay was performed by Reaction Biology Corp. (Malvern, PA, USA) according
50 to their validated protocol. To evaluate the potency and selectivity of T-3796106 and T-
51 3793168, HDAC panel assay was carried out by Reaction Biology Corp. Briefly, the
52 deacetylation reaction was performed in buffer conditions of 50 mM Tris-HCl pH 8.0, 137 mM
53 NaCl, 2.7 mM KCl, 1 mM MgCl₂, and 1 mg/mL bovine serum albumin (BSA), and 1% DMSO.
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3 The fluorogenic peptide, RHK-K(Ac)-AMC, is used as substrate for Class1 and 2B HDACs,
4 RHK(Ac)K(Ac)-AMC for HDAC8, and Boc-Lys(trifluoroacetyl)-AMC for Class2A HDACs. After
5 the reaction, by Developer with Trichostatin A, a fluorescence signal (Ex. 360 nm/Em. 460
6 nm) developed.
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11 **Animals**

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14 C57BL/6J0laHsd mice were obtained from Harlan UK (Bicester, UK). All animal work was
15 carried out in accordance with the Animals (Scientific Procedures) Act, 1986, under Project
16 License 70/7620.
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21 **Cell culture**

22 *Explant SCG cultures*

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25 SCGs were dissected from 0 to 2 days old C57BL/6 (wild-type) mouse pups. Cleaned explants
26 were placed in the centre of 3.5 cm tissue culture dishes pre-coated with poly-L-lysine (20
27 mg/mL for 1–2 h; Sigma) and laminin (20 mg/mL for 1–2 h; Sigma). Explants were cultured in
28 Dulbecco's Modified Eagle's Medium (DMEM) with 4,500 mg/L glucose and 110 mg/L sodium
29 pyruvate (Sigma), 2 mM glutamine (Invitrogen), 1% penicillin/streptomycin (Invitrogen), 100
30 ng/mL 7S NGF (Invitrogen), and 10% fetal bovine serum (Sigma). Four μ M aphidicolin
31 (Calbiochem) was used to reduce proliferation and viability of small numbers of non-neuronal
32 cells. Cultures were used after 6 days.
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45 *Dissociated SCG cultures*

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48 SCG ganglia were dissociated by incubation in 0.025% trypsin (Sigma) in PBS (without CaCl_2
49 and MgCl_2) for 30 min followed by 0.2% collagenase type II (Gibco) in PBS for a further 20
50 min. Ganglia were then gently triturated using a pipette. After a 2 h pre-plating stage to remove
51 non-neuronal cells, 5,000–10,000 dissociated neurons were plated in a 1 cm^2 poly-L-lysine
52 and laminin-coated area in the centre of 3.5 cm ibidi μ -dishes (Thistle Scientific, Glasgow, UK)
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3 for microinjection experiments or in the centre of 3.5 cm tissue culture dishes for analysis by
4 western blotting. Dissociated cultures were maintained the same as explant cultures.
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7 **HDAC6 inhibitors treatment**

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10 The SCG explants and dissociated cultures were treated for 24 h at 37°C with compound
11 dosages ranging from 1 μ M to 100 μ M for toxicity experiments and 1nM to 250 nM for testing
12 the dose-response study of α -tubulin acetylation. For axonal transport rescue experiments,
13 dissociated SCG cultures were treated with either 100 nM T-3796106, 250 nM T-3793168, 1
14 μ M Tubastatin A or an equivalent amount of DMSO.
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22 **Plasmid constructs**

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24 The S135F and P182L mutations were introduced separately by QuikChange II site-directed
25 mutagenesis (Stratagene) into the complete open reading frame of human HSPB1 isoform
26 cloned into expression vector pCMV-Tag2 (Stratagene). The mito-EGFP construct was kindly
27 provided by Dr Andrea Loreto.
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34 **Microinjection**

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36 Microinjection was performed using a Zeiss Axiovert 200 microscope with an Eppendorf 5171
37 transjector and 5246 micromanipulator system and Eppendorf Femtotips. Microinjection mixes
38 of plasmid DNA were prepared in 0.5 \times PBS(-), passed through a Spin-X filter (Costar,
39 Glasgow, UK) Eppendorf and injected directly into the nuclei of SCG neurons in dissociated
40 cultures. Femtotips were loaded with the microinjection mix and injection was performed using
41 an Eppendorf 5171 transjector and 5246 micromanipulator system on a Zeiss Axiovert 200
42 microscope. All injections were carried out directly into the nuclei of dissociated SCG neurons.
43 A maximum total DNA concentration of 0.05 μ g/ μ L in the injection mix was used. Forty cells
44 were injected per dish and imaging was performed 12 hours after microinjection.
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56 **Western Blotting**

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3 Following treatment, ganglia and neurites were collected and washed in PBS with complete,
4 ethylenediaminetetraacetic acid (EDTA)-free protease inhibitor cocktail tablets (Sigma-
5 Aldrich), and lysed directly into 2x Laemmli sample buffer. A total of 10 μ L of each sample
6 were separated on a 12% SDS-PAGE and transferred to PVDF membrane (Millipore) using
7 the Bio-Rad Mini-PROTEAN III wet transfer system. Blots were blocked and incubated with
8 primary antibodies overnight (in 1xTBS pH. 8.3, with 0.05% Tween 20 and 5% milk powder or
9 5% BSA). The antibodies were directed against α -tubulin (1/5,000; ab 15246, Abcam) and
10 acetylated α -tubulin (1/5,000; T7451, Sigma) and detected with mouse-700 (Life
11 Technologies) and rabbit-800 (LI-COR) secondary antibodies. Blots were then scanned and
12 quantified using the Odyssey imaging system (LI-COR Biosciences, Lincoln, North Carolina).
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25 **Live imaging of mitochondrial transport and image analysis**

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27 Mitochondria were labelled by microinjection of mito-EGFP and their movement along the
28 neurites was recorded with an inverted spinning-disk confocal microscope Olympus IX70
29 using a 100x 1.49 NA oil immersion objective (Olympus), and controlled with MetaMorph 7.7
30 software (Molecular Devices). The environment was controlled with a stage top incubator
31 (model INUBG2E-ZILCS; Tokai Hit), set at 37°C and CO₂ set to 5%. Time lapse images of
32 mitochondrial movements were acquired every 1 s for 2 min (120 frames in total). A total of
33 4-5 movies from different neurons were captured from each culture dish. Individual neurites
34 were straightened using the Straighten plugin in ImageJ software version 1.44 (Rasband,
35 W.S., ImageJ, U. S. National Institutes of Health, Bethesda, MD; <http://imagej.nih.gov/ij/>,
36 1997e2012). Transport parameters were determined for individual neurites using the
37 Difference Tracker set of ImageJ plugins ⁰⁶⁰. The principal output of these plugins is the
38 number of moving particles identified in each frame of the image, normalized to 1000 pixels
39 (Figure 3A).
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56 **Human whole blood assay**

57 *Study design*

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3 In human whole blood assay, whole human blood was collected from healthy volunteers after
4 informed consent at Takeda Pharmaceutical Company Limited. 25 μ L of the collected human
5 whole blood was put into each well of a 96-well round-bottom plate. The whole blood was
6 treated with 10 μ L of diluted compounds in 10% FBS containing RPMI 1640 medium (Gibco).
7
8 In the control group, 0.1% DMSO was added as a final concentration. After that, the treated
9 whole blood was incubated at 37°C for 30 minutes at 5% CO₂. Next, 65 μ L of RPMI 1640
10 medium was added onto each well, and the samples were incubated at 37°C for 3.5 hours. T-
11 3796106, T-3793168, and ACY-1215 were dissolved in 100% DMSO (to a stock concentration
12 of 10-300 mM for our in vitro studies).
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23 *Measurements*

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25 For flow cytometry analyses, the compound-treated whole blood samples were transferred to
26 an assay block (Costar). Diluted Lyse/Fix buffer (BD Biosciences) in dH₂O was added to each
27 sample with pipetting well. The samples were put at RT for 10 minutes and then centrifuged
28 at 400xg for 5 minutes. After centrifugation, the supernatant was removed by aspiration. 250
29 μ L of Perm/Wash buffer I (BD Biosciences) was added to each well and the samples were
30 transferred into a 96-well V-bottom plate, and they were incubated on ice for 20 minutes.
31 These samples were centrifuged at 400xg for 5 minutes at RT, and the supernatant was
32 removed by aspiration. The cells were stained with Zenon conjugated AF647 acetylated α -
33 tubulin (ab179484, Abcam) or matched isotype control (ab172730, Abcam) for 20-30 minutes
34 on ice. Zenon Rabbit IgG Labeling Kit, AF647 (Molecular Probes) was used according to the
35 provided protocol. The cells were centrifuged at 400xg at 5 minutes, and the supernatant was
36 removed, and then washed with 200 μ L of Perm/Wash buffer I. After re-centrifugation and
37 removal of the supernatant, the samples were suspended with 200 μ L of FACS stain buffer
38 (1% FBS/PBS). The samples were analyzed using lymphocyte gate by BD Fortessa, and the
39 results were analyzed with FlowJo software. Therefore, only lymphocytes were analyzed
40 neither erythrocytes nor thrombocytes were included in the assay.
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Statistical analysis

Statistical tests, as described in the figure legends, were performed using Prism software (GraphPad Software Inc, La Jolla, CA, USA). A p value of >0.05 was considered not significant (ns) and *p < 0.05 was significant.

SUPPORTING INFORMATION

Supplementary method and extended western blot results.

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Writing the manuscript: R.A., M.P.C. and M.T.M.

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Notes

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14
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16
17 development of human whole blood assay..
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20 **ABBREVIATIONS**

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23 HDAC, histone deacetylase; KIF5A, kinesin heavy chain isoform 5A; A β ₁₋₄₂, amyloid beta
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25 peptide 42; HMN, hereditary motor neuropathy; NMNAT2, nicotinamide mononucleotide
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27 adenylyltransferase 2; DMSO, dimethyl sulfoxide; FBS, fetal bovine serum.
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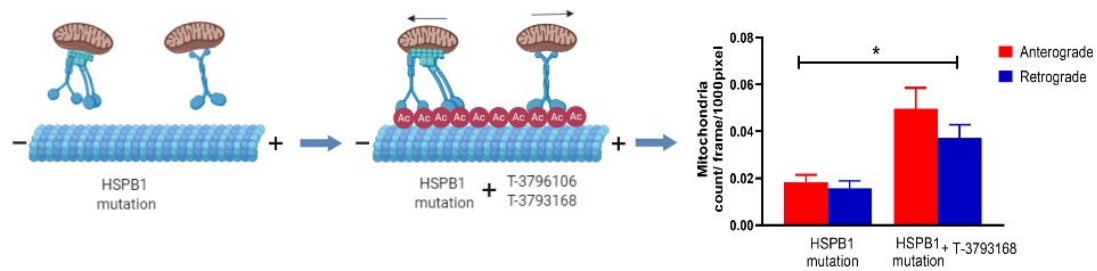
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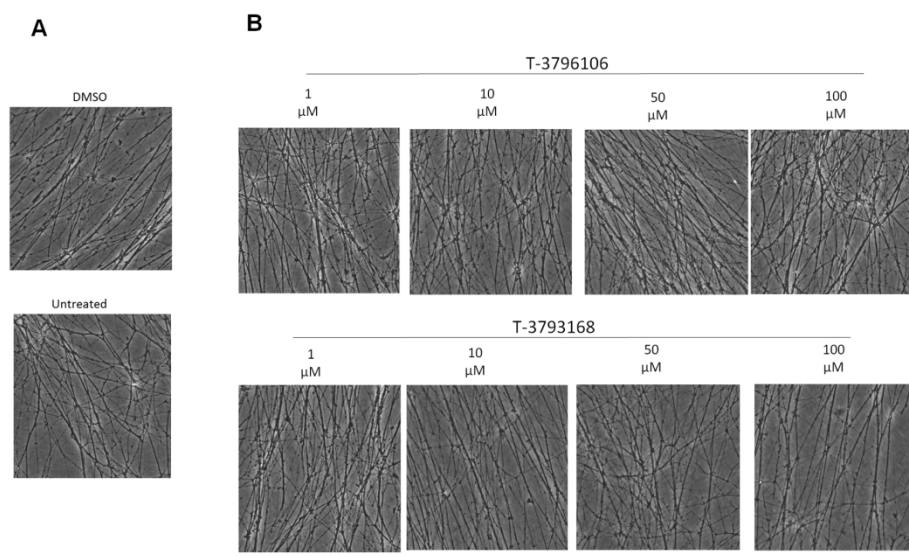


Figure 1

Figure 1. Morphologically normal neurites of SCG explants treated with high concentrations of T-3796106 and T-3793168. (A) Vehicle treated and untreated cultures have healthy looking neurites with no signs of fragmentation or blebbing. (B) Cultures treated with indicated compound concentrations look morphologically similar with control cultures, which suggest no cytotoxic effect. Scale bar 50 μm .

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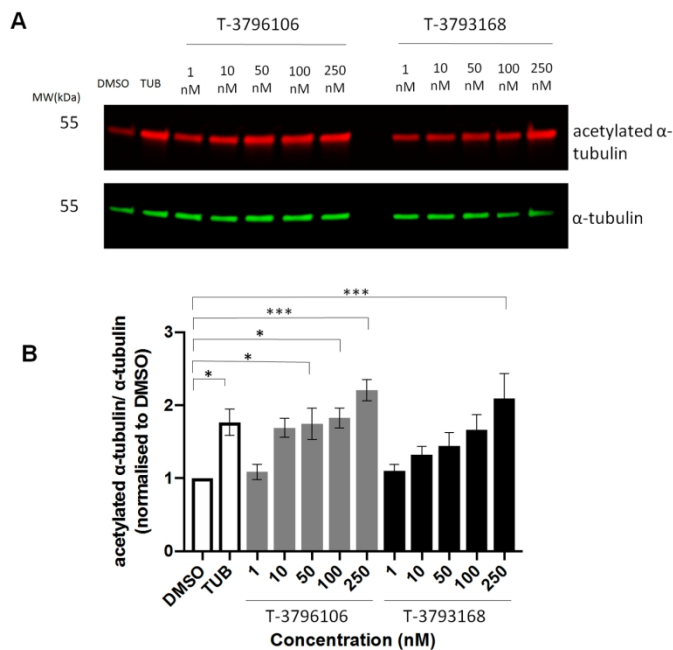


Figure 2

Figure 2. Pharmacological properties of T-3796106 and T-3793168 in SCG neurons. Western blot showing dose response of T-3796106 and T-3793168 (A) on acetylation of α -tubulin. 1 μ M Tubastatin A (Tub) was used as positive control. Quantification of the ratio of acetylated α -tubulin to total tubulin levels on Western blot for T-3796106 and T-3793168 (B). Statistically significant difference between groups is indicated (* $p < 0.05$, 1-way analysis of variance). Data are presented as means \pm s.e.m. $n=4-5$.

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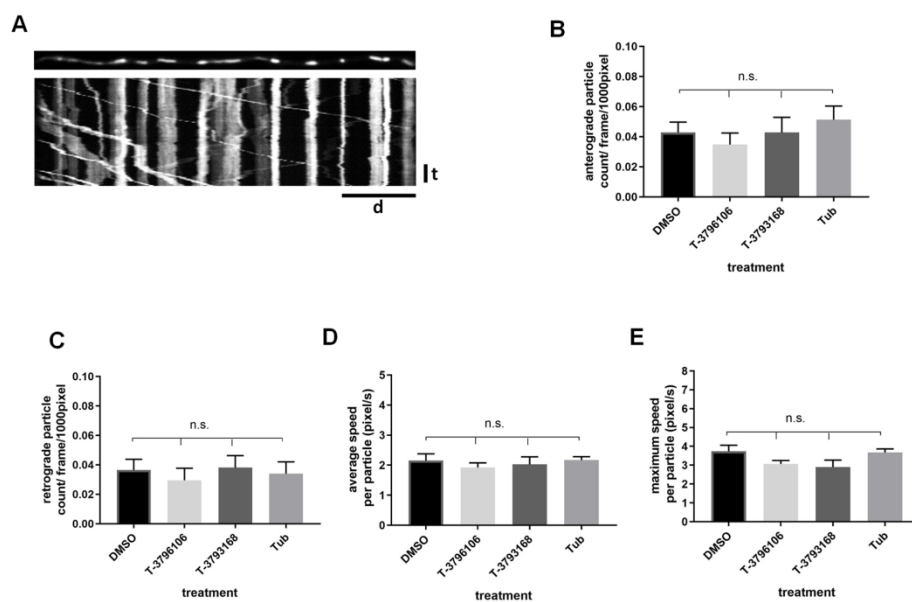


Figure 3

Figure 3. Mitochondrial transport remains unaltered in wild type SCGs after treatment. Representative kymograph obtained from a SCG neuron grown in dissociated cultures (A). The top image shows a straightened neurite. Vertical lines indicate stationary mitochondria while lines deflecting to the right or left represent anterograde or retrograde moving mitochondria. Time (t) scale bar: 120 s; distance (d) scale bar: 10 μ m. Quantification of anterograde (B) and retrograde (C) mitochondria transport, average (D) and maximum speed (E) of mitochondria movement in neurons treated with the indicated compounds and concentrations. Data are presented as means \pm s.e.m. n=12-14.

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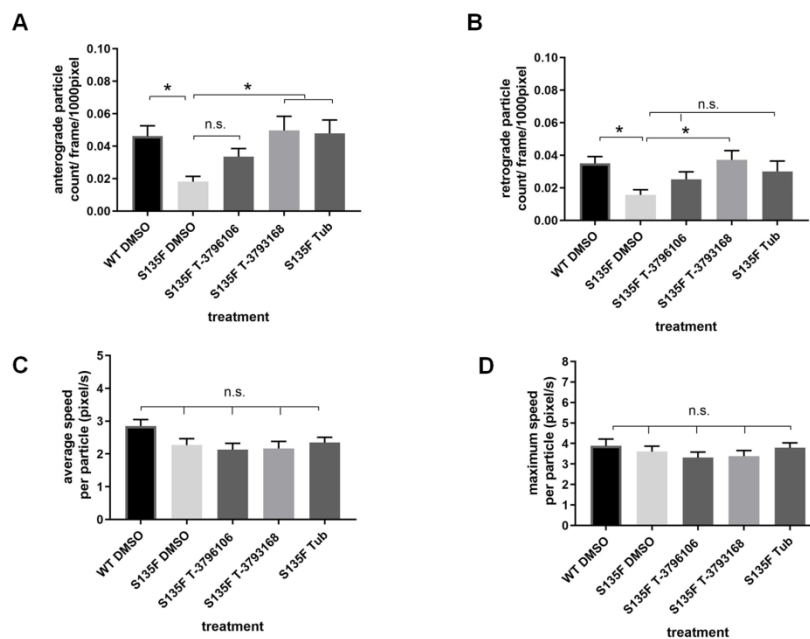


Figure 4

Figure 4. T-3793168 reverses axonal transport deficit in SCGs with HSPB1S135F mutation. Quantification of anterograde (A) and retrograde (B) mitochondria transport, average (C) and maximum speed (D) of movement with genotypes and treatments indicated. Statistically significant differences between groups is indicated (* $p < 0.05$, 1-way analysis of variance). Data are presented as means \pm s.e.m. $n=12-17$.

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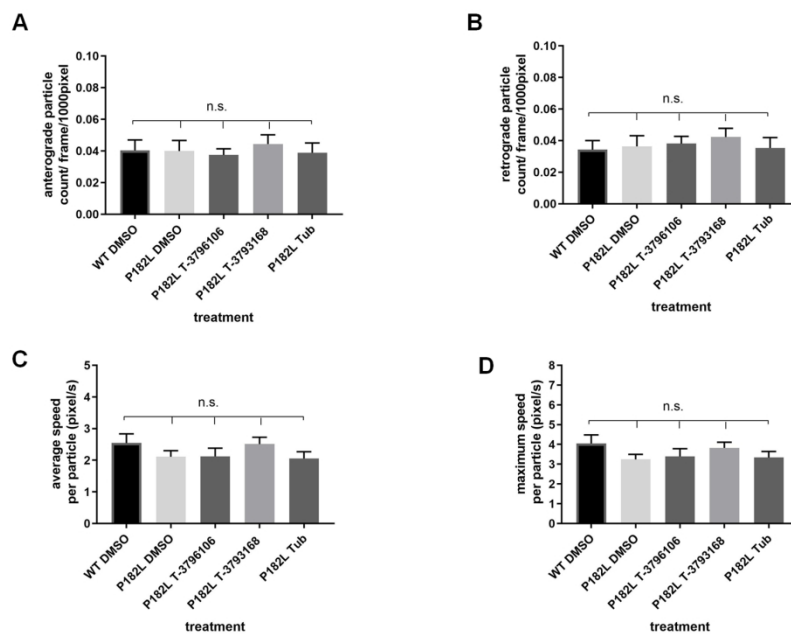


Figure 5

Figure 5. No changes in axonal transport of SCGs with HSPB1P182L mutation. Quantification of anterograde (A), retrograde (B), average (C) and maximum speed (D) of mitochondria transport with genotypes and treatments indicated. Data are presented as means \pm s.e.m. n=12-14.

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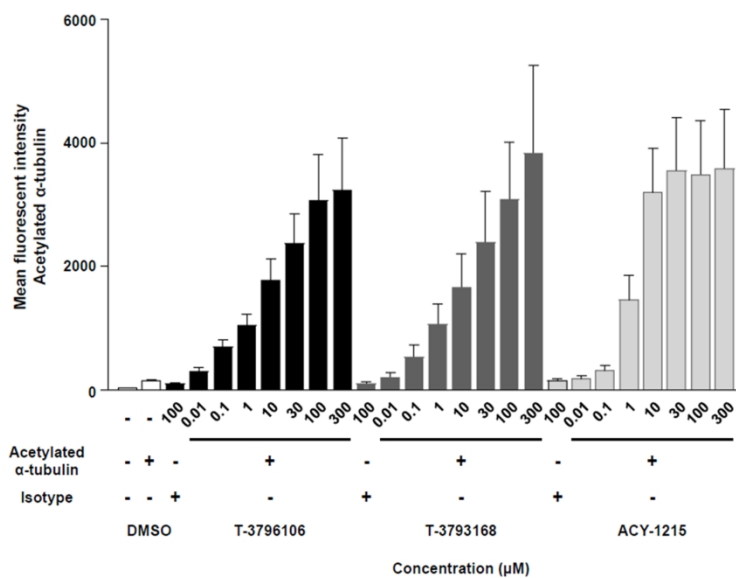


Figure 6

Figure 6. Increased acetylation levels of α -tubulin in human whole blood. Quantification of acetylated α -tubulin in whole blood treated with the indicated compounds and concentrations. Data are presented as means \pm s.e.m. $n=3$.

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