Discovery of Tirasemtiv, the First Direct Fast Skeletal Muscle Troponin Activator

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Supporting Information

ABSTRACT: The identification and optimization of the first activators of fast skeletal muscle are reported. Compound 1 was identified from high-throughput screening (HTS) and subsequently found to improve muscle function via interaction with the troponin complex. Optimization of 1 for potency, metabolic stability, and physical properties led to the discovery of tirasemtiv (25), which has been extensively characterized in clinical trials for the treatment of amyotrophic lateral sclerosis.

KEYWORDS: Amyotrophic lateral sclerosis (ALS), skeletal muscle activator, troponin, tirasemtiv, imidazo[4,5-b]pyrazin-2-one

Direct activation of skeletal muscle has the potential to improve physical performance in patients with compromised muscle function, such as those suffering from amyotrophic lateral sclerosis (ALS), spinal muscular atrophy, and Charcot-Marie-Tooth disease. Muscle impairment in these disease states occurs from diminished neuromuscular input and can lead to disability and increased mortality. Current treatment options are limited or nonexistent and represent a significant unmet medical need.

Precedent for activation of muscle function exists with omecamtiv mecarbil, a direct cardiac muscle activator that improves contractility and is undergoing clinical trials for the treatment of amyotrophic lateral sclerosis.

This report describes the identification and optimization of the first direct activators of fast skeletal muscle, culminating in the discovery of tirasemtiv (25), a compound that was found to selectively sensitize the sarcomere of fast skeletal muscle to calcium and increase its force production via interaction with the regulatory troponin complex.

Skeletal muscle is classified into two main forms, fast skeletal muscle (Type II fibers) and slow skeletal muscle (Type I), which differ in usage, metabolism, and sarcomere composition. Fast skeletal muscle fibers have unique myosin and troponin components compared to cardiac and slow skeletal muscle, while slow skeletal muscle fibers have the same myosin and troponin C (TnC, a subunit of the regulatory complex) as cardiac muscle. Myosin is the molecular motor responsible for contractility where chemical energy from adenosine triphosphate (ATP) hydrolysis is coupled to force generation. The troponin complex plays a regulatory role by acting as a calcium sensor and resides in the thin filament along with actin and tropomyosin. The sequence homology between fast and slow muscle myosin and troponins C, T, and I is about 50%, providing a good opportunity to find selective skeletal muscle troponin activators.

Because skeletal muscle is mainly responsible for voluntary movement and has unique myosin and troponin components compared to other muscle types, fast skeletal muscle represents a compelling target for small molecule intervention.

Compounds with the potential to modulate human muscle function were identified using a high-throughput screen (HTS) that measures the rate of ATP hydrolysis in rabbit fast skeletal muscle myofibrils as a surrogate for force generation in skeletal muscle.

Biochemical activity (AC_{40}) is defined as the compound concentration that produces a 40% increase in myofibril ATPase activity at 25% of the calcium concentration needed for a maximal response in control preparations. Among the hits was imidazopyrazinone 1 (CK-1303249). Compound 1 displayed modest biochemical potency (AC_{40} = 7.0 μM), ligand and lipophilic efficiencies (LE and LipE) of 0.32 and 5.01, respectively, and specificity for fast skeletal muscle over slow skeletal, cardiac, and smooth muscle (Table 1). To determine whether activation occurs via the myosin or regulatory complex, mixed sarcomeres were created using myosin and thin filament preparations from fast, slow, and cardiac muscle sources, and each was then assayed in the presence of 1. Because activation was observed only in combinations containing fast skeletal thin filaments, 1 was determined to be a direct troponin complex activator.

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Table 1. Activity and Selectivity of HTS Hit and Close-in Analogues

| # | R   | FSK AC₄₀ (µM) | CV AC₄₀ (µM) | SSK AC₄₀ (µM) |
|---|-----|---------------|---------------|---------------|
| 1 | S-Me | 7             | >49           | >39           |
| 2 | R-Me | >49           |               |              |
| 3 | H    | >49           |               |              |

**FSK** = fast skeletal muscle; **CV** = cardiac muscle; **SSK** = slow skeletal muscle.

The observed increase in ATP hydrolysis in the presence of 1 was subsequently shown using the skinned fiber assay to translate to improved force generation in a skinned, fast muscle fiber (Figure 1). This assay is performed by mounting a muscle fiber (pretreated with detergent to remove the membranes and cytoplasm) to a force transducer rig to enable force measurement across the muscle. Treatment of primarily fast rabbit psoas muscle fibers with 1 increased the calcium sensitivity of the muscle fibers without increasing the maximum force or the shape of the curve (Figure 1A). A related control experiment using skinned, primarily slow, rabbit semimembranosus muscle fibers failed to show increased calcium sensitivity in the presence of 1 (Figure 1B). For potency comparison across compounds, DF₃₀ is defined as the compound concentration that produces a 30% increase in tension at 25% of the calcium concentration needed for a maximal response in control preparations.

The initial optimization goals were based loosely on our previous experience with the optimization of cardiac muscle activators. The biochemical potency needed for compounds to show physiologically significant cardiac muscle activation was between 0.1—5 µM. The desired pharmacokinetic profile would ensure that efficacious concentrations could be sustained by dosing orally once or twice per day. Therefore, the target compound profile included biochemical potency (AC₄₀) less than 1 µM, minimal in vitro clearance, and good in vivo exposure with oral dosing.

Figure 1. Calcium-force relationship of 1 on skinned skeletal fibers from (A) rabbit psoas muscle (fast) and (B) rabbit semimembranosus muscle (slow) plotted as % of maximal tension vs (−)log of calcium concentration (pCa).

Synthesis of the imidazo[4,5-b]pyrazin-2-one analogues of 1 was achieved using Scheme 1. Addition of primary amines to 2-

Scheme 1. Synthesis of Imidazopyrazinone Derivatives

(a) NHR¹. (b) CDI, THF. (c) (R² = furyl or phenyl) R²B(OH)₃, Cl₂Pd(dppf), Na₂CO₃, dioxane/water. (d) (R² = SMe) NaSMe, NMP. (e) (R² = OMe) NaOMe, NMP. (f) (R² = CN) Zn(CN)₂, (Ph₃P)₄Pd, DMF. (g) (R² = Ac) tributyl(1-ethoxyvinyl)tin, Cl₂Pd(dppf), dioxane. (h) (R² = H) H₂, Pd/C, EtOH. (i) (R² = Me) trimethylboroxine, Cl₂Pd(dppf), Na₂CO₃, dioxane/water. (j) (R² = Ac) tributyl(1-ethoxyvinyl)tin, Cl₂Pd(dppf), dioxane. (k) (R² = H) H₂, Pd/C, EtOH. (l) (R² = vinyl) trimethylboroxine, Cl₂Pd(dppf), Na₂CO₃, dioxane/water. (m) (R² = vinyl) trimethylboroxine, Cl₂Pd(dppf), Na₂CO₃, dioxane/water. (n) (R² = Et) from R² = vinyl, H₂, Pd/C, EtOH.

The SAR around the N1-position (R¹) was examined in the context of a bromine substituent at the 6-position (Table 3). The focus was on defining a minimum pharmacophore at this position.

amino-3,5-halopyrazine (31) gave diaminopyrazines 32 in good yields, followed by ring closure using carbonyl diimidazole (CDI) to afford imidazo[4,5-b]pyrazin-2-ones 33. The targeted analogues were produced in one or two steps from 34. Having established calcium sensitization in isolated muscle fibers, we were interested in demonstrating improved functional activity in an animal model. We initially focused on replacing the furan in 1 to improve the metabolic stability and physical properties while exploring the structure–activity relationship (SAR) at that position. Overall, the tolerance for substitution at R² was found to be somewhat limited (Table 2). The observed potency improvement with vinyl (5) but not phenyl (4) illustrated the limited space in that region and hinted at the potential for more potent, lower molecular weight structures. The combination of reduced potency with ethyl (4) and methoxy (5) but improved potency with SMe 8 and bromo 9, suggesting a preference for polarizable electron-rich groups. In contrast, analogues with greater polarity (10—12) were much less active, pointing to intolerance for certain polar interactions at this position.

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position and reducing oxidative potential for the series (as measured from microsome clearance experiments). Replacement of the phenyl group from the methylbenzyl substituent of 9 with an ethyl group (10) improved stability in rat and human microsomes. A 3-pentyl group provided similar microsomal stability to R1 = 10, standard deviation = 0.2).

The SAR exploration at the R2 position in the context of a 3-pentyl R1 group revealed specific steric and electronic preferences (Table 4). Compact, electron-rich lipophilic groups were generally preferred in this position, as exemplified by the potency loss observed when bromine (14) or chloro (21) substituent was replaced with a trifluoromethyl (22) or cyano (27) group. Vinyl analogue 24 was among the more potent compounds, but the strongest preference was for an ethynyl group (25, CK-2017357), which was 3-fold more potent than bromo 14. Notably, propynyl analogue 26 was much less active, again indicating a steric limitation in that part of the binding pocket. The microsomal stability of 14, 21, and 25 was roughly equivalent, with 25 displaying the most promising combination of potency and metabolic stability.

The importance of the hydrogen bond donor/acceptor region of 25 to biochemical potency is highlighted in Figure 2. Compound 25 may exist as a mixture of tautomers in solution (25a, 25b). Removal of the H-bond donor by either isosteric replacement of nitrogen with oxygen (29) or methylation at nitrogen (28) or oxygen (30) led to a dramatic reduction in potency.

The rat pharmacokinetic profiles for compounds with optimal potency and microsomal stability were assessed to determine their suitability for in situ live fiber experiments (Table 5). In rats, analogues 14, 21, and 25 all showed low to moderate clearance values and good oral bioavailability when dosed as crystalline suspensions. The free fraction (F) for 25 in both human and rat plasma was greater than those for 14 and 21. Based on superior activity and oral free exposure, 25 was selected for advanced in vivo testing.

Table 3. Assessment of N-1 position (R1)

| #  | R1      | AC50 (μM) | DF50 (μM) | Re (r) | Re (h)  |
|----|---------|-----------|-----------|--------|---------|
| 9  | (S)-Me-benzyl | 1.6       | 3.0       | 0.5    | <0.3    |
| 13 | (S)-2-butyl   | 1.0       | 5.1       | 0.3    | <0.2    |
| 14 | 3-pentyl      | 0.3       | 0.4       | 0.4    | <0.2    |
| 15 | i-Pr         | 6.0       | 2.2       | NT     | NT      |
| 16 | t-butyl       | 0.8       | 4.6       | 0.3    | 0.2     |
| 17 | cyclopropyl   | >49       | NT        | NT     | NT      |
| 18 | cyclohexyl    | 10        | NT        | NT     | NT      |

*Re = ratio of extraction in microsomes (r = rat, h = human). Compound 14 was tested multiple times in the skinned fiber assay (n = 10, standard deviation = 0.2).

Figure 2. SAR assessment of the donor/acceptor region of 25.
The sensitization of fast skeletal muscle to calcium in the presence of 25 was demonstrated using an in situ preparation of the extensor digitorum longus (EDL) muscle in a rat model where the nerve and the blood supply are intact. Systemic infusions of 25 resulted in rapid, dose-dependent increases in muscle force at submaximal stimulation rates (Figure 3) without increasing the maximum tension at tetanic stimulation rates. This result was consistent with the effects of 25 on the force—calcium relationship of skinned rabbit psoas muscle fibers and the increase in ATPase rate (a surrogate for force production) in the biochemical assay.

To support pharmacology and safety studies as well as human PK and dose projections, a pharmacokinetic assessment of 25 was performed (Table 6). In a Caco-2 cell monolayer system, 25 demonstrated high permeability (permeability coefficient of >50 × 10⁻⁶ cm/s) and no apparent efflux. It showed low to moderate liver microsomal stability and relatively low exposure following oral administration of the clinical dose in humans.20

Table 6. In Vitro and in Vivo Pharmacokinetics of 25

| species | Re ² | CL (mL/min/kg) | Vss (L/kg) | F (%) |
|---------|------|----------------|------------|------|
| human   | <0.21| 4.3            | 1.7        | 93   |
| dog     | <0.21| 2.6            | 0.8        | 98   |
| rat     | 0.26 | 13.9           | 1.0        | 43   |

²Re = ratio of extraction in microsomes; PK dose = 1 mg/kg.

Compound 25 exhibited moderate inhibition and time-dependent inhibition of CYP1A2 (IC₅₀ = 20 μM, IC₅₀ shift of >3 following a 30 min preincubation).21 In clinical studies involving ALS patients taking both 25 and Riluzole, a drug–drug interaction (DDI) occurred that resulted in an approximate doubling of Riluzole concentrations.22 Riluzole is a commonly prescribed drug for ALS that is cleared mainly by CYP1A2 and CYP1A1 mediated metabolism.23 This DDI was managed in the clinic by reducing the dose of Riluzole from 100 mg/day b.i.d. to 50 mg/day q.d.

The effect of compound 25 (tirasemtiv) on muscle performance has been evaluated in preclinical and clinical models. In a mouse model of ALS with functional deficits (B6SJL-SOD1<sup>1093A</sup>), a single dose of tirasemtiv significantly increased submaximal isometric force, forelimb grip strength, grid hang time, and rotarod performance.24 Tirasemtiv also increased grip strength in rats treated with the anti-nAchR antibody in the PT-EAMG (passive transfer experimental autoimmune myasthenia gravis) rat model.16 In a clinical study performed in healthy humans, tirasemtiv increased force produced by the tibialis anterior in a dose-, concentration-, and frequency-dependent manner with the largest increases produced at subtetanic nerve stimulation frequencies.25 This effect was similar to what was observed in rat EDL studies.

In conclusion, HTS hit imidazopyrazinone 1 was identified using a fast skeletal ATPase assay and demonstrated improved calcium sensitivity and force production in muscle fibers. To enable in vivo pharmacological testing, 1 was optimized for biochemical potency and oral exposure, leading to compound 25 (tirasemtiv). Upon intravenous administration in preclinical models, tirasemtiv showed a concentration-dependent improvement in muscle force production in situ. In a Phase 3 clinical trial (VITALITY-ALS, NCT02496767) in patients with ALS, tirasemtiv did not meet its primary end point, change from baseline in slow vital capacity (SVC) at 24 weeks, or any of the secondary end points. Though not reaching statistical significance, the decline in SVC was smaller in patients who received any dose of tirasemtiv compared to patients receiving placebo. The largest differences from placebo were observed in patients randomized to the mid- and high-dose groups of tirasemtiv who could tolerate and remain on their target dose, suggestive of pharmacologic activity for the mechanism of action in ALS. More patients discontinued double-blind treatment on tirasemtiv than on placebo primarily due to nonserious adverse events related to tolerability, potentially confounding the assessment of its efficacy. The limitations of tirasemtiv may be addressed with CK-2127107, the next-generation fast skeletal muscle troponin activator currently being assessed in several midstage clinical trials.26

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Supporting Information
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Experimental procedures and analytical data (PDF)

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