Inhibition of CERS1 in skeletal muscle exacerbates age-related muscle dysfunction

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Keywords: sphingolipids, ceramide synthase 1 (Cers1), skeletal muscle, aging, sarcopenia, myoblast differentiation, myogenesis, P053, muscle genetics
ABSTRACT
Age-related muscle wasting and dysfunction render the elderly population vulnerable and incapacitated, while underlying mechanisms are poorly understood. Here, we implicate the CERS1 enzyme of the de novo sphingolipid synthesis pathway in the pathogenesis of age-related skeletal muscle impairment. In humans, CERS1 abundance declines with aging in skeletal muscle cells and, correlates with biological pathways involved in muscle function and myogenesis. Furthermore, CERS1 is upregulated during myogenic differentiation. Pharmacological or genetic inhibition of CERS1 in aged mice blunts myogenesis and deteriorates aged skeletal muscle mass and function, which is associated with the occurrence of morphological features typical of inflammation and fibrosis. Ablation of the CERS1 orthologue lagr-1 in C. elegans similarly exacerbates the age-associated decline in muscle function and integrity. We discover genetic variants reducing CERS1 expression in human skeletal muscle and Mendelian randomization analysis in the UK biobank cohort shows that these variants reduce muscle grip strength and overall health. In summary, our findings link age-related impairments in muscle function to a reduction in CERS1, thereby underlining the importance of the sphingolipid biosynthesis pathway in age-related muscle homeostasis.
INTRODUCTION

We are currently experiencing the biggest demographic shift in human history with a prospective doubling of the 65+ year old population by the year 2050, thereby constituting almost a quarter of the world’s population for the first time. While the increase in human lifespan could provide unprecedented opportunities for aged individuals, healthspan is not quite following this trend: One-fifth of an individual’s life will be lived with morbidity and reduced mobility. Health burden escalates with age, which not only incapacitates the elderly but also puts tremendous stress on the healthcare system exemplified by the 20% increased healthcare spending due to the growing elderly population in the last 10 years alone. Hence, understanding the underpinnings of aging is an important first step to inform development of biological tools to promote healthy aging.

Skeletal muscle is one of the organs most affected by aging. 3-8% muscle mass and strength are lost yearly after 30 years of age and 6-15% after 65 years of age during normal human aging. Exacerbated age-related loss of muscle mass and strength has recently been recognized as the disease sarcopenia, which predicts and correlates with mortality. The importance of sarcopenia is further highlighted by the high prevalence of sarcopenia in the common 60+ year old population (10-20%) especially considering that sarcopenia is one of the leading causes for frailty, loss of independence and reduced quality of life. Several mechanisms have been implicated in the skeletal muscle aging process, including reduced mitochondrial bioenergetics coupled with increased levels of reactive oxygen species, dysfunctional proteostasis, reduced hormone levels and signaling, nutritional intake, increased inflammation, loss of motor units, as well as reduced capacities for muscle regeneration and maturation/synthesis (for a review, see ). Recent evidence further suggests the involvement of sphingolipids in skeletal muscle homeostasis upon aging.

Sphingolipids are bioactive lipids exerting pleiotropic cellular functions such as inflammation, proliferation, myelination, cell growth, and cell death. Ceramides are the building blocks of most complex sphingolipids, which contain a sphingoid base (typically 18 carbon dihydrosphingosine or sphingosine) attached to a variable length fatty acyl side-chain. While sphingolipids have been shown to be involved in skeletal muscle bioenergetics, insulin sensitivity and diabetes, the contribution of ceramides to skeletal muscle aging is not well understood. We recently reported that the first enzyme of the de novo sphingolipid biosynthesis pathway (SPT) is involved in skeletal muscle aging. However, less is known about the involvement of ceramide synthases (Cers) in muscle aging, despite their prominent role in membrane homeostasis and cellular signaling. Six ceramide synthases (Cers1-6) catalyze ceramide synthesis in mammals resulting in a range of ceramides from C14:0 to C36:0. Depending on the enzyme, different length fatty acyl-coenzyme A (CoA) are transferred to the amine group of the sphingoid base. For example, Cers1 only uses 18 carbon (C18) fatty acids to create C18 (d18:1/18:0) ceramides, whereas Cers2 predominantly synthesizes d18:1/24:0 (C24:0) and d18:1/24:1 (C24:1) ceramides and Cers5 uses C16 acyl-CoA substrates for acylation of the free primary amine group of sphingoid bases to form the corresponding C16:0 ceramides. Cers1 and Cers5 are the two most expressed ceramide synthases in skeletal muscle. Reducing Cers1 or Cers5 function by genetic or pharmacological means has been shown to reduce adiposity and body weight upon exposing mice to a high fat diet. Moreover, BMI was found to correlate with C18:0 ceramide species, pointing towards Cers1 as a desirable target for metabolic interventions. On the other hand, recent evidence demonstrates that skeletal muscle-specific deletion of Cers1 in young mice reduces muscle fiber size and force, suggesting that Cers1 is needed for proper skeletal muscle function in young mice. However, Cers1’s involvement during organisal aging, its mechanistic function in skeletal muscle, and its relevance in human muscle are not well characterized.

In the present study, we show that CERS1 is involved in the age-related loss in muscle mass and function. In line with previous data, we find reduced CERS1 expression in old human donor myoblasts compared to myoblasts from young donors. Performing unbiased gene set enrichment analyses of CERS1-correlating
transcripts, we discover that CERS1 abundance in human skeletal muscle biopsies correlates with muscle function and myogenesis. Our experiments validate the increase in CERS1 abundance in mouse and human skeletal muscle cells during myogenic differentiation. Conversely, the selective pharmacological inhibitor of CERS1, P053\textsuperscript{11}, impairs myogenic maturation across several timepoints by blunting expression of myogenic regulatory factors and consequently, myosin heavy- and light chains. In aged mice, P053 treatment or AAV9-mediated silencing of Cers1 in skeletal muscle attenuates C18:0/C18:1 ceramide and dihydroceramide species, and exacerbates the age-related decline in muscle fiber size while elevating inflammation and fibrosis. These morphological alterations coincide with reduced myogenesis and reduced muscle mass function and mass. Lenti- or adenovirus-mediated CERS1 inhibition specifically in mouse or human muscle cells recapitulates maturation defects of muscle cells. Pharmacological or genetic inhibition of the CERS1 orthologue lagr-1 in C. elegans similarly impairs age-related muscle function and morphology suggesting conserved CERS1 function across species. Finally, we identify single nucleotide variants in the genomic CERS1 locus, which reduce CERS1 levels in skeletal muscle and reduce muscle grip strength as well as overall health in humans.

RESULTS

CERS1 expression correlates with muscle contraction and myogenesis

The sphingolipid synthesis pathway produces ceramides and other sphingolipids by using fatty acids and amino acids as substrates (Figure 1A). SPT converts L-serine and palmitoyl-CoA to 3-ketosphinganine, which is rapidly converted to sphinganine. Coupling of sphinganine to long-chain fatty acid is accomplished by one of 6 distinct mammalian ceramide syntheses. Among these, Cers1 expression in skeletal muscle tissue was shown to correlate negatively with aging\textsuperscript{21}. To expand on this finding and assess whether the age-related decline in CERS1 abundance is mediated specifically by skeletal muscle cells, we measured CERS1 expression in primary muscle cells isolated from young and old human donors. We found reduced CERS1 transcript levels in aged compared to young myoblasts which is in line with the previous report in whole muscle tissue\textsuperscript{21} (Figure 1B).

To study the role of CERS1 in human skeletal muscle, we performed unbiased gene set enrichment analyses of CERS1 correlating transcripts in human skeletal muscle biopsies. Molecular pathways involved in muscle contraction and myogenesis were among the strongest correlating pathways (Figure 1C). In line with this finding, we also observed significant positive correlations of CERS1 expression with transcripts involved in myogenesis (MYF5, MYH2, MYH7, MYL1; Figure 1D) and muscle contraction (TPM2, ACTA1, TMOD2, TNN1; Figure S1A). Assaying the myogenic differentiation process in mouse and human muscle cells confirmed the upregulation of Cers1/CERS1 at several timepoints once muscle differentiation was induced (Figure S1B and Figure 1E). CERS1 has been shown to synthesize C18:0 and C18:1 ceramides\textsuperscript{17,27,28}. Measurement of ceramide levels during myogenic differentiation confirmed particularly the increase of C18:1 during mouse myoblast maturation (Figure S1C), whereas C14:0, C22:0, as well as C18:0 ceramide species were found to be induced during human myoblast differentiation suggesting some species-specific effects (Figure S1D).

Importantly, the CERS1-specific pharmacological inhibitor P053, which has been shown to reduce fat mass\textsuperscript{11}, inhibited mouse and human skeletal muscle cell differentiation as evidenced by downregulated expression of the early expressed myogenic transcription factors Myog/MYOG and Myf6/MYF6 (Figure S2A and Figure 1F) as well as of myosin light- and heavy chains (Figure S2B and Figure S2C). In aggregate, these findings suggest that CERS1 is involved in skeletal muscle cell maturation.

Pharmacological inhibition of Cers1 impairs skeletal muscle in aged mice

We next assessed the systemic effect of the pharmacological Cers1 inhibitor P053 on skeletal muscle homeostasis in aged mice. As expected, systemic administration of P053 over 6 months (Figure 2A) reduced
C18:0 and C18:1 ceramides in skeletal muscle tissue (Figure 2B), suggesting that P053 indeed targeted Cers1 as previously shown\(^{11}\). Interestingly, this was coupled with an increase in C24:0/C24:1 ceramides (Figure 2B) and C24:0/C24:1 dihydroceramides (Figure S3A), which are synthesized by Cers2\(^{17,29,30}\). This might reflect a compensatory mechanism to maintain the flow of sphinganine and fatty acid substrates. An increase in C24:0/C24:1 ceramide and dihydroceramides upon Cers1 inhibition is in line with a previous reports on Cers1 inhibition\(^{11,21}\). Laminin staining (Figure 2C) followed by minimum ferret diameter measurement (Figure S3B) and fiber size distribution (Figure 2D) showed that fiber size was reduced upon aging with a frequency shift towards smaller muscle fibers. Treatment with P053 exacerbated this age-related reduction in muscle fiber size. We next automated the pipeline to detect skeletal muscle fibers in mouse tissue sections by custom training and application of the recently described deep learning-based segmentation algorithm cellpose\(^{31}\). Ground truth data, model training and validation, as well as the script to run on the data are publicly available on Zenodo (DOI: 10.5281/zenodo.7041137). Using our trained model in muscle sections stained with laminin and CD45, we were able to capture most muscle fibers (Average Precision: 0.97 ±0.02, n=4, Table S1). Results showed an age-related decrease in cross sectional area, which worsened with P053 treatment (Figure 2E, top and Figure S3C) and is in agreement with our manual curation of fiber diameter (Figure 2C-D). We also observed an increase in CD45\(^+\) neighboring muscle fibers with age and P053 treatment (Figure 2E, bottom). The effect of age and P053 on muscle inflammation was confirmed by another CD45 staining with DAPI showing an increased proportion of CD45\(^+\) cells upon aging, which was exacerbated by P053 administration (Figure S3D). Analyses of other morphological markers of skeletal muscle aging, fibrosis and centralized nuclei, revealed that pharmacological inhibition of Cers1 led to increased fibrosis (Figure 2F) and occurrence of centralized nuclei (Figure 2G). In line with our findings of differentiating skeletal muscle cell cultures (Figure 1F and Figure S2A-C), P053 treatment downregulated expression of myogenic regulatory factors and of myosin heavy chain transcripts in mice (Figure S4A). Of note, P053-mediated alterations in skeletal muscle morphology and expression of myogenic factors were correlated with impairments in skeletal muscle mass and function. Lean body mass was reduced upon aging and aged mice treated with P053 presented further reductions in muscle mass (Figure 2H), which is in line with reduced fiber diameters (Figure 2C-D) and cross section area (Figure S3C). Measurement of functional muscle parameters revealed that P053 administration worsened the effect of aging by reducing grip strength (Figure 2I) and treadmill running performance in aged mice (Figure S4B-C). Our observation of pharmacologic Cers1 inhibition in aged mice is in line with a previous study where genetic knockout of Cers1 in young mice impaired muscle contractility\(^{21}\).

**Muscle-specific genetic inhibition of CERS1 impairs aged muscle and blunts myogenesis**

Systemic administration of P053 in aged mice might have exerted effects on other tissues that affect aged skeletal muscle. Moreover, pharmacological strategies might have undesired off target effects. Therefore, we investigated the effect of direct genetic inhibition of Cers1 in aged mouse skeletal muscle. A single intramuscular injection of AAV9 with short hairpin RNA (shRNA) against mouse Cers1 (Figure 3A) in 18-month-old mice reduced Cers1 mRNA expression by \(\approx 50\%\) at 24 months of age (Figure S5A) and hence, reduced C18:0 and C18:1 ceramide metabolites (Figure 3B). Similarly to our observation in P053-treated animals, genetic inhibition of Cers1 in aged skeletal muscle tended to increase Cers2-derived very long chain ceramides (Figure 3B) and dihydroceramides (Figure S5B). Genetic silencing of Cers1 reduced average skeletal muscle fiber size (Figure 3C and Figure S5C), shifting muscle fibers towards smaller fibers (Figure 3D). This was in line with our custom cellpose model for fiber detection in muscle cross sections showing that Cers1-deficient, aged muscles had smaller cross-sectional area (Figure 3E, top and Figure S5D), and showed more signs of inflammation (Figure 3E, bottom and Figure S5E). Furthermore, ablation of Cers1 in aged muscle showed a trend towards increased fibrosis (Figure 3F) and more centralized nuclei (Figure 3G). We also found that genetic targeting of Cers1 downregulated expression of myogenic regulatory factors and myosin heavy chains (Figure S6A). These findings were coupled with reduced muscle mass (Figure 3H), as well as reduced muscle grip strength (Figure 3I) and running performance in Cers1-deficient aged mice (Figure
Reduced grip strength upon Cers1 inhibition in aged mice is in line with reduced ex vivo contraction capacity of young muscles deficient of Cers1. Overall, our results indicate that genetic, AAV9-mediated silencing of Cers1 in skeletal muscle of aged mice deteriorates skeletal muscle morphology and function and therefore, recapitulate the effects observed by pharmacological inhibition of Cers1.

Cers1 mediated deterioration of myogenesis is specific to skeletal muscle cells

Several different cell types reside within human and mouse skeletal muscle tissue. To investigate whether the observed detrimental effects of genetic Cers1 inhibition on muscle cell maturation are specific to skeletal muscle cells, we generated mouse and human skeletal muscle cells deficient of Cers1/CERS1. Lentivirus-mediated silencing of Cers1 using shRNA in mouse C2C12 muscle progenitor cells reduced Cers1 expression (Figure S7A) and with it, C18:0/C18:1 ceramide species (Figure S7B). This was coupled by an increase in potentially toxic very long chain ceramides (Figure S7B) and dihydroceramides (Figure S7C). Differentiation of these Cers1 deficient myoblasts towards myotubes revealed diminished myogenesis evidenced by immunostaining (Figure S7D), which shows reduced myotube diameter, myotube area and number of multinucleated myotubes (Figure S7E). Expression profiling revealed reduced transcript expression of myogenic differentiation markers (Figure S7F), further suggesting that Cers1 is indispensable for proper myogenic differentiation of mouse muscle cells. We next silenced CERS1 in isolated primary human myoblasts using adenovirus-mediated delivery of a shRNA construct targeting the human CERS1 transcript (Figure S8A). As expected, reducing CERS1 expression in human primary myoblasts reduced C18:0/C18:1 ceramides (Figure 4A). Similar to mouse tissue and cells, CERS1 inhibition increased very long chain C24:0/C24:1 ceramides (Figure 4A) and C24:0/C24:1 dihydroceramides (Figure S8B) and impaired myogenic differentiation, as indicated by immunostaining (Figure 4B) of differentiating myoblasts showing smaller myotube diameter, myotube area and reduction in multinucleated myotubes (Figure S8C-E) and expression profiling (Figure 4C). Therefore, CERS1 appears indispensable for intact myogenic maturation across human and mouse myoblasts.

Inhibition of the CERS1 orthologue lagr-1 in C. elegans deteriorates healthspan

Maintaining proper muscle function is vital to healthy aging and potent modulators of this process are typically conserved across species. Hence, the effect of inhibiting Cers1 on skeletal muscle function and morphology upon C. elegans aging was evaluated. We used the RW1596 transgenic worm strain expressing GFP under the control of the muscle-specific myo3p worm promoter, which allowed us to visualize muscle fibers as shown previously. Exposure of the RW1596 C. elegans strain to P053 mixed within the agar (50uM and 100uM) reduced muscle function at the onset of worm aging (day 5) when movement distance and speed are declining (Figure 4D and Figure S9A). In line with our observation of reduced muscle function when using P053 in mice, imaging muscle fibers of P053-treated worms showed deteriorated muscle morphology at day 5 as shown by the presence of rigid fibers compared to the smoother muscle fibers observed in the control worms that were exposed to DMSO vehicle (Figure 4E). We next evaluated the effect of genetic inhibition of the CERS1 orthologue lagr-1 on worm muscle function and morphology. Feeding RW1596 worms with an interference RNA (RNAi) targeting the lagr-1 transcript reduced its expression by ≈80% (Figure S9B), and reduced worm muscle function as measured by travelled distance and movement speed (Figure 4F and Figure S9C). Assessment of muscle fiber morphology revealed deteriorated muscle fibers in worms treated with lagr-1 RNAi at day 5 (Figure 4G). Taken together, these results suggest that pharmacological or genetic inhibition of the Cers1 orthologue impairs certain hallmarks of healthy worm aging as suggested by reduced motility and muscle fiber morphology. This is most evident at the onset of worm aging, suggesting that observed effects in Cers1 function might be conserved across species.

CERS1 genetic variants reduce muscle CERS1 expression and muscle function in humans
We next sought to relate our functional mouse and worm findings of Cers1/CERS1 inhibition impacting on muscle function to humans. To this end, we searched for both common (allele frequency > 1%) and rare genetic variants (allele frequency <1%) that alter CERS1 levels either indirectly via regulatory elements, or directly by mutations in the CERS1 coding region, respectively.

By exploring the Genotype-Tissue Expression (GTEx) dataset, we discovered three independent (linkage disequilibrium r²<0.1; Figure S9D), common genetic variants that significantly alter CERS1 expression in human skeletal muscle (Figure S9E). The C-allele at rs117558072 and the G-allele at rs1122821 reduce muscle CERS1 expression, while the T-allele at rs71332140 increases muscle CERS1 expression. We next evaluated how these skeletal muscle expression quantitative trait loci (eQTLs) might affect overall health, and muscle function by performing Mendelian randomization analysis using the UK biobank traits “overall health” (data-field 2178), “right grip strength” (data-field 47) and “left grip strength” (data-field 46). Combined results of the Mendelian randomization analysis showed that a decrease in muscular CERS1 expression leads to impaired overall health rating and grip strength (Figure 5A). In particular, the CERS1 muscle expression reducing alleles (rs117558072-C; rs1122821-G) decrease overall health, as well as right and left grip strength. Notably, we find a linear relationship between the eQTL-mediated effect sizes on muscle CERS1 expression and the eQTL-mediated effect sizes on the respective phenotypes (Figure 5B). Importantly, some of the CERS1 expression-reducing alleles are common in the human population, as indicated by their allele frequencies ranging from rs117558072-C(2%) to rs1122821-G(44%) and rs71332140-C(77%) and might therefore, contribute to skeletal muscle health in the broad population.

We next evaluated the effect of rare coding variants within the protein-coding exons of CERS1 on the same phenotypes in the UK biobank cohort. Assessing the association of such coding variants with overall health rating, right grip strength and left grip strength, we found that the large majority of the significant (suggestive significant association, p<0.05) variants in the CERS1 coding region (9 out of 11) showed negative effect sizes on overall health rating, right grip strength and left grip strength (Figure 5C). These data support the genetic finding of our Mendelian randomization analysis of common variants and therefore, underline the notion that CERS1 might affect overall health and muscle function in humans.

In conclusion, our study reveals the necessity of intact CERS1 expression for muscle cells to undergo muscle maturation, and that disruption of CERS1 expression might contribute to muscle dysfunction across different species. Due to our consistent observation of increased very long chain C24:0/C24:1 ceramides and dihydroceramides upon Cers1/CERS1 inhibition, we suggest a molecular network in which the compensatory upregulation of CERS2 derived very long chain ceramide and dihydroceramide C24:0/C24:1 species in aging might accelerate age-related muscle wasting and dysfunction, hence contributing to sarcopenia (Figure 5D).

DISCUSSION

In the present study, we sought to investigate the contribution of CERS1 to the skeletal muscle aging process. Our main findings show that I)CERS1 expression correlates positively with skeletal myogenesis and function in human muscle biopsies; II)intact CERS1 is necessary for proper muscle cell maturation; III)Cers1 is indispensable for intact organismal muscle aging, shown by deterioration of muscle mass, function and morphology upon genetic/pharmacologic inhibition of Cers1; and IV)common (5-70% allele frequency) and rare variants in humans affect muscle CERS1 and hence, muscle function and perception of one’s own health.

Our findings of reduced fiber size and muscle weight together with reduced muscle strength upon Cers1 knockdown in aged mice are in line with previous findings by Tosetti et al21 who show that Cers1 knockdown in young mice reduced muscle fiber size and contractility. Ceramide levels, and particularly Cers1-derived
C18-ceramides were previously found elevated in skeletal muscle upon high-fat diet\textsuperscript{34}, streptozotocin-induced diabetes\textsuperscript{35} and in obese, diabetic patients\textsuperscript{36}. \textit{Cers1} would therefore appear as an attractive target to improve metabolic outcomes due to its effect on C18 ceramide species. Indeed, the isoform-specific \textit{Cers1} inhibitor P053 reduced C18 ceramide species (while increasing C24 ceramides) exclusively in skeletal muscle, which reduced fat mass in mice fed a high-fat diet\textsuperscript{11}. However, our study shows that the improvements in fat mass loss might come at the expense of muscle function/myogenesis, particularly upon aging. We show that inhibition of \textit{Cers1} reduces myogenesis and exacerbates age-related muscle inflammation and fibrosis – key parameters associated with muscle aging. Furthermore, intact \textit{Cers1} might not only be important for skeletal muscle homeostasis in aging; \textit{Cers1} deficiency from birth was previously also reported to cause defects in foliation, progressive shrinkage, and neuronal apoptosis in the cerebellum, which translated to functional deficits including impaired exploration of novel objects, locomotion, and motor coordination\textsuperscript{27}.

An interesting observation was that \textit{Cers1} inhibition upregulated the levels of C24:0/C24:1 ceramides in all our \textit{Cers1} targeting strategies and muscle models. This is consistent with previous literature\textsuperscript{11,21} and suggests a possible compensatory upregulation of \textit{Cers2} to make up for \textit{Cers1} deficiency. Of note, compensation effects in brain and liver have also been reported in \textit{Cers2} knockout mice\textsuperscript{37,38}, and upregulation of \textit{Cers2} to compensate for \textit{Cers1} deficiency in neurons reduced long chain C24:0/C24:1 ceramides and hence, ameliorated \textit{Cers1} deficiency-mediated neurodegeneration\textsuperscript{39}. These studies combined with ours, therefore suggest that the long chain ceramides C24:0/C24:1 might exert detrimental effects at least in brain and skeletal muscle tissues. Confirmation of this hypothesis would open new avenues for targeted drug development, as \textit{Cers1} targeting shows conflicting effects with regards to metabolic and muscle health.

In summary, our study links \textit{Cers1} to both the muscle maturation process in mouse and humans and muscle aging. Disrupting \textit{Cers1}, and hence the myogenic potential in aged skeletal muscle, exacerbates the age-related decline in skeletal muscle mass, function, and morphology. Given the highly beneficial effects of Sptlc1 inactivation on healthy aging\textsuperscript{15}, our current study demonstrates that this mechanism does not involve \textit{Cers1}, and thus serves as an important signpost to rule out \textit{Cers1} inhibition as a therapeutic strategy for sarcopenia and age-related muscle dysfunction.
METHODS

In vivo mouse and worm studies

Animal experiments. Young (3mo) and aged (18-month-old) male C57BL/6JRj mice were obtained from Janvier Labs. Mice were randomized to the respective treatment groups according to their body weight. Animals were fed a standard chow diet. All animals were housed in micro-isolator cages in a room illuminated from 7:00AM to 7:00PM with ad libitum access to diet and water. The dose of P053 or DMSO was 5 mg/kg 3 times a week. Adeno-associated virus (AAV) 9 was bilaterally injected in gastrocnemius muscle at 18-month of age with 2 x 10^{11} viral particles (Vector Biolabs, United States) containing scramble short hairpin RNA (shRNA) or Cers1-targeting shRNA. 5-6 months after pharmacological or gene-modifying injections, phenotypic tests were performed on the animals. At sacrifice, muscles were removed for histochemical analyses or snap frozen in liquid N2 for biochemical assays. Use of animals for all experimental studies were approved by animal licenses 2890.1 and 3341 in Canton of Vaud, Switzerland and were in compliance with the 1964 Declaration of Helsinki and its later amendments.

Endurance running test. After familiarizing with the treadmill, the exercise regimen started at a speed of 15 cm/s. Every 12 minutes, the speed was increased by 3 cm/s. Mice were considered to have reached their peak exercise capacity, and removed from the treadmill if they received 7 or more shocks (0.2 mA) per minute for two consecutive minutes. The distance traveled and time before exhaustion were registered as maximal running distance and time as shown previously^{22}.

Grip strength. Muscle strength was assessed by the grip strength test as previously described^{23}. Grip strength of each mouse was measured on a pulldown grid assembly connected to a grip strength meter (Columbus Instruments). The mouse was drawn along a straight line parallel to the grip, providing peak force. The experiment was repeated three times, and the highest value was included in the analysis.

Measurement of sphingolipids. Cell pellets (~1.0e6 cells) were extracted by the addition of 100 μL of MeOH spiked with the stable isotope-labelled internal standards (Spa(d17:0), Cer(d18:1/16:0)-d9, Cer(d18:1/18:0)-d7, Cer(d18:1/24:0)-d7 and Cer(d18:1/24:1)-d7)). Sample homogenization was performed in the Cryolys Precellys Tissue Homogenizer (2 x 20 seconds at 10000 rpm, Bertin Technologies, Rockville, MD, US) with ceramic beads. The bead beater was air-cooled down at a flow rate of 110 L/min at 6 bar. Homogenized extracts were centrifuged for 15 minutes at 4000 g at 4°C and the resulting supernatants were collected for the LC-MS/MS analysis. Sphingolipids were quantified by LC-MS/MS analysis in positive ionization mode using a 6495 triple quadrupole system (QqQ) interfaced with 1290 UHPLC system (Agilent Technologies), adapted from Checa et.al.^{24}. Briefly, the chromatographic separation was carried out in a Zorbax Eclipse plus C8 column (1.8 μm, 100 mm × 2.1 mm I.D) (Agilent technologies). Mobile phase was composed of A = 5 mM ammonium formate and 0.2 % formic acid in water and B = 5 mM ammonium formate and 0.2% formic acid in MeOH at a flow rate of 400 μL/min. Column temperature was 40 °C and sample injection volume 2μL. The linear gradient elution starting from 80% to 100% of B (in 8 min) was applied and held until 14 min. The column was then equilibrated to initial conditions. ESI source conditions were set as follows: dry gas temperature 230 °C, nebulizer 35 psi and flow 14 L/min, sheath gas temperature 400 °C and flow 12 L/min, nozzle voltage 500 V, and capillary voltage 4000 V. Dynamic Multiple Reaction Monitoring (dMRM) was used as acquisition mode with a total cycle time of 500 ms. Optimized collision energies for each metabolite were applied. Raw LC-MS/MS data was processed using the Agilent Quantitative analysis software (version B.07.00, MassHunter Agilent technologies). For absolute quantification, calibration curves and the stable isotope-labelled internal standards (IS) were used to determine the response factor. Linearity of the standard curves was evaluated for each metabolite using a 12-point range, in addition, peak area integration was manually curated and corrected when necessary. Sphingolipid concentrations were reported to protein concentrations measured in protein pellets (with BCA, Thermo Scientific) following the metabolite extraction.

Histology. Skeletal muscles were harvested from anesthetized mice, and immediately embedded in Thermo Scientific™ Shandon™ Cryomatrix™ and frozen in isopentane, cooled in liquid nitrogen, for 1 min before
being transferred to dry ice and stored at -80 °C. 8 µm cryosections were incubated in 4% PFA for 15 min, washed three times for 10 min with PBS, counterstained with DAPI, laminin (1:200, Sigma), CD45 (1:200, Life Technologies), coupled with Alexa-488 or Alexa-568 fluorochromes (Life Technology) and mounted with Dako Mounting Medium. Images were performed with VS120-S6-W slides scanner (Olympus) acquired using a 20x/0.75 air UPLS APO objective and an Olympus XM10 CCD camera. 3 channel images (DAPI, Laminin, and CD45-A568) were acquired defining a focus map based on the Laminin signal for fluorescence imaging whereas hematoxylin and eosin and Sirius Red were imaged with brightfield. Resulting VSI images could be opened in Fiji/ImageJ and QuPath using the BioFormats library. Minimum Feret diameter, and cross-sectional area were determined using the ImageJ software. For fiber size distribution, a minimum of 2,000 fibers were used for each condition and measurement. The minimum Feret diameter is defined as the minimum distance between two parallel tangents at opposing borders of the muscle fiber. This measure has been found to be resistant to deviations away from the optimal cross-sectioning profile during the sectioning process.

**Deep learning-based muscle segmentation and quantification.** A subset of 4 muscle cross-section images was manually annotated in multiple regions to serve as ground-truth for cellpose training. In total, 15 areas were used for training and 4 for validation. Annotation and training was done through QuPath using the QuPath cellpose extension (https://github.com/BIOP/qupath-extension-cellpose). Original images were exported as “raw image plus labeled image” pairs and downsampled 4x in X and Y (Final pixel size for training: 1.2876 um/px). The cellpose model was trained with the starting weights of cellpose’s ‘cyto2’ model, for 500 epochs. Predicted validation images were further run through a validation notebook derived from ZeroCostDL4Mic<sup>25</sup> in order to assess the quality of the predictions (see Table S1). Segmentation of muscle fibers was performed in two steps. First, the whole cross-section was detected though a gaussian-blurred version of the Laminin signal (sigma=5 um), followed by an absolute threshold (threshold value: 5). The resulting region was then fed into the QuPath cellpose extension along with our custom model for individual fiber segmentation. Training images, training and validation scripts and Jupyter notebooks, as well as an example QuPath project are available on Zenodo (DOI: 10.5281/zenodo.7041137).

**C. elegans movement and muscle morphology.** *C. elegans* strains were cultured at 20°C on nematode growth medium (NGM) agar plates seeded with *E. coli* strain OP50 unless stated otherwise. Strains used in this study were the transgenic RW1596 strain, which was provided by the Caenorhabditis Genetics Center (University of Minnesota). RW1596 expresses GFP under the control of the worm skeletal muscle promoter myo3 (myo3p-GFP). The *Cers* orthologue *lagr-1* was found on wormbase (https://wormbase.org/). For RNAi experiments, worms were exposed to *lagr-1* RNAi or an empty vector control plasmid using maternal treatment to ensure robust knock down. P053 was dissolved in DMSO, and used at a final concentration of 50uM and 100uM. Worms were exposed to P053 starting from L4 stage. P053 or DMSO was added to agar before preparing plates. To ensure permanent exposure to the compound, plates were changed twice a week. *C. elegans* movement analysis was performed using the Movement Tracker software as done previously<sup>23</sup>. The experiments were repeated at least twice. A population of ~50 worms was washed in M9, immobilized using a 7.5 mM solution of tetramisole hydrochloride (Sigma-Aldrich) and immediately imaged afterwards as shown previously. Confocal images were acquired with Zeiss LSM 700 Upright confocal microscope (Carl Zeiss AG) under non-saturating exposure conditions. Image processing was performed with the Fiji software.

**In vitro mouse and human studies**

**Cell culture and cell transfection.** The C2C12 mouse myoblast cell line was obtained from the American Type Culture Collection (CRL-1772TM). C2C12 cells or clones were cultured in growth medium consisting of Dulbecco’s modified Eagle’s medium (Gibco, 41966-029), 20 % Fetal Bovine Serum (Gibco, 10270-106) and 100 U/mL penicillin and 100 mg/mL streptomycin (Gibco, 15140-122). To induce differentiation, FBS was substituted with 2% horse serum (Gibco, 16050-122). Trypsin-EDTA 0.05% (GIBCO, 25300-062) was used to detach cells. Primary human skeletal muscle cells were obtained from Lonza (SKMC, #CC-2561) and the Hospices Civils de Lyon and were cultured in growth medium consisting of DMEM/F12 (Gibco,
10565018), 20 % Fetal Bovine Serum (Gibco, 10270-106) and 100 U/mL penicillin and 100 mg/mL streptomycin (Gibco, 15140-122). To induce differentiation, FBS was exchanged with 2 % horse serum (Euroclone) and kept in culture. All cells were maintained at 37 °C with 5% CO2. Cell transfections were done using TransIT-T2 (Mirus) according to the manufacturer’s protocol with a 3:1 ratio of transfection agent to DNA. C2C12 cells were grown confluent, and 1 μM P053 or DMSO was added, and cells were kept in growth medium for another 3 days to measure muscle cell differentiation. Cell lines were regularly tested for mycoplasma contamination.

**RNA isolation and real-time qPCR.** Tissues were homogenized with Trizol, whereas cells were homogenized, and RNA isolated using the RNeasy Mini kit (Qiagen, 74106). Reverse transcription was performed with the High-Capacity RNA-to-cDNA Kit (4387406 Thermofisher scientific) as shown previously26. Gene expression was measured by quantitative reverse transcription PCR (qPCR) using LightCycler 480 SYBR Green I Master (50-720-3180 Roche). Quantitative polymerase chain reaction (PCR) results were calculated relative to the mean of the housekeeping gene Gapdh. The average of two technical replicates was used for each biological data point. Primer sets for qPCR analyses are shown in the Table S2.

**Lentivirus production and transduction.** Lentiviruses were produced by cotransfecting HEK293T cells with lenti plasmids expressing scramble shRNA or Cers1 targeting shRNA (see plasmid list in Table S3), the packaging plasmid psPAX2 (addgene #12260) and the envelope plasmid pMD2G (addgene #12259), in a ratio of 4:3:1, respectively. Transfection medium was removed 24h after transfection and fresh medium was added to the plate. Virus containing medium was collected at 48h and concentrated using Lenti-X™ Concentrator (Takara). C2C12 mouse myoblasts were seeded 24h prior to infection and then transduced with virus-containing supernatant supplemented with 8μg/mL polybrene (Millipore) as shown previously22. For the delivery of a shRNA construct targeting CERS1 in human primary myoblasts, adenovirus containing Ad-U6-h-CERS1-shRNA or Ad-U6-scrmb-shRNA (Vector Biolabs) was infected using 1uL adenovirus per mL cell culture medium at a titer of 5x10^10 PFU/ml.

**Immunocytochemistry.** C2C12 cells or Lonza SkMU cells cultured on a sterilized coverslip in 6-well plates (Greiner bio-one, CELLSTAR, 657160) were fixed in Fixx solution (Thermo Scientific, 9990244) for 15 min and permeabilized in 0.1% Triton X-100 (Amresco, 0694) solution for 15min at 20°C. Cells were blocked in 3% BSA for 1h at 20°C to avoid unspecific antibody binding and then incubated with primary antibody over night at 4°C with gentle shaking. MyHC was stained using the MF20 primary antibody (1:200, Invitrogen, 14-6503-82) for C2C12 cells and in Lonza muscle cells with a MYL1 antibody (1:140, Thermofisher, PA5-29635). Antibodies used in this study are shown in Table S4. The next day cells were incubated with secondary antibody (Thermo Fisher #A10037 for MF20 and #A-21206 for MYL1) for 1h at 20°C and nuclei were labeled with DAPI. The immunofluorescence images were acquired using either fluorescence or confocal microscopy. The myofusion index was calculated as the ratio of nuclei within myotubes to total nuclei. Myotube diameter was measured for 8 myotubes per image using ImageJ. Myotube area was calculated as the total area covered by myotubes.

**Human studies**

**Skeletal muscle gene expression in Genotype-Tissue Expression (GTEx) project.** The Genotype-Tissue Expression (GTEx) project version 8 was accessed through accession number dbGaP phs000424.v8.p2 (approved request #10143-AgingX). Transcript expressions and covariates from male human muscle of 469 individuals were extracted from the GTEx Portal (https://gtexportal.org). To remove the known and hidden factors, the Probabilistic Estimation of Expression Residuals (PEER) was applied the expression residuals were used to further analysis. For gene set enrichment analysis, genes were ranked by Spearman correlation coefficients with the gene expression level of CERS1 based on the Spearman correlation approach, and the CERS1 enriched gene sets were calculated by Gene Set Enrichment Analysis (GSEA) using clusterProfiler R package (version 3.10.1). Gene ontology (GO), Kyoto Encyclopedia of Genes and Genomes (KEGG),
Reactome and Hallmark gene sets were retrieved from the Molecular Signatures Database (MSigDB) using the msigdb R package (version 7.2.1).

**Mendelian randomization.** We used inverse variance-weighted Mendelian randomization to assess the causal effect of *CERS1* expression on muscle-related traits in humans. As exposures, we used the GTEx version 8 cis-eQTLs that were mapped by the GTEx consortium based on the normalized gene expressions in skeletal muscle from 706 deceased human individuals. These summary statistics are publicly available and can be accessed on the internet at the online GTEx webpage portal through [https://storage.googleapis.com/gtex_analysis_v8/single_tissue_qtl_data/GTEx_Analysis_v8_eQTL.tar](https://storage.googleapis.com/gtex_analysis_v8/single_tissue_qtl_data/GTEx_Analysis_v8_eQTL.tar). As outcomes, we used 10648 publicly available summary statistics obtained from various sources through the “available_outcomes” function from the TwoSampleMR R package. Outcome IDs can be queried from the ieu gwas database at [https://gwas.mrcieu.ac.uk/](https://gwas.mrcieu.ac.uk/). We restricted the analysis to independent instrumental variables (genetic variants). We therefore followed an iterative pruning workflow whereby we included the variant with the most significant effect on *CERS1* expression, followed by pruning with a 0.05 squared correlation coefficient (r²) threshold, then again selecting the most significant variant, and so on until no variants are left. This procedure retained three genetic variants: rs117558072 (A > C), rs1122821 (A > G), and rs71332140 (C > T). The causal effect of altered *CERS1* expression was inferred on the 10648 outcomes through inverse-variance weighted Mendelian randomization. The false discovery rate was controlled at the 5% significance level with the Benjamini-Hochberg procedure.

**UK Biobank (UKBB) rare variant analysis.** We performed genetic association analyses for hand grip strength and overall health in UK Biobank participants from 379,530 unrelated individuals. To run GWAS on the UK Biobank 450k Whole Exome Sequencing release (data field 23150) we filtered variants based on allele frequency and Hardy-Weinberg equilibrium. We then used Regenie v3.1.1 and accounted for sex, age, and body mass index. First, population structure estimation was performed using array sequencing to equally represent the whole genome. Second, GWAS burden tests were employed to measure SNP effects on the UKBB phenotypes overall health, right grip strength and left grip strength using the UKBB whole exon sequencing data set. The UKBB data was accessed under the application #48020.

**Statistical analyses.** For experimental conditions in which there were two independent factors and multiple comparisons, a factorial ANOVA with subsequent post hoc analysis was performed. Where appropriate, one-way ANOVA with post hoc analysis or t-tests was performed. All statistical analyses were performed using GraphPad Prism (9.4.0, San Diego, CA, USA). Results are reported as means ± standard error of the mean. Statistical significance was set to P < 0.05.
Acknowledgments

We wish to thank the staff of EPFL histology, bioimaging and optics (BIOP), flow cytometry, and animal facilities for technical assistance. We also wish to acknowledge Tony Teav (from Metabolomics Platform at UNIL) for his help with the sphingolipid measurements. This research has been conducted using the UK Biobank Resource. The work in the JA laboratory was supported by grants from the Ecole Polytechnique Federale de Lausanne (EPFL), the European Research Council (ERC-AdG-787702), the Swiss National Science Foundation (SNSF 31003A_179435), the Fondation Suisse de Recherche sur les Maladies Musculaires (FSRMM) and the Fondation Marcel Levaillant (190917). MW’s position was supported by Central Norway Regional Health Authority. BC was supported by FAPESP (2019/22512-0). LJEG was supported by the European Union’s Horizon 2020 research and innovation program through the Marie Sklodowska-Curie Individual Fellowship “AmyloAge” (grant agreement no. 896042).

Author contributions

The study was conceived and designed by MW, PPL and JA. Human and mouse bioinformatics analyses were performed by XL, GA and MW. Mouse phenotyping and injections were carried out by MW and PPL. In vitro work was performed by MW, BC, RM. HGA and JI performed sphingolipid measurements. Histological analyses and ex vivo measurements were performed by BC, RM and MW. Genetic analyses were performed by LJEG, GA and MW. MW and JA wrote the manuscript, and all authors gave critical comments on it. JA supervised the work.

Conflict of interest

PPL and JA are inventors on a patent application filed by EPFL covering the use of ceramide synthesis inhibitors for muscle disorders. The other authors do not declare a conflict of interest.
FIGURE LEGENDS

Figure 1. CERS1-correlated pathways in human skeletal muscle associate with muscle function and myogenesis. (A) Overview over the de novo sphingolipid biosynthesis pathway. (B) CERS1 expression in young (22 ± 3.61yo) and old (74 ± 8.4yo) human skeletal muscle cells (n=3-5 biological replicates). (C) Gene set enrichment analysis of CERS1 mRNA correlated transcripts in the human skeletal muscle from the Genotype-Tissue Expression (GTEx) dataset (n=469). Normalized effect size (NES). (D) Spearman correlation between skeletal muscle expression of CERS1 and the first principal component (PC1) of the myogenesis pathway (left) from (C) and expression of key genes involved in myogenesis (right). (E-F) Gene expression of enzymes involved in the sphingolipid de novo synthesis pathway (E) and myogenic regulatory factors (F) upon horse serum induced differentiation of primary human skeletal muscle cells (n=2-3 biological replicates). The pharmacological inhibitor P053 or DMSO was used at 1µM in (E-F). Data: mean ± SEM. *P < 0.05, **P < 0.01, ****P < 0.0001.

Figure 2. P053 administration inhibiting Cers1 in aged mice deteriorates skeletal muscle function and morphology. (A) Overview over the experimental pipeline that was used to administer P053 in aged C57BL/6J mice using intraperitoneal injections (I.P) three times a week for 6 months. (B) Skeletal muscle ceramide levels in aged mice treated with DMSO or P053 (n=7-8 per group). (C) Representative images of laminin-stained tibialis anterior of young control or mice injected with DMSO or P053 (n=5-6 per group). Scale bar, 50µM. (D) Quantification of minimal Ferret diameter distribution in skeletal muscle cross sections from young control or mice injected with DMSO or P053 (n=5-6 per group). (E) Automated detection of laminin/CD45+ stained muscle fibers from young control or aged mice injected with DMSO or P053 showing cross sectional area (top) and inflammation-adjacent signal intensities (bottom). (F) Representative brightfield images and quantification of Sirus red stained muscle sections from young and aged mice treated with DMSO or P053 (n=6 per group). (G) Representative brightfield images and quantification of muscle cross sections from young and aged mice treated with DMSO or P053 stained with hematoxylin/eosin (n=6 per group). (H-I) Phenotyping measurements of young control and aged mice injected with DMSO or P053 showing (H) lean body mass and (I) grip strength (n=7-14 per group). Data: mean ± SEM. *P < 0.05, **P < 0.01, ***P < 0.001.

Figure 3. Adeno-associated virus 9 (AAV9) mediated knockdown of Cers1 expression in aged skeletal muscle reduces skeletal muscle function and morphology. (A) Schematic showing a single gastrocnemius intramuscular injection of adeno-associated virus particles containing short hairpin RNA against Cers1 in aged C57BL/6J mice. (B) Skeletal muscle ceramide levels in aged mice intramuscularly injected with shRNA targeting Cers1 (n=6-8). (C) Representative images of laminin-stained aged mice muscle cross sections intramuscularly injected with shRNA targeting Cers1 (n=6 per group). Scale bar, 50µM. (D) Quantification of minimal Ferret diameter distribution in aged mice intramuscularly injected with shRNA targeting Cers1 (n=6 per group). (E) Automated detection of laminin/CD45+ stained skeletal muscle fibers from aged mice intramuscularly injected with shRNA-Cers1 showing cross sectional area (top) and inflammation-adjacent signal intensities (bottom). (F) Representative brightfield images and quantification of Sirus red stained skeletal muscle cross sections from mice intramuscularly injected with AAV9 particles containing scramble, or shRNA targeting Cers1 (n=6 per group). (G) Representative brightfield images and quantification of hematoxylin/eosin-stained muscle cross sections from mice intramuscularly injected with AAV9 particles containing scramble, or shRNA targeting Cers1 (n=6 per group). (H-I) Phenotyping measurements of aged mice intramuscularly injected with shRNA-Cers1 or a scramble shRNA with (H) gastrocnemius muscle weight and (I) grip strength (n=6-8 per group). Data: mean ± SEM. *P < 0.05, **P < 0.01, ***P < 0.001.

Figure 4. Inhibition of Cers1 in muscle cells impairs myogenic differentiation and C. elegans healthspan. (A) Primary human skeletal muscle cell ceramide levels injected with adenovirus containing silencing RNA targeting CERS1 (n=5-6 per group). (B) Representative confocal immunocytochemistry images of differentiating human primary muscle cells deficient of CERS1 (n=4 per group, left). (C) Gene expression profiling of differentiating human primary muscle cells deficient of CERS1 (n=4-5 per group, right). Scale bar, 50µM. (D) Travelled distance in transgenic RW1596 C. elegans (myo3p-GFP) treated with DMSO, 50uM P053 or 100uM P053 (n=40-57 per group). (E) Representative confocal muscle images of RW1596 C. elegans (myo3p-GFP) treated with DMSO or 100uM P053 (n=6 per group). (F) Travelled distance in transgenic RW1596 C. elegans (myo3p-GFP) treated with EV control, or RNAi against lagr-1 (n=46-56 per group). (G)
Representative confocal muscle images of RW1596 C. elegans (myo3p-GFP) treated with control, or RNAi against lagr-1 (n=6 per group). Data: mean ± SEM. *P < 0.05, **P < 0.01, ***P < 0.001, ****P<0.0001.

Figure 5. Common and rare genetic variants in CERS1 affect muscle function and health in humans. (A) Overall result of the Mendelian randomization analysis in the UK biobank cohort using skeletal muscle expression quantitative trait loci (cis-eQTL) of CERS1. (B) Scatter plot showing the effect of the three independent (R²<0.1, see Figure S9D) cis-eQTLs rs117558072, rs1122821, rs71332140 on overall health, right grip strength, and left grip strength in the UK biobank. The slope of the regression line depicts the estimated causal effect with the inverse-variance weighted Mendelian randomization method. (C) Lollipop plot depicting rare variants in the coding region of CERS1 and their effects on overall health, right grip strength, and left grip strength in the UK biobank. Colors indicate phenotypes, dot size indicates effect size, red dotted line indicates the suggested cut-off p<0.05. (D) Overview over the de novo sphingolipid biosynthesis pathway highlighting the hypothesis that CERS1 inhibition leads to a compensatory upregulation of CERS2, which might inhibit muscle function in aging. Data: mean ± SEM. P values are shown in the figure panels.
FIGURE LEGENDS (Supplementary Figures)

**Figure S1.** Cers1 regulates muscle contraction in human muscle and myogenesis in mouse myoblasts. (A) Spearman correlation between skeletal muscle expression of CERS1 and the first principal component (PC1) of the muscle contraction pathway (left) and expression of key genes involved in muscle contraction (right). (B) Gene expression of enzymes involved in the sphingolipid de novo synthesis pathway upon horse serum induced differentiation of mouse C2C12 myoblasts (n=3-4). (C-D) Ceramide levels during myogenic differentiation of mouse C2C12 myoblasts (n=3-4) (C) and human primary skeletal muscle cells (n=2-3) (D). Data: mean ± SEM. ****P<0.0001.

**Figure S2.** The Cers1 inhibitor P053 deteriorates myoblast maturation. (A) Gene expression of myogenic regulatory factors upon horse serum induced differentiation of mouse C2C12 myoblasts (n=3-4). (B-C) Gene expression of myosin light- and heavy chains upon horse serum induced differentiation of isolated human primary myoblasts (n=2-3 biological replicates) (B) and mouse C2C12 (n=3-4) muscle cells (C) upon horse serum induced myoblast differentiation. The pharmacological inhibitor P053 or DMSO was used at 1μM in (A-C). Data: mean ± SEM. *P < 0.05, **P < 0.01, ***P < 0.001, ****P<0.0001.

**Figure S3.** The Cers1 inhibitor P053 exacerbates mouse skeletal muscle aging. (A) Skeletal muscle dihydroceramide levels in aged mice treated with DMSO or P053 (n=7-8 per group). (B) Average minimal Ferret diameter in muscle of young and aged mice treated with DMSO or P053 (n=5-6 per group). (C) Quantification of muscle cross-sectional area in young and aged mice treated with DMSO or P053 (n=5-6 per group). (D) Representative immunohistochemistry images of muscle cross sections from young and aged mice treated with DMSO or P053 stained with DAPI and CD45 (n=5-6 per group). Scale bar, 50μM. Data: mean ± SEM. *P < 0.05, **P < 0.01, ***P < 0.001.

**Figure S4.** The Cers1 inhibitor P053 deteriorates mouse skeletal muscle aging. (A) Gene expression in young or aged muscle of mice treated with DMSO or P053 (n=5-6 per group). (B-C) Phenotyping of young control and aged mice injected with DMSO or P053 showing running time (B), and running distance (C) (n=7-14 per group). Data: mean ± SEM. *P < 0.05, **P < 0.01.

**Figure S5.** Genetic inhibition of Cers1 exacerbates mouse skeletal muscle aging. (A) Cers1 RNA expression in aged mouse muscle intramuscularly injected with scramble, or shRNA targeting Cers1 (n=6 per group). (B) Dihydroceramide levels in gastrocnemius muscle injected with AAV9 particles containing scramble, or shRNA targeting Cers1 (n=6-8 per group). (C) Average Ferret diameter in skeletal muscle of aged mice intramuscularly injected with scramble, or shRNA targeting Cers1 (n=6 per group). (D) Quantification of cross-sectional area in aged mice treated with AAV9-shRNA Cers1 (n=6 per group). (E) Representative immunohistochemistry images of muscle cross sections of CD45-stained muscle from mice intramuscularly injected with AAV9 particles containing scramble, or shRNA targeting Cers1 (n=6 per group). Data: mean ± SEM. *P < 0.05, **P < 0.01.

**Figure S6.** Genetic inhibition of Cers1 deteriorates mouse skeletal muscle aging. (A) Muscle gene expression in of mice intramuscularly injected with AAV9 particles containing scramble, or shRNA targeting Cers1 (n=6 per group). (B-C) Phenotyping measurements of aged mice intramuscularly injected with shRNA-Cers1 shows running time (B) and running distance (C) (n=6-8 per group). Data: mean ± SEM. *P < 0.05.

**Figure S7.** Lentivirus mediated silencing of Cers1 reduces ceramides and blunts myogenesis in mouse muscle cells. (A) Cers1 gene expression in C2C12 mouse muscle cells transduced with lentivirus containing scramble, or shRNA targeting Cers1 (n=5-6 biological replicates per group). (B) C2C12 myoblast ceramide levels upon lentivirus mediated silencing of Cers1 (n=3 per group). (C) Dihydroceramide levels in C2C12 cells transduced with lentivirus containing shRNA targeting Cers1 (n=3 per group). (D) Representative immunocytochemistry images of differentiating C2C12 muscle cells deficient of Cers1 (n=6 per group). Scale bar, 50μM. (E) Quantification of (D) showing myotube diameter (left), myotube area (middle) and % of multinucleated myotubes (right) in C2C12 mouse muscle cells transduced with lentivirus containing scramble, or shRNA targeting Cers1 (n=6 per group). (F) Gene expression profiling of differentiating C2C12 muscle cells deficient of Cers1 (n=8-12 per group). Data: mean ± SEM. *P < 0.05, **P < 0.01, ***P < 0.001, ****P<0.0001.
Figure S8. Adenovirus mediated silencing of CERS1 reduces ceramides and blunts myogenesis in human muscle cells. (A) CERS1 gene expression in isolated primary human muscle cells transduced with adenovirus containing scramble, or shRNA targeting CERS1 (n=3 per group). (B) Dihydroceramide levels in differentiated primary human muscle cells transduced with adenovirus containing scramble, or shRNA targeting CERS1 (n=5-6 per group). (C-E) Quantification of myotube diameter (C), myotube area (D), and % of multinucleated myotubes (E) in differentiated primary human muscle cells transduced with adenovirus containing scramble, or shRNA targeting CERS1 (n=4 per group). Data: mean ± SEM. *P < 0.05, **P < 0.01.

Figure S9. CERS1 inhibition is associated with impaired health span. (A) Mean speed measured in transgenic RW1596 C. elegans (myo3p-GFP) treated with DMSO, 50uM P053 or 100uM P053 (n=40-57 per group). (B) Gene expression of the Cers1 orthologue lagr-1 in transgenic RW1596 C. elegans (myo3p-GFP) treated with empty vector control, or RNAi targeting lagr-1 (n=3 biological replicates per group). (C) Mean speed measured in transgenic RW1596 C. elegans (myo3p-GFP) treated with EV control, or RNAi against the Cers1 orthologue lagr-1 (n=46-56 per group). (D) Linkage disequilibrium of the common CERS1 expression quantitative trait loci (cis-eQTL) rs117558072, rs1122821, rs71332140 measured as r² in all populations (https://ldlink.nci.nih.gov). (E) Violin plots showing the allelic effect of rs117558072 (left), rs1122821 (middle), rs71332140 (right) on CERS1 expression in human skeletal muscle in the Genotype-Tissue Expression (GTEx) dataset (n=706 human in total). Data: mean ± SEM. *P < 0.05, **P < 0.01, ***P < 0.001.
**Table S1.** Cellpose training quality control.

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**Table S2.** List of mouse and human qPCR primers.

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**Table S3.** List of plasmids.

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<td>pMD2G</td>
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## Table S4. List of antibodies.

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<td>Laminin</td>
<td>Sigma</td>
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<td>CD45 (1:200, eBioscience, eFluor450 conjugated)</td>
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<td>Donkey anti-Mouse IgG secondary antibody</td>
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REFERENCES

**Figure 1**

(A) Serine + palmitoyl-CoA → 3-ketosphinganine → Sphinganine → Dihydroceramide → Ceramide

(B) CERS1 gene expression (FC) vs. YOUNG vs. OLD

(C) Reactome neurotransmitter release cycle

(D) R = 0.70, p < 2.2^{-16}

(E) CERS1***, DEGS1*, SPTLC1***, CERS2*

(F) Gene expression (FC) vs. Time (h)

-log10(q-values)

CERS1 expression

Ratios:

- MYOD Control
- MYOG Control
- MYF5 Control
- MYF6 Control

Correlations:

- MYF5 vs. MYH2: R = 0.23, p = 5.4^{-7}
- MYF5 vs. MYH7: R = 0.65, p < 2.2^{-16}
- MYF5 vs. MYL1: R = 0.58, p < 2.2^{-16}
- MYF5 vs. MYF6: R = 0.54, p < 2.2^{-16}
- MYF5 vs. MYOD: R = 0.85, p < 2.2^{-16}

Hedgehog Pathway:

- Hedgehog on state
- Hedgehog off state
- Degradation of axin

Nurse Cell Endocytosis:

- Cross presentation of soluble exogenous antigens endosomes
- Transition to the post-replicative state
- Degradation of axin

Other Pathways:

- Apoptosis by caspases
- Activation of the mTOR pathway
- Regulation of the cell cycle

Cellular Responses:

- Apoptosis by caspases
- Activation of the mTOR pathway
- Regulation of the cell cycle
Figure 2

(A) P053 Cers1 inhibitor I.P.

(B) Phenotyping

(C) Young

(D) Lean body mass (%)**

(E) CD45 intensity (AU)

(F) Fibrosis score

(G) % centralized nuclei

(H) Grip strength (g)***

(I) Lean body mass (%)***

(A) (B) (C) (D) (E) (F) (G) (H) (I)
Figure 3

(A) AAV9-Cers1 shRNA

(B) Muscle weight (g)

(C) Aged + AAV con

(D) % fibers

(E) CSA (µm²)

(F) Aged + AAV con

(G) Fibrosis score

(H) Muscle weight (g)

(I) Grip Strength (g)

* p < 0.05
** p < 0.01
*** p < 0.001
**** p < 0.0001

(CD45 (AU))

(L) % centralized nuclei

* p < 0.05
**Figure 4**

(A) Bar graph showing ceramide levels (nmol/mg protein) for different ceramide species (C14:0, C16:0, C18:0, C18:1, C20:0, C22:0, C24:0, C24:1) across different experimental conditions. The graph includes data for adControl shRNA and adCERS1 shRNA.

(B) Images of human cells stained for MYL1 and DAPI, showing control and adCERS1 shRNA conditions.

(C) Gene expression analysis (fold change, FC) for MYOD, MYOG, MYH7, MYH2, and MYH4 genes under control and adCERS1 shRNA conditions.

(D) Line graph depicting the distance (AU) from control and treated samples (P053 50 uM and P053 100 uM) over different days.

(E) Images of myo3p-GFP expression under control, P053 100uM, and P053 100uM conditions.

(F) Line graph showing the distance (AU) over different days for EV con RNAi and lagr-1 RNAi conditions.

(G) Images of myo3p-GFP expression under EV con RNAi and lagr-1 RNAi conditions.
Figure 5

(A) Common CERS1 variants (AF>1%)

(B) Rare CERS1 variants (AF<1%)

(D) Metabolite pathway

- **Serine + palmitoyl-CoA**
  - SPTLC1-3
  - 3-ketosphinganine
  - CERS1
  - Sphinganine
  - Dihydroceramide
  - Ceramide

- **Beta**
  - 0.5
  - 1.0
  - 2.0

- **Traits**
  - Overall health rating
  - Hand grip strength (right)
  - Hand grip strength (left)

- **r-values**
  - rs117558072: A > C
  - rs1122821: A > G
  - rs71332140: C > T

- **-log10(p)** values

- **Beta** values

- **q-values**
  - 1e-05
  - 1e-04
  - 0.001
  - 0.01
  - 0.1
Supplementary Figure 4

(A) Gene expression (FC) of MyoD, MyoG, Myh7, Myh2, Myh4.

(B) Running time (min).

(C) Running distance (m).

Legend:
- Young
- Aged + DMSO
- Aged + P053

**p < 0.01
*p < 0.05
Supplementary Figure 5

A. Cers1 expression (FC) comparison between Aged AAV con and Aged AAV Cers1KD groups.

B. DhCer levels across different ceramide species (dhCer C16:0, dhCer C18:0, dhCer C20, dhCer C22, dhCer C24:0, dhCer C24:1).

C. Ferret diameter comparison between Aged + AAV con and Aged + AAV Cers1KD groups.

D. CSA (μm²) comparison between Aged + AAV con and Aged + AAV Cers1KD groups.

E. Immunofluorescence images showing CD45 (green), AAV (red), and DAPI (blue) staining.

* indicates p < 0.05, ** indicates p < 0.01, *** indicates p < 0.001.
Supplementary Figure 6

(A) Gene expression (FC) of MyoD, MyoG, Myh7, Myh2, and Myh4 in Aged AAV con and Aged AAV Cers1KD. * indicates significant difference.

(B) Running time (min) for Aged AAV con and Aged AAV Cers1KD. * indicates significant difference.

(C) Running distance (m) for Aged AAV con and Aged AAV Cers1KD. * indicates significant difference.
Supplementary Figure 7

A) Cers1 expression (FC)

B) myotube diameter (µm)

C) myotube area (µm²)

D) Mouse

E) gene expression (FC)

F) multinucleation (%)
Supplementary Figure 8

**A**
- CERS1 expression (FC)
- adControl shRNA
- adCERS1 shRNA

**B**
- nmol/g protein
- dhCer C16:0
- dhCer C18:0
- dhCer C20
- dhCer C22
- dhCer C24:0
- dhCer C24:1

**C**
- myotube diameter (um)
- adControl shRNA
- adCERS1 shRNA

**D**
- myotube area (µm²)
- adControl shRNA
- adCERS1 shRNA

**E**
- multinucleation (%)
- adControl shRNA
- adCERS1 shRNA

* p < 0.05
** p < 0.01
Supplementary Figure 9

A

![Graph A](image)

B

![Graph B](image)

C

![Graph C](image)

D

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Linkage disequilibrium ($r^2$)

E

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- AA (662)
- AC (42)
- CC (2)

rs1122821 (chr19:18940148: A>G)  

- AA (261)
- AG (332)
- GG (113)

rs71332140 (chr19:18886298: C>T)  

- CC (423)
- CT (232)
- TT (51)