RESEARCH ARTICLE

Spastin mutations impair coordination between lipid droplet dispersion and reticulum

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Abstract

Lipid droplets (LD) are affected in multiple human disorders. These highly dynamic organelles are involved in many cellular roles. While their intracellular dispersion is crucial to ensure their function and other organelles-contact, underlying mechanisms are still unclear. Here we show that Spastin, one of the major proteins involved in Hereditary Spastic Paraplegia (HSP), controls LD dispersion. Spastin depletion in zebrafish affects metabolic properties and organelle dynamics. These functions are ensured by a conserved complex set of splice variants. M1 isoforms determine LD dispersion in the cell by orchestrating endoplasmic reticulum (ER) shape along microtubules (MTs). To further impact LD fate, Spastin modulates transcripts levels and subcellular location of other HSP key players, notably Seipin and REEP1. In pathological conditions, mutations in human Spastin M1 disrupt this mechanism and impacts LD network. Spastin depletion influences not only key proteins but also modulates specific neutral lipids and phospholipids, revealing an impact on membrane and organelle components. Altogether our results show that Spastin and its partners converge in a common machinery that coordinates LD dispersion and ER shape along MTs. Any alteration of this system results in HSP clinical features and impacts lipids profile, thus opening new avenues for novel biomarkers of HSP.

Author summary

Hereditary spastic paraplegia is a group of inherited diseases characterized by spasticity and gait disorder. Spastin mutations represent the most common cause of hereditary spastic paraplegia. While Spastin was initially defined as a microtubule severing protein, alternative isoforms impact organelles. In the present work, we identify how Spastin combines the specificity of different splice variants to coordinate endoplasmic reticulum shape and lipid droplet dispersion. When mutated, Spastin impairs this synchronization between organelle dynamics, and affects other proteins involved in hereditary spastic paraplegia.
notably REEP1 and Seipin. The coordination of lipid droplet dispersion and reticulum shaping under the control of Spastin is a unique trafficking model, affected in the different forms of hereditary spastic paraplegia. The impact of Spastin on this machinery results in the alteration of the lipid profile, particularly phospholipids and neutral lipids in muscle and brain. These potential biomarkers open new therapeutic avenues in hereditary spastic paraplegia.

Introduction

Lipid droplets (LDs) have long been considered as inert organelles, limited to fat storage. However, a wide range of evidence recently highlighted the multiple functions of LDs in energy supply, embryogenesis, reactive oxygen species management or pathogen invasion [1–6]. Present in bacteria, yeast, plants and animals, LDs are not limited to adipose tissue in mammals. LDs play determinant functions in many cells, including glial cells, neurons or myocytes [4, 7]. To face cell-specific needs, LDs adapt in size and number through complex machineries [8], comparable to mitochondrial dynamics. The proteins and mechanisms underlying these pathways are partially elucidated [9]. LD biogenesis starts with the accumulation of neutral lipids in the intermembrane space of the endoplasmic reticulum (ER). Among key proteins, Seipin has been shown to control the budding of nascent LD [10–13]. The nuclear membrane recently appeared as a second source of LD [14]. LDs can adapt their caliber through fusion and fission [15], notably via the role of cell death-inducing DFF45-like effector family proteins [16]. Degradation is supported by two complementary mechanisms: enzymatic lipolysis and lipophagy [17].

LDs are well-connected organelles with multiple inter-organelar communications [8, 18]. Beyond the biogenesis process, mature LDs remain in contact with the ER [19, 20], allowing the transfer of lipids and proteins [21]. Contact sites with mitochondria [22, 23] and peroxisomes [24] enhance fatty acid exchanges and modulate metabolic functions. These inter-organelar interactions suggest the existence of a finely tuned trafficking. The first evidence of LD transport comes from the drosophila model and highlights the importance of the gene kIarsicht [25]. Further studies identified Halo as a cofactor of Kinesin-1, controlling LD transport along Microtubules (MTs) in drosophila embryos [26]. However, Halo and Klar have no orthologue in vertebrates. The importance of MTs for LD trafficking has been confirmed in mammalian cells, particularly to adapt to nutritional states [27]. In rat liver cells, Kinesin-1 has been shown to be recruited at the LD surface in fed conditions suggesting the existence of active transport of these organelles [28].

The exact processes and the molecular actors that lead to the distribution and spreading of LDs in vertebrate cells are still unknown. Similarly to Miro-1 and Milton, which are required for mitochondria trafficking along MTs [29], proteins involved in LD transport should exhibit targeting to both MTs and LDs. Until now, none of the LD coating proteins exhibit interactions with the cytoskeleton. The identification of Spastin isoform M1 at the LD surface makes it a great candidate [30]. Spastin is an evolutionary conserved protein that exhibits the unique capacity of severing MTs [31–33]. Mutations of the SPAST gene are responsible for autosomal dominant cases of Hereditary Spastic Paraplegia (HSP); a group of neurodegenerative disorders affecting upper motor neurons [34, 35]. The canonical isoform of Spastin, referred to as M87 in human, exhibits a strong MT-targeting domain in the N-terminus and an AAA ATPase enzymatic function of the C-terminus. The latter participates in the severing activity [32], which has been extensively described in wild-type and mutant conditions [33, 36]. The
existence of an alternative ATG initiator revealed a second transcript encoding for the isoform M1 with a hydrophobic N-terminal domain [37]. This long isoform conserves the MT-targeting domain and strongly binds LD in mammalian cells [30]. Mutations in Spastin isoform M1 have been described to alter axonal transport and intracellular organelles distribution [38–40].

In this report, we propose an extensive description of Spastin splice isoforms, from zebrafish to human. The generation of a CRISPR-Cas9 knockout model supports a role for Spastin in LD dynamics. Spastin deletion affects metabolic properties and organelle characteristics in fish. We further confirm in human cells that Spastin regulates LD formation and dispersion through the reorganization of ER along MTs. Spastin isoforms ensure their function in synergy with other HSP-related proteins such as REEP1 and Seipin. In mutant conditions, Spastin generates a reorganization of ER with a subsequent impact on LDs. Altogether, our results provide evidence that Spastin and other HSP-related proteins synchronize the shaping of ER and MTs stability to determine the dispersion of LDs in the cell, converging in comparable clinical features in case of alteration.

Results

Spastin, a complex set of splice isoforms in zebrafish

Spastin is an ancient protein, putatively present in plants, that exhibits a high conservation of amino-acid identity in mammals [30, 31, 41–43] (Fig 1A). Due to the direct involvement of Spastin (OMIM# 604277) in more than 40% of autosomal dominant HSP (SPG4, OMIM# 182601) [44, 45], its impact on motor neuron and cytoskeleton has been extensively characterized in different animal models [31, 38, 43]. Zebrafish has been shown to be an excellent model to investigate the role of Spastin in motor neuron organization [46, 47]. The conservation of the two initiator sites of the gene has been confirmed in fish [48], suggesting that the different transcripts of Spastin play a key role through evolution. First, to characterize the whole-body expression pattern of Spastin, we performed in situ hybridization at key stages of zebrafish embryogenesis (Fig 1B). Antisense probe staining exhibited an ubiquitous expression of Spastin in early stages of development from 14 to 28 hours post-fertilization (hpf) followed by an accumulation of the transcript in the central nervous system at the end of embryogenesis (72 hpf). In parallel, we used RT-PCR to decipher the existence of different transcripts during zebrafish embryogenesis and in the adult (Fig 1C). This approach revealed a stable expression of the short and long transcripts, respectively M61 and M1, throughout embryonic and larval development. In adult fish, Spastin isoforms were ubiquitous but more abundantly expressed in brain, fat and skeletal muscle (Fig 1C). As described by Jardin et al. [48], a splice variant affecting exon 4 was detectable with the use of specific primers amplifying exon 3 and 5. This splicing event was particularly strong in fat and during the first half of embryogenesis from 6 to 24 hpf (Fig 1C).

Cloning Spastin isoforms from 24 hpf embryos cDNA confirmed the co-existence of four transcripts. All the resulting Spastin proteins conserved the MT interacting and trafficking domain, the MT binding domain and the c-terminal AAA ATPase domain, but differed by the presence of exon 4 and the initiator methionine (Fig 1D). Expression of the different Spastin variants in zebrafish embryonic cells confirmed that short isoforms M61 and M61Δex4 shared a strong severing activity, as overexpression of these isoforms nearly eliminated the MT network compared to un-transfected cells (Fig 1E). The long isoforms M1 and M1Δex4 targeted LDs (Fig 1E), which is explained by a hydrophobic region generated by the alternative initiator codon (Fig 1D), while also partially conserving the severing property.

Further, we found evidence for a third pair of splice variants lacking a large region from exon 2 to 10 (S1A Fig). Detected both in embryos and adult tissues, these splice variants lost
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the severing activity but exhibited a strong location in ER and LDs (S1B and S1C Fig). Although alternative isoforms have been described in human Spastin corresponding to the suppression of exon 8 and 15 [49], this splicing event has not yet been reported in mammals which is why we have decided to focus on the variants presented in Fig 1D.

Taken together, these results highlighted the complex role of Spastin, as a function of its transcripts, and confirmed that the zebrafish is a powerful model to investigate the alternative roles of this protein in the whole organism.

Spastin depletion impacts metabolic properties in knockout animals

Because of the localization of some isoforms to LDs, we next investigated the impact of Spastin on metabolic properties by designing a knockout (KO) zebrafish line based on CRISPR/Cas9 strategy. The gRNA targeted the second exon, immediately after the second initiator codon (M61), resulting in a truncation affecting the four main isoforms (Fig 2A and 2B). In embryos, the lack of Spastin did not affect development or morphology, but altered dechorionation, which was delayed in 30% of the animals at 60 hpf (Fig 2C). Respirometry, performed with the Seahorse XF24 Extracellular Flux Analyzer, revealed a reduction of basal oxygen consumption in 48 hpf KO embryos and a similar tendency in maximal respiration (Fig 2D), indicating reduced mitochondrial respiration. The extracellular acidification rate was also decreased in Spastin depleted embryos compared with wild-type fish (Fig 2E), suggesting a reduced reliance on glycolysis [50]. In adults, lack of Spastin reduced size, weight and BMI in 3 months old KO fish compared with wild-type littermates (Fig 2F–2H). These results highlight the metabolic function of Spastin.

Spastin depletion affects LDs and ER patterning

Given the impact of Spastin on metabolism and the potentially important role of M1 isoforms targeting LDs [30], we continued with the exploration of subcellular components. Here, we incubated embryonic cells from 24 hpf zebrafish embryos for 18h with 300µM of oleic acid (OA) to stimulate the production of LDs. Immunolabeling of Tubulin did not show any differences in MT structure in KO conditions compared with control. BODIPY staining pinpointed dramatic modifications of the LD network in Spa-/- cells (Fig 3A). Lack of Spastin decreased LD size by two-folds and increased their number per cell from 131±13 to 330±34 (Fig 3B and 3C). This increase was confirmed by measuring LD density in 100µm² (Fig 3D). Further, the dispersion from the nucleus center was significantly higher with a mean distance between LD and nucleus of 18.8±0.9µm in wild-type cells and 23.1±0.9µm in KO cells (Fig 3E). In summary, Spastin depletion induced a higher number of LDs, with smaller caliber, that were more disseminated and dispersed through the cytoplasm.

To decipher the impact of LD reorganization in vivo, we followed fat distribution in 6 dpf larvae with Oil Red O staining [51]. At this developmental stage, neither adipose depots nor guts are yet developed. In basal conditions, Spastin KO larvae exhibited lipid accumulation around the swim bladder compared with control animals (Fig 3F). Administration of OA [52] from 5 to 6 dpf triggered an accumulation of lipids in the head, around the yolk and in the...
vessels (Fig 3F) in Spastin KO fish, while control larvae succeeded in buffering the accumulation of fat (Fig 3F).

To determine functional consequences, we measured locomotor performances in 6 dpf larvae (Fig 3G–3J). In basal conditions, wild-type and Spa-/- fish exhibited similar swimming capacities. While OA administration increased the distance swam by control fish from 1206 ±76 to 1873±184mm in 30min, Spastin KO larvae presented a significant reduction from 1324 ±119 to 799±224mm (Fig 3H). OA treatment also reduced net velocity and increased pausing in Spa-/- animals (Fig 3I and 3J). Given that the impact of OA on Spastin mutant could be partially mediated by ER stress [52, 53], we evaluated the expression of X-box Binding Protein 1 (XBP1) by RT-PCR [54]. Indeed, the presence of spliced XBP1 (sXBP1), in addition to the canonical form (uXBP1), is an indicator of ER stress as shown in 6 hpf embryos [54], here...
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A. Bodipy staining showing lipid droplets in Ctrl and Spa/- cells. 50μm and 30μm images are shown.

B. LD Area (μm²) comparison between Ctrl and Spa/-.

C. LD number/cell comparison between Ctrl and Spa/-.

D. LD number/100μm² comparison between Ctrl and Spa/-.

E. Distance LD-nucleus comparison between Ctrl and Spa/-.

F. 6 dpf comparison between Ctrl and Spa/-, and 6 dpf + OA.

G. Diagram showing Ctrl, Spa, + TM, and + OA marked with circles.

H. Graph showing Distance (mm) between Ctrl, Spa, Ctrl + OA, Spa + OA, Ctrl + TM, and Spa + TM.

I. Graph showing Net velocity (mm/s) comparison between Ctrl, Spa, Ctrl + OA, Spa + OA, Ctrl + TM, and Spa + TM.

J. Pausing time (%) comparison between Ctrl, Spa, Ctrl + OA, Spa + OA, Ctrl + TM, and Spa + TM.

K. 6 hpf and 6 dpf comparison of uXBP1 and sXBP1 with GAPDH.

L. Diagram showing slow and fast fibers.

M. Images showing slow fiber and fast fiber differences between Ctrl and Spa/-.
used as positive controls (Fig 3K). In Spastin KO larvae, the detection of sXBP1 indicated the activation of ER stress response (Fig 3K), which was not present in wild-type controls. We further confirmed the higher sensitivity to ER stress of Spastin KO treating the larvae for 24h with 2µg/ml of Tunicamycin (TM). TM administration did not affect the distance swam in control larvae but impacted Spastin KO which exhibited a large reduction of mean distance and net velocity (Fig 3G and 3I) with a reciprocal higher pausing time (Fig 3J). This confirms that Spastin KO larvae are more sensitive to ER stress than controls.

This impact of Spastin on lipid management and ER stress at the larval stage called for the investigation of organelles in adult fish with a particular interest in skeletal muscle. Indeed, the role of Spastin in non-neuronal tissues remains poorly characterized, despite its ubiquitous pattern of expression (Fig 1C) and the previous description of Spastin effect on human skeletal muscle transcriptome [55]. We used electron microscopy to decipher the role of Spastin on zebrafish skeletal muscle ultrastructure (Fig 3L). In control animals, slow muscle was enriched in mitochondria around fibers and between sarcomeres (i.e. the contractile units), while fast muscle mitochondria were mostly at the periphery (Fig 3L). Lack of Spastin did not affect mitochondria appearance (Fig 3M). In wild-type slow and fast muscle, sarcomeres were surrounded by budding ER (Fig 3M). Lack of Spastin completely flattened ER in slow fibers and accumulated ER in hives at the edges of fast fibers (Fig 3M). These observations confirmed the importance of Spastin in muscle and in organelle organization. Taken together, these experiments, performed in the zebrafish model, confirmed the important impact of Spastin on LDs and ER in different tissues and cell types.

Human Spastin M1 isoforms conserve features of the zebrafish counterparts

To emphasize the observations obtained in zebrafish, the impact of human Spastin isoforms on organelles had to be investigated. Analysis of cDNA from human muscle confirmed the coexistence of the two initiator codons and the splice modifications that suppressed exon 4 as described in zebrafish (Fig 4A). The absence of exon 4 was systematically accompanied by the lack of Alanine 139 (A139), which is encoded by the overlapping sequence formed by exon 1 and 2 (Fig 4A). The lack of exon 4 abolished the prediction of a strong coiled-coil domain in human Spastin (Fig 4B). The removal of A139 residue had a minor impact on this secondary structure (Fig 4B). Given these observations, we pursued the investigations on the full-length isoforms M1 and M87, as well as their counterpart missing both exon4 and A139 (hereafter referred as to M1Δex4 and M87Δex4).

In HeLa cells treated with 300µM of OA, M87 and M87Δex4 exhibited a diffuse location and triggered the complete severing of acetylated Tubulin in all the transfected cells, comparable to a 12h treatment with nocodazole. M1 and M1Δex4 presented an exclusive targeting to LDs and partially conserved the severing activity (Fig 4C). These results pointed out strong and evolutionarily-conserved functions of the different isoforms of Spastin.
Spastin and lipid droplets

A

Spastin M1

Spastin M87

Hydrophobic Region
Exon 4
AAA ATPase Domain
Microtubule Interacting and Trafficking Domain
Microtubule Binding Domain

B

Spastin M1 (616aa)
Spastin M1ΔA139 (615aa)
Spastin M1Δex4 (584aa)
Spastin M1Δex4 ΔA139 (583aa)

C

+ OA

| Ø    | Noco | Taxol | Spastin M87 | Spastin M87Δex4 | Spastin M1 | Spastin M1Δex4 |
|------|------|------|-------------|-----------------|-------------|----------------|

Spastin

Ac Tub

Bodipy

Merge

10μm
To decipher the putative involvement of the long form M1 and M1Δex4 in HSP, we reproduced the mutation C488Y in our constructs, as well as a new modification that we identified in human cDNA H455R. In HeLa cells, both mutations converged in strong effects targeting both LDs and MTs, similarly for M1 and M1Δex4 (S2A Fig). LDs were accumulated around the nucleus. In presence of mutant constructs, acetylated Tubulin formed large concentric bundles. Mutant Spastin presented a specific pattern at the cortical extremity of MTs, forming an inverse gradient of intensities with acetylated Tubulin (S2A Fig), i.e. when one fluorophore was intense the other was moderate and vice-versa.

In challenged conditions, mutant M1H455R was accumulated on MTs around the nucleus in presence of Taxol. Nocodazole led to a strict relocation of Spastin on LDs. M1H455R maintained MTs-targeting despite a short cold exposure that suppressed the acetylated Tubulin signal [56] (S2B Fig). In summary, mutations in Spastin long isoforms converged on a common gain of function that induced bundled MTs, equivalent to an abnormal stabilization, and modifications of the LD network.

**Spastin isoforms modulate organelles contact sites**

To explore the fate of other organelles with a quantitative approach, we used multispectral analysis in transfected U2OS cells [20] (Fig 5A–5M). Overexpression of Spastin isoforms increased LD size (Fig 5C and 5D). M1 and M1Δex4 decreased peroxisome number (Fig 5E) but did not modify their size (Fig 5F). Lysosome number and size were not affected (Fig 5A and 5B). ER and mitochondria area fraction remained stable (Fig 5G and 5H) even when ER morphology was altered (Fig 5K–5M).

In addition to these modifications of individual organelles, Spastin isoforms modulated contact sites. Wild-type M1 reduced the contact between ER/mitochondria and ER/peroxisome (Fig 5I). Similarly, M1 decreased contact between LD/peroxisome and LD/ER (Fig 5J). In the absence of exon 4 or the presence of the mutation H455R, the impact of Spastin M1 on organelle contacts was not observed (Fig 5I and 5J).

To complete this approach, we used a quantitative analysis in HeLa cells treated with OA to enhance LD biogenesis. Expression of wild-type M1 and M1Δex4 increased LD size (Fig 6A–6C). In a pathological context, the presence of H455R mutation decreased LD number and increased LD size. These observations were coherent with the effect of Spastin KO described in zebrafish cells (Fig 3A–3E). M87 and M87Δex4 did not impact LDs features, similarly to the administration of nocodazole, suggesting that the effect of Spastin could not be attributed to the severing activity on MTs (Fig 6A–6C). Stabilization of MTs by Taxol administration reduced LD number without changing their size, confirming that Spastin effect was not limited to Tubulin modulations (Fig 6A–6C).

In addition to the modification of LD size and number, Spastin long isoforms triggered a redistribution of LDs that accumulated around the nucleus (Fig 6D). High magnification confirmed a close contact between Spastin rings and Lamin (Fig 6A). Expression of M1 and M1Δex4 increased the contact between LDs and nucleus membrane, but also enhanced the accumulation of intranuclear LDs that represented respectively 5.6±1.6 and 6.2±2.1% of the
total pool compared to 2.6±0.6% in control conditions (Fig 6D). Mutations of Spastin M1 massively increased intranuclear LDs to 18.8±4.7%. The impact of Spastin on nuclear LDs was in line with a former study reporting an effect of the protein on nuclear membrane [57].
Spastin impacts LDs through effects on ER

The impact of Spastin on the LD network cannot be explained by MTs modulation but could be due to an alteration of biogenesis mechanisms. It is now well established that nascent LDs appear through neutral lipid accumulation within the ER in a complex budding process mediated by key players such as Seipin [11, 12]. The modulation of this machinery is still poorly characterized. To follow up on the multispectral analysis pointing to a decreased contact between ER/LD (Fig 5J), we performed a complementary staining of Spastin, Calreticulin and acetylated Tubulin in HeLa cells (Fig 6E and 6F). While wild-type Spastin M1 and M1Δex4 resulted in a retraction of ER around the nucleus, mutant counterparts broadly reorganized the network in filamentous structures (Fig 6E). Co-staining of acetylated Tubulin confirmed that Calcineurin labeling colocalized with mutated Spastin along the cortical extremity of MTs (Fig 6F). Our observations are in line with previous works showing that Spastin M1 isoform can modulate ER shaping through the anchoring of its hydrophobic N-terminus [58, 59]. Here, we propose that the retraction or elongation of ER along MTs determine the position of LD biogenesis in the cell (Fig 6G). As the majority of LDs remain attached to the ER [20], the impact of Spastin on ER spreading may directly define LD dispersion. In wild-type context, long Spastin isoforms modulated ER shaping by its retraction around the nucleus, explaining the accumulation of fewer but bigger LDs, further participating in the presence of perinuclear LDs. Mutations spread ER along MTs to form excessive tubular shapes, resulting in the same LDs pattern and size than wild-type Spastin (Fig 6G).

Spastin modulates other players involved in HSP at both transcript and protein levels

ER shaping is known to be modulated by other proteins involved in HSP such as Atlastin1 and REEP1 [60–63]. Our results reinforce the hypothesis of a global function shared by HSP-associated proteins to coordinate ER shaping and MTs dynamics [64]. To determine the potential connection in a common pathway, we investigated the impact of Spastin depletion on other genes transcripts. We chose to focus on four HSP-associated proteins: Seipin (OMIM# 606158, involved in SPG17 OMIM# 270658) and Spartin (OMIM# 607111, involved in SPG20 OMIM# 275900) given their role in LD fate [65], as well as Atlastin1 (OMIM# 606439, involved in SPG3A OMIM# 182600) and REEP1 (OMIM# 609139, involved in SPG31 OMIM# 610250) given that both had been shown to impact LD size [64, 66, 67]. Thus, we followed the expression of Seipin, Spartin, Atlastin1 and REEP1 (also referred as to SPG17, SPG20, SPG3A and SPG31 respectively) in Spastin KO zebrafish (Fig 7A and 7B). In brain, REEP1 and Atlastin1 presented a twofold increase in absence of Spastin. Seipin and Spartin exhibited a similar trend (Fig 7A). In skeletal muscle, Spastin depletion induced a threefold increase of Spartin and Atlastin1. Seipin and REEP1 transcripts appeared respectively 7 and 4
times more present in Spa-/- than in controls (Fig 7B). These results point to a transcriptional compensation of Seipin, Spartin, Atlastin1 and REEP1 to face Spastin depletion, thus providing evidence for a genetic link between Spastin and other HSP-related genes.

In addition to the transcriptomic dialogue between HSP genes, we examined the influence of Spastin on these candidates at the protein level defining their localization in HeLa cells. Wild-type Spastin M1 and M1Δex4 modulated Seipin through ER shaping, decreasing its location around LDs. Conversely, mutated Spastin proteins spread Seipin along MTs to the cell periphery (Fig 7C and 7D). This effect was specific to Spastin mutations, as MT-modifying drugs were unable to relocate Seipin (Fig 7E). The impact of Spastin was also observable for a longer splice variant of Seipin [11] (S3A and S3B Fig). The relocation of Seipin in presence of wild-type or mutant Spastin confirmed that the reticulum reorganization affects the machinery responsible of LD biogenesis, explaining the dispersion or the retraction around the nucleus.

Spastin also affected other key players of the ER/LD dialogue. Expression of wild-type Spastin M1 or M1Δex4 triggered the relocation of REEP1 (Fig 7F) depending of the presence or absence of OA. In the absence of OA, REEP1 relocated along the ER, thus around the nucleus (Fig 7F). Expression of mutant Spastin M1H455R and M1Δex4H455R redistributed REEP1 along thick tubules of ER (Fig 7F). Remarkably, in HeLa cells, REEP1 property to shape tubular ER was very weak, especially compared with mutant Spastin. OA administration totally relocated the protein around giant LDs close to the nucleus (Fig 7G) revealing an unexpected LD-targeting of REEP1, which also enhanced the volume of these organelles (S3C Fig). This property was not dependent on Spastin, as REEP1 still targeted LDs in zebrafish embryonic cell depleted for Spastin (S3D Fig). In presence of OA, wild-type Spastin M1 and REEP1 colocalized on LD surface, with REEP1 exhibiting an exclusive pattern at the inter-LD contact sites. Mutated Spastin M1H455R (Fig 7G) was excluded from the LD surface in presence of REEP1 (Fig 7G).

Wild-type and mutant Spastin M1 relocated Atlastin1 as a function of ER shaping (S4A Fig). Spartin expression generated clusters of LDs in which wild-type and mutant Spastin M1 were localized (S4B Fig). Altogether these elements point to a strong interdependence between HSP-associated proteins. However, Spastin represents the core of this machinery and plays a crucial role in ER shaping and subsequent LD location.

**Spastin depletion triggers alterations in the lipid profile**

The effect of Spastin on LDs and ER suggested a potential impact on neutral lipids or membrane components. Using state-of-the-art mass spectrometry, we analyzed lipidomics in brain and skeletal muscle from wild-type or Spastin KO zebrafish. Quantification of non-esterified cholesterols (Chol), esterified cholesterols (Est Chol) and triacylglycerides (TG) did not reveal any alterations of neutral lipids in Spa-/- brain (Fig 8A and S1 Table). In skeletal muscle, Spastin KO induced reductions of Chol, Est Chol and total TG (Fig 8B and S1 Table). Analysis of saturated, mono- and poly-unsaturated fatty acids (respectively SAFA, MUFA and PUFA) did not indicate any variations between control and Spa-/- brain but were affected in skeletal muscle (S1 Table). Among the different molecular species, only fatty acids composed of 16 carbons...
Fig 8. Spastin depletion modifies lipid profile in adult zebrafish brain and skeletal muscle. (A-B) Unesterified cholesterol (Chol), total esterified cholesterol (Est Chol) and total triacylglycerides (Tot TG) in brain (A) and in muscle (B). (C) Fatty acids composed of C16 in muscle. (D-E) Ceramides (Cer) in brain (D) and muscle (E). (F) Phosphatidylethanoamines (PE) in brain. All values correspond to lipid quantity per total protein amount. Bars are mean ± SEM (n = 3 per group), *P<0.05, **P<0.01, ***P<0.001 (unpaired t-test between Ctrl and Spa-/-). See also S1 Table.

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were reduced in the absence of Spastin (Fig 8C). Altogether, these results highlighted the impact of Spastin on neutral lipids and associated fatty acids in skeletal muscle. The broad reduction of TG and Est Chol may be associated with LDs alteration and may represent a new hallmark of Spastin depletion.

Due to the involvement of Spastin in organelle dynamics and contacts, we also investigated its impact on structural lipids. Analysis of phospholipids demonstrated that phosphatidylcholines (PC) were not affected in Spastin KO fish (S1 Table). Sphingomyelins (SM) and Phosphatidylinositol (PI) exhibited few modifications in some specific species. SM d18:1/16:0 presented a twofold reduction in Spastin KO skeletal muscle, while PI 38:2 increased in KO brain (S1 Table). Beside these specific modifications, Ceramides (Cer) were broadly affected by Spastin depletion with a reduction of Cer d18:1/18:0 in brain (Fig 8D) and Cer d18:1/22:0, Cer d18:1/26:0 and Cer d18:1/26:1 in muscle (Fig 8E). Phosphatidylethanolamines (PE) also revealed dramatic alterations in Spastin KO fish, specifically in the nervous system. PE 32:0, PE 34:0, PE 34:1, PE 36:1, PE 36:2, PE 40:3 and PE 40:7 followed a comparable increase in Spa-/- brain (Fig 8F and S1 Table). Altogether, these finding confirmed the impact of Spastin on structural lipids with organ specificities. Interestingly, Spastin affected particularly PE and Cer, two major components of organelle membranes known to impact membrane fluidity and folding, notably in the ER [68, 69].

Discussion

Spastin isoforms coordinate cytoskeleton and organelles

While the role and location of a protein are historically characterized from its canonical isoform, recent advances reveal that alternative splice variants can represent a source of function diversity [70]. In pathological conditions, splice isoforms are associated to specific disorders as described for Tau [71]. The co-existence of different Spastin variants has been described from worm to human [43]. In human, former studies mainly focused on the short M87 protein, known to carry the severing property [32, 33, 72]. However, recent advances highlighted that the M1 long isoform harbors a LD targeting sequence [30]. In addition to the existence of the two start codons, an alternative splicing of exon 4 is also conserved from fish [48] (Fig 1C–1E) suggesting a specific function for this variation. The presence of exon 4 represents a strong coiled-coil domain (Fig 4B) and has been hypothesized to modify Spastin functions [73]. Indeed, while M1 and M1Δex4 share similar effect on peroxisome number and LDs size (Fig 5C–5F and 6B–6D), M1 exhibits a unique propensity to reduce organelle contact area (Fig 5I and 5J).

Beyond these major isoforms, the existence of uncharacterized splice events present in low concentration may confer new levels of complexity. In zebrafish, a new splice variant of M1, deleted from exon 2 to 10, only conserves LD targeting properties (S1 Fig) and may play a role of competitor for the full-length M1. Further investigations should confirm the existence of this variant in mammals. Conversely, low abundance isoforms lacking exon 8 and 15 have been described in human, but their functions remain unknown [74]. The alteration of AAA domains may confer to these natural variants a MT-bundling function similar to mutant Spastin. These results highlight the complexity of Spastin variants.

Spastin plays an important role in muscle

While different papers claimed a specific expression of Spastin, notably M1 isoform, in the central nervous system [75], we provide evidence of the ubiquitous pattern of this protein. Spastin depletion affects respiration, locomotion and body measurements. At the cellular level, KO fish reveal an impairment of sarcoplasmic reticulum (Fig 3M) and confirm an impact of
Spastin on skeletal muscle [55]. Lipid profile analyses highlight a reduction of neutral lipids in muscle, while the nervous system is not affected. C16 fatty acids and associated TG are specifically altered in skeletal muscle and may represent a new biomarker in HSP patients. Altogether these observations indicate the involvement of Spastin in muscle. The action of Spastin and other HSP genes on organelles may affect tissue metabolism and participate in the muscular symptoms described in patients [35].

**Spastin coordinates MTs and ER shaping to modulate LDs dispersion**

Spastin isoforms reveal a global impact on organelles (Figs 5 and 6) that cannot be limited to their action on MTs dynamics. Spastin depletion alters ER morphology in fish skeletal muscle while overexpression of M1 and M1Δex4 condensate ER around the nucleus in mammalian cells. A direct role of Spastin on ER shaping has been recently evidenced [58, 59]. Here we confirm this function in vivo (Fig 3M) and implicate Spastin deletion in causing ER stress (Fig 3F–3K).

The reorganization of the ER network affects contacts and dynamics of other organelles, mainly LDs. In KO zebrafish cells, lack of Spastin enhances the dispersion of small LDs (Fig 3A and 3B), while overexpression of M1 isoforms in HeLa cells promote LD accumulation around the nucleus (Figs 4C, S2A and 6A–6D). Spastin appears as an inhibitor of LDs dispersion. Although MTs have been shown to impact LD location in the cell [27], Taxol and nocodazole administration do not mimic Spastin effects (Fig 6B–6D) confirming that the impact of the protein on LD dispersion is not mediated by its action on cytoskeleton. As the large majority of LDs remain in contact with ER after biogenesis [20], the reorganization of reticulum along MTs network subsequently determines the dispersion of LD. We here propose the concept that by controlling the spreading of ER, Spastin confers to LDs a specific trafficking process, different from other organelles that move directly along MTs tracks. LDs spread in the cell in function of ER/MTs dynamics (Fig 6G). Given that the elongation of ER along MTs represents an evolutionary conserved mechanism that has been described in plants [76], the subsequent spreading of LDs is probably an ancestral process that should be further investigated in eukaryotes. In mammals, this dispersion may be crucial to deliver LDs to the extremity of long neurons such as upper motor neurons. However, LDs also play key functions in other cell types [1, 2, 5, 9, 77] explaining the impact of Spastin on many tissues.

Structurally, modulations of ER shape and LD dynamics result in organ-specific alterations of the lipid profile. Spastin impacts not only LD number but also affects their composition (Fig 8). Structural lipids are also modified, notably Cer and PE. These phospholipids, which are particularly important to define membrane fluidity, may participate in ER shaping and organelle contact. PEs are also chaperone lipids that ensure protein folding and participate in ER stress [68, 69]. These findings show that Spastin functions are linked with the modulation of the lipid profile in neuronal and muscle tissues.

**Mutations in Spastin massively affect organelles through M1 isoforms gain of function**

HSP-related mutations were reported in the different domains of Spastin [34]. These mutations lead to the loss of M87 function, which represented a discrepancy with the dominant transmission of the disease, did not affected only the AAA domain and did not explain HSP pathological features [78]. Other studies highlight the importance of M1 isoforms in the disease onset [40] with mutations conferring to this isoform a bundling activity that stabilizes MTs [79]. Here, we reinforce the hypothesis that long Spastin isoforms exhibit a gain of function and may play a central part in HSP physiopathology [78, 79]. In mutant conditions, M1
and M1Δex4 converge on an abnormal stabilization of MTs (S2 Fig). This propensity to bundle MTs may co-exist with the severing activity in the wild-type protein [73], but becomes dominant in mutants.

Spastin M1 gain of function is not only associated with cytoskeleton stability. Mutations in Spastin long isoforms also impacted LD dispersion and morphology (Fig 6A–6D). The abnormal formation of tubular ER [59] (Fig 6E–6G) and the relocation of other proteins involved in LD biogenesis (Fig 7D) resulted in the reorganization of organelle networks, notably LDs. Surprisingly, excessive ER spreading and retraction converged on similar consequences for LDs.

Mutated M1 also increased the accumulation of LD around the nucleus (Fig 6A and 6D). The alteration of ER shaping may enhance the import of LDs inside the nucleus or promote biogenesis directly inside the nuclear membrane. Former studies highlighted an action of Spastin on nuclear envelope integrity [57] that may participate in LD accumulation in the nucleus. Regarding the numerous functions of LDs in the nucleus [80], it could be hypothesized that their abnormal accumulation may play a role in HSP pathophysiology.

Spastin is involved in a multi-protein system controlling organelle fate

Based on multiple domains and isoforms, Spastin coordinates MTs, ER and LDs in basal and pathological conditions. Strikingly, proteins sharing similar functions are all involved in the same group of diseases. Spastin influences HSP-related proteins at the transcriptomic level (Fig 7A and 7B) and determines their relocation in the cell (Fig 7C–7G).

REEP1 [62–64] and Atlastin1 [61, 81] are known to influence ER shape by creating tubular structures, similarly to mutant Spastin M1. By controlling the location and the transcription of these HSP-related proteins (Figs 7 and S4A), Spastin coordinates ER shape. The convergent functions between Spastin and REEP members are probably crucial for neuronal and muscle tissues. In cardiac muscle from rat, inactivation of REEP5 (OMIM# 125265) induces an alteration of sarcoplasmic reticulum morphology [82], comparable to the effect of Spastin deletion in fish muscle (Fig 3M). Among other Spastin partners, Seipin may represent the main actor to control LD dynamics. Transcriptional modulation of Seipin (Fig 7A and 7B) and relocation in function of ER state (Figs7C–7E and S3A and S3B) affect LD biogenesis and subsequent dispersion through the cell.

Spastin uses MT tracks to remodel the ER and this shaping defines LD dispersion. While the impact of Spastin on MTs is well described, other HSP-related proteins also revealed an action on cytoskeleton. In drosophila, Atlastin and Spartin have been shown to modulate MTs stability [83]. Similarly to Spastin, Kinesins KIF1A (OMIM# 601255), another protein involved in HSP [84], accumulates on specific MTs regions when mutated in its ATPase domain [85]. Interestingly, Kinesins have been shown to modulate reticulum spreading along MTs [86, 87]. Their effect on LD transport notably described in drosophila may be due to an indirect action on ER shaping [26].

Beside its impact on ER shaping, Spastin controls HSP-related partners directly involved in LD dynamics. Spastin follows the clustering of LDs in presence of Spartin (S4B Fig), a protein initially described at the mitochondria level [88] but also reported as a LD modulator [65]. Here, we also describe a strong targeting of REEP1 to LDs, increasing their size (Figs 7G and S3C) in presence of fatty acid stimulation. While REEP1 depletion has been previously described to modulate LD size and to be associated with lipodystrophy in mouse model [89], this report provides evidence for the first time of a direct targeting of the wild-type protein to LDs. The access of REEP1 on LDs probably depends on fatty acids concentration, as other studies described the protein on LDs only in mutant context [66]. Wild-type and mutant Spastin triggers subtle modification in REEP1 location.
Further investigations are needed to explore the participation in this dialogue between ER and LDs of other HSP-related proteins known to impact ER shaping [45], such as RTN2 (OMIM# 603183, involved in SPG12 OMIM# 604805) [90], Protrudin (OMIM# 610243, involved in SPG33 OMIM# 610244) [91] or ARL16IP (OMIM# 607669, involved in SPG61 OMIM# 615685) [92]. Indeed, we can hypothesize that by modulating ER shape, these proteins may impact LD dispersion. Furthermore, KIF1A may also play a role in this machinery, potentially representing the engine that actively spreads ER along microtubules as observed in plant and mammalian cells, with a subsequent reorganization of the LD network [76, 86].

Altogether these HSP-related proteins constitute a scaffold ensuring membrane shaping and organelle dialogues. Their numerous functions are supported by splice-mediated complexity, as illustrated by Spastin or Paraplegin variants [93]. This machinery converges to control LD dynamics. At the crossroad of MTs, ER and LDs, Spastin appears as a key player of this machinery and illustrates the functional variability carried by splice isoforms.

Conclusion

In summary, the characterization of Spastin isoforms reveals the complex functions of this multi-faceted protein. Coordinating MTs and ER dynamics, Spastin modulates the LD network and dispersion in the cell. These crucial functions are based on key partners similarly involved in the physiopathology of HSP. In the light of this complex machinery, HSP appears as the first group of diseases directly due to a disorganization of organelles dynamics. A major consequence is the modification of the lipid profile in absence of Spastin. These alterations exposed in the zebrafish model suggest that lipidomic analyses may supply new biomarkers for HSP patients and open new therapeutic targets.

Materials and methods

Ethics statement

This study protocol was approved by the cantonal veterinary authorities (approval number VD3391).

Zebrafish Husbandry and strains

Zebrafish (Danio rerio, Oregon AB) were housed at the Zebrafish facility of the School of Biology and Medicine, maintained at 28.5°C and on a 14:10h light:dark cycle. Embryos were staged by hours (h) or days (d) post fertilization according to Kimmel et al. [94].

Generation of CRISPR/Cas9 knockout

The guide RNA targeting zebrafish Spastin (Q6NW58) was designed on https://chopchop.cbu.uib.no. The target sequence was selected in exon 2 with a predicted efficiency of 0.74: TGTAGGTGATAAGCAGAAGGCGG. gRNA was produced and purified with Gene Art precision gRNA synthesis kit (A29377, Invitrogen, ThermoFischer Scientific, Waltham, MA, United States). gRNA and recombinant Cas9 nuclease (B25641, Invitrogen) were mixed at respectively 100ng/µl and 500pg/nl in a 200mM solution of KCl. 1nl of the mix was injected in the first cell of AB embryos.

F1 heterozygous animals were identified after fin clipping and PCR amplification of Spastin fragment using the following primer: For 5-ACCTGTGATGTGAACGTTTCT-3; Rev 5-TTGTAATAAGCCGTGTAATAA-3. The sequence modification was visualized on Bis-acrylamide gels. The exact nature of the CRISPR/Cas9-induced mutation was confirmed by Sanger sequencing on fish from the F2 generation. Among different lines deleting the Spastin
gene, fish exhibiting an insertion of 7 nucleotides at the beginning of exon 2 were used for the present study.

**Locomotion assays on zebrafish larvae**

Zebrafish spontaneous activity was monitored at 6 dpf using a Zebrabox recording system (Viewpoint, Lyon, France). One fish was placed per well on a 96 well plate. Recording was performed during 30 min in the dark. Velocity thresholds were set to distinguish between slow and fast velocity (respectively v in green, and V in red) as follow: $3 < v < 6 \text{mm/s}$ and $V > 6 \text{mm/s}$ [95]. Net velocity was defined as the global larval speed, excluding pausing time. 2µg/ml of Tunicamycin (T7765, Sigma-Aldrich, St. Louis, MO, United States) and 0.2mM of OA (O1383-1G, Sigma Aldrich) [52] were added to the fish water from 5 to 6 dpf and washed prior to the locomotion test.

**RNA extraction and qPCR**

Total RNA extraction, RT-PCR and RT-qPCR were performed as previously described [54, 96]. All protocols are detailed in the supporting information in S1 Text. All primers are listed in S2 Table.

**In situ hybridization**

Probe templates corresponding to the 500 first nucleotides of Spastin M1 (Q6NW58) were obtained by PCR (see primers in S2 Table) and amplicons were flanked with T7 promoter. Sense and anti-sense probes were synthetized by reverse transcription (mMESSAGE mMACHINE T7 Transcription Kit, AM1344, Invitrogen) and purified on Microspin G50 columns (27533001, GE Healthcare, Chicago, IL, United States). In situ hybridizations were performed as previously described [95].

**Cloning**

CDS were cloned from cDNA obtained from human muscle, brain, or 24 hpf zebrafish embryos and amplified by PCR with ATTB flanking sequences (see primers in S2 Table) to use in the Gateway cloning system. PCR products were purified on agarose gels (MinElute Gel Extraction, 28604, Qiagen, Hilden, Germany) and inserted in Gateways pDon221 vector by recombination using BP clonase II enzyme mix (11789, Invitrogen). For expression in eukaryote cells, CDS were transferred in destination vectors pcDNA-CMV-CherryN or pCI-CMV-3XflagN with LR clonase II enzyme mix (11791, Invitrogen).

**Embryonic cells culture**

24 hpf embryos were anesthetized in Tricaine (E10521MS-222, Sigma-Aldrich) and digested in trypsin-EDTA 0.05% (25300, GIBCO, ThermoFischer Scientific). Cell culture was processed at 28°C in L15 complete culture medium (11415–049, GIBCO) as described in Vallone et al. [97]. When indicated, 300µM of OA (O1383-1G, Sigma-Aldrich) was added in L15 Medium/2% BSA for 18h prior to fixation.

**Cells transfection and Immunolabeling**

HeLa cells were grown in DMEM1X GlutaMAX medium (31966–021, GIBCO) supplemented with 10% FBS (10270–016, GIBCO) and penicillin/streptomycin. Transfection was performed with Lipofectamine 3000 (L3000-01, Invitrogen) following supplier instructions. Immunostaining procedure was performed 48h later. When indicated, 300µM of OA were added in
DMEM/2% BSA for 18h before the immunostaining protocol. Similarly, 1µM of nocodazole (M1404, Sigma-Aldrich) and 20µM of Taxol (Paclitaxel, P3456, Invitrogen) were added for 12h prior to the immunolabeling protocol. Cells were fixed 10min in PBS1X/PFA4% then permeabilized in PBX1X/Triton 0.25% for 5min. Primary antibodies were applied in 1% BSA blocking solution for 90min at room temperature. All antibodies and concentrations are listed in the supporting information in S1 Text.

Confocal imaging and quantification

Image acquisition was performed on a confocal laser-scanning microscope (LSM510, Zeiss, Stockholm, Sweden) at the Cell Imaging Facility of the School of Biology and Medicine. Quantification of LDs number and size was performed with ImageJ (NIH, Bethesda, MA, United States), using the Analyze Particles function. For the dispersion analysis, Cartesian coordinates from individual LD (X_{LD}, Y_{LD}) and from nucleus center (X_{Nu}, Y_{Nu}) were determined with ImageJ. Distances (d) were calculated using the formula \( d = \sqrt{(X_{Nu} - X_{LD})^2 + (Y_{Nu} - Y_{LD})^2} \).

Electron microscopy

Electron microscopy was performed at the Electron Microscopy Facility of the School of Biology and Medicine. Zebrafish skeletal muscle were dissected immediately after euthanasia and pre-fixed in PB 0.1M/2.5% glutaraldehyde/PFA4% solution at room temperature for 30min. 1mm cross-sectional sections were cut with a razor blade and fixation was prolonged until 2h. The samples were washed in PB three times (5min distilled water) and dehydrated in acetone solution (Sigma). Careful positioning was performed with binoculars on Aclar film (EMS, Hatfield, PA, United States) inside a Gene Frame (ThermoFisher Scientific) in 65µl of Epon resin and finally polymerized for 48h at 60˚C in oven. Sagittal ultrathin sections of 50nm were cut on a Leica Ultracut (Leica Mikrosysteme GmbH, Vienna, Austria) and picked up on a copper slot grid 2x1mm (EMS) coated with a polystyrene film (Sigma). Sections were post stained with 4% uranyl acetate (Sigma). Micrographs were taken with a transmission electron microscope Philips CM100 (ThermoFisher Scientific) with a TVIPS TemCam-F416 digital camera (TVIPS GmbH, Gauting, Germany).

Seahorse analyzer respirometry

Mitochondrial function was determined with an XFe24 extracellular flux analyzer (Seahorse Bioscience, Billerica, MA, United States). Oxygen consumption rate (OCR) and extracellular acidification rate (ECAR) were measured in dechorionated embryos at 48 hpf. Embryos were staged and placed one per well on an islet capture microplate filled with E3 egg water. The plate was incubated in an incubator without CO2 at 28˚C for 30min. After measuring baseline OCR as an indication for basal respiration, OCR was measured after an injection of 2µM of FCCP to determine maximal respiration.

Multi spectral analysis

U2OS cells were prepared for multispectral imaging as previously described [20, 98]. Details are available in the supporting information in S1 Text.

Lipidomics

Seven months old male zebrafish were anesthetized on ice and killed by decapitation. Brain and skeletal muscle were immediately dissected and flash frozen in liquid nitrogen. Extended
protocols for lipid extraction and analyses are described in the supporting information in S1 Text.

Supporting information

S1 Fig. Identification of an uncharacterized splice variant in zebrafish Spastin. (A) Schematic representation of alternative splice suppressing exon 2 to 10 in zebrafish Spastin. (B-C) Confocal microscopy images of zebrafish embryonic cells (B) and HeLa cells (C) overexpressing Spastin splice variant (treated with 300µM oleic acid for 18h before acquisition). Cherry-tagged Spastin appears in red. Tubulin labeling corresponds to microtubules (cyan), Bodipy to LDs (yellow) and Hoechst to nucleus (blue).

S2 Fig. Mutations in Spastin M1 and M1Δex4 converge on similar gain of function. (A-B) Confocal microscopy images of HeLa cells overexpressing Spastin splice variants treated with OA for 18h. Cherry-tagged Spastin appears in red. Acetylated Tubulin labeling corresponds to microtubules (cyan), Bodipy to LDs (yellow) and Hoechst to nucleus (blue). In (B) cells were also submitted to 20µM Taxol, 1µM nocodazole (Noco) treatment for 12h, or to cold exposure for 15 min before staining protocol.

S3 Fig. Seipin and REEP1 participate to ER/LD dynamics. (A) Schematic representation of human Seipin variant α and β. (B) Confocal microscopy pictures of HeLa cells overexpressing human Seipin α and β with Spastin M1 isoforms (after 18h administration of OA). Cherry-tagged Spastin appears in red, Seipin α and β in green, LDs (Bodipy) in magenta and nucleus (Hoechst) in blue. (C) Confocal microscopy images of HeLa cells overexpressing human REEP1 after 18h administration of OA. REEP1 appears in green, LDs (Bodipy) in magenta and nucleus (Hoechst) in blue. (D) Confocal microscopy images of zebrafish embryonic cells from wild-type and Spastin KO animals (Ctrl and Spa −/−). Cells were transfected with human REEP1 and treated with 300µM oleic acid for 18h. REEP1 labeling (green) was counterstained by bodipy (Magenta).

S4 Fig. Spastin influences Atlastin1 pattern and colocalizes with clustered LDs associated with Spartin. (A) Confocal microscopy images of HeLa cells overexpressing human Atlastin1 with Spastin M1 isoforms (after 18h administration of OA). Cherry-tagged Spastin appears in red, Atlastin1 in green, LDs (Bodipy) in magenta and nucleus (Hoechst) in blue. (B) Confocal microscopy images of HeLa cells overexpressing human Spartin with Spastin M1 isoforms (after 18h administration of OA). Cherry-tagged Spastin appears in red, Spartin in green, LDs (Bodipy) in magenta and nucleus (Hoechst) in blue.

S1 Table. Quantitative analysis of neutral lipids and phospholipids in brain and muscle from wild-type and Spa−/- zebrafish measured by mass-spectrometry. (A) Unesterified cholesterol (referred as Cholesterol), esterified cholesterol (Chol-C16, Chol-C18 and total) and triacylglycerides (TG) with specific fatty acids composition and carbon total number. (B) Individual and total saturated (SAFA), mono-unsaturated (MUFA) and Poly-unsaturated (PUFA) fatty acids. (C) Individual and total ceramides (Cer), phosphatidylcholines (PC), phosphatidyl-ethanolamines (PE), sphingomyelins (SM) and phosphatidylinositols (PI) with specific fatty acids compositions. All values correspond to lipid quantity per total protein amount. Numbers are mean ± SEM (n = 3 per group). *P<0.08, *P<0.05, P<0.01, ***P<0.001 (unpaired t-
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