Neurocalcin Delta Suppression Protects against Spinal Muscular Atrophy in Humans and across Species by Restoring Impaired Endocytosis

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Homozygous *SMN1* loss causes spinal muscular atrophy (SMA), the most common lethal genetic childhood motor neuron disease. *SMN1* encodes SMN, a ubiquitous housekeeping protein, which makes the primarily motor neuron-specific phenotype rather unexpected. SMA-affected individuals harbor low SMN expression from one to six *SMN2* copies, which is insufficient to functionally compensate for *SMN1* loss. However, rarely individuals with homozygous absence of *SMN1* and only three to four *SMN2* copies are fully asymptomatic, suggesting protection through genetic modifier(s). Previously, we identified plastin 3 (PLS3) overexpression as an SMA protective modifier in humans and showed that SMN deficit impairs endocytosis, which is rescued by elevated PLS3 levels. Here, we identify reduction of the neuronal calcium sensor Neurocalcin delta (NCALD) as a protective SMA modifier in five asymptomatic *SMN1*-deleted individuals carrying only four *SMN2* copies. We demonstrate that NCALD is a Ca²⁺-dependent negative regulator of endocytosis, as NCALD knockdown improves endocytosis in SMA models and ameliorates pharmacologically induced endocytosis defects in zebrafish. Importantly, NCALD knockdown effectively ameliorates SMA-associated pathological defects across species, including worm, zebrafish, and mouse. In conclusion, our study identifies a previously unknown protective SMA modifier in humans, demonstrates modifier impact in three different SMA animal models, and suggests a potential combinatorial therapeutic strategy to efficiently treat SMA. Since both protective modifiers restore endocytosis, our results confirm that endocytosis is a major cellular mechanism perturbed in SMA and emphasize the power of protective modifiers for understanding disease mechanism and developing therapies.

Introduction

In monogenic disorders, genetic modifiers can influence disease-causing mechanisms resulting in incomplete penetrance.¹ Identification of such modifiers is of utmost relevance since they can uncover regulatory networks and pathological mechanisms, as well as allow identification of therapeutic pathways. For recessive disorders, full protection through modifiers is extremely rare, making their identification highly challenging.

Spinal muscular atrophy (SMA), a motor neuron disease, is one of the most common and devastating autosomalrecessive disorders, for which no treatment is available yet. However, various clinical trials using antisense oligonucleotides (ASOs), small molecules, or gene therapy show highly promising ameliorations.² Most SMA individuals show homozygous absence of exon 7 of survival

motor neuron 1 (SMN1 [MIM: 600354]),³ allowing easy and efficient genetic testing.⁴ SMN1 encodes SMN, a housekeeping protein involved in snRNP biogenesis and splicing, microRNA biogenesis, transcription and translation regulation, and others;^{5–8} full absence of SMN causes embryonic lethality.9 Only humans have an almost identical copy, SMN2 (MIM: 601627), but this produces only ~10% correctly spliced full-length transcript and protein, due to a single silent mutation affecting an exonic splicing enhancer and creating a new splice silencer.^{10–12} In SMA-affected individuals, SMN2 is the only source of SMN, so its copy number (between 1 and 6) determines SMA severity.¹³ In type 1 SMA (SMA1 [MIM: 253300]), the severe and most common form (60%), the majority of individuals carry two SMN2 copies and die within the first 2 years of life. Most type 2 SMA (SMA2 [MIM: 253550])-affected individuals carry three SMN2 copies

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and are never able to walk. In type 3 SMA (SMA3 [MIM: 253400]), the mild form, most individuals carry four *SMN2* copies and are able to walk, but often become wheel-chair bound.¹⁴

Despite the important housekeeping function of SMN, reduced levels primarily cause spinal motor neuron (MN) dysfunction in all types of SMA.¹⁴ Thus, MN loss, impaired maturation and maintenance of neuromuscular junctions (NMJs), and decreased proprioceptive inputs on MN soma are hallmarks of SMA.^{15–17} Nonetheless, dramatic reduction of SMN below a certain threshold, as seen in severely affected SMA individuals or animal models, compromises almost every organ and many different cellular processes, which is in line with the essential function of SMN in all cell types.^{18,19} Therefore, we reasoned that the search for the main cellular pathway specifically driving MN dysfunction has to be carried out in mildly affected SMA individuals, in whom only motor neuron function is impaired and moreover, that protective modifiers identified in these individuals may reveal the critical underlying cellular mechanism.

To do so, we took advantage of very rarely occurring SMA-discordant families, in which relatives of SMA-affected individuals carry a homozygous *SMN1* deletion together with three or four *SMN2* copies but are clinically asymptomatic.²⁰⁻²³ In seven of these families, we previously identified the Ca²⁺-dependent protein Plastin 3 (PLS3) as a protective modifier.^{24,25} PLS3 overexpression (OE) rescues SMA across species and is specifically upregulated in MNs of asymptomatic individuals produced from induced pluripotent stem cells.^{24,26–29} Moreover, PLS3 together with the second modifier found in this study pointed us toward endocytosis as the key disturbed cellular mechanism in SMA.²⁹

Here, we report the identification of Neurocalcin delta (NCALD [MIM: 606722]), which encodes a neuronal Ca^{2+} sensor protein, as an SMA-protective modifier in humans. We show that NCALD acts as a negative regulator of endocytosis, which is in contrast to PLS3 acting as its positive regulator. We show Ca²⁺-dependent interaction of NCALD with clathrin, a protein essential in endocytic vesicles coating. We demonstrate that low SMN levels reduce voltage-dependent Ca²⁺ influx and that NCALD binds clathrin at low Ca²⁺ levels, thereby acting as a Ca²⁺-sensitive inhibitor of endocytosis. Our results, obtained from multiple in vitro and in vivo systems, show that NCALD suppression reestablishes synaptic function, most likely by restoring endocytosis. Most importantly, we prove that NCALD knockdown (KD) in various SMA animal models ameliorates major functional SMA disturbances, such as motor axon development in zebrafish or MN circuitry and presynaptic function of neuromuscular junction (NMJ) in mice. Moreover, we introduce a mild SMA mouse model generated by combined lowdose SMN-ASO treatment and heterozygous loss of Ncald that show restored motoric function. Our data support the notion that genetic modifiers reveal additional valuable treatment options, beyond existing therapies.

Material and Methods

Individuals' DNA, Fibroblast Cell Lines, and Lymphoblastoid Cell Lines

Informed written consent was obtained from each subject or their legal guardians for all biological samples according to the Declaration of Helsinki. The study has been approved by the Ethical Committee of University of Cologne (04-138). Human fibroblast and EBV-transformed lymphoblastoid cell lines (LBs) from SMA-affected individuals, carriers, and asymptomatic SMN1-deleted individuals used in this work are listed in Table S1. DNA was extracted from EDTA blood samples, primary fibroblast cell lines, and LBs using standard protocols. SMN1 and SMN2 copy number were determined by qRT-PCR or MLPA lysis (MRC Holland) as described.³⁰ For haplotype analysis, polymorphic markers Ag1-CA (D5S1556), C212 (D5F149S1/S2), VS19A (D5S435), and MIT-I105 (D5S351) were analyzed as described.³¹ SMN2 coding region was sequenced in qRT-PCR products obtained from LB-isolated RNA as described.³² PLS3 expression was analyzed as described.²⁴ All cell lines used were tested for mycoplasma contamination.

Genome-wide Linkage Analysis

Genome-wide scan was performed in 14 individuals of the Utah family using Affymetrix GeneChip Human Mapping 10K Array 2.0, which comprises 10,024 SNPs with a mean intermarker distance of 258 kb, equivalent to 0.36 cM (Affymetrix). Parametric linkage analysis was performed by ALLEGRO program³³ assuming autosomal-dominant inheritance with full penetrance and 0.0001 disease allele frequency. Haplotypes were reconstructed with ALLEGRO and presented graphically with HaploPainter.³⁴ All data handling was performed using the graphical user interface ALOHOMORA.³⁵

Transcriptome Analysis

For expression profiling, 400 ng total RNA were amplified and biotinylated using Illumina TotalPrepTMRNA Amplification Kits (Ambion) according to manufacturer's protocol. Human HT-12v3 bead arrays (Illumina) were hybridized with 750 ng cRNA for 18 hr at 58°C according to Illumina Whole-Genome Gene Expression with IntelliHyb SealSystem Manual. Arrays were washed with E1BC buffer, High-Temp Wash Buffer, and 100% ethanol, stained with streptavidine-Cy3, and washed with E1BC buffer. Fluorescence intensities were recorded on BeadArray Reader GX (Illumina). Average signal intensities without background correction³⁶ were performed with BeadStudio3.1 (Illumina). All data analysis steps were performed in the statistical environment R (v.2.10-0) with several bioconductor packages (v.2.6.1). Signal intensities were normalized with VSN (variance stabilizing and normalization quantification method³⁷) and non-informative probes were removed based on p values. Signals were averaged for individual subgroups and a linear model was designed capturing the influence of the asymptomatic group on gene expression levels.³⁸ Differences between subgroups were extracted as contrasts and analyzed with the moderated F-test (empirical Bayes method) including a correction step for multiple testing with 5%-FDR-based method.³⁹ To attribute significant regulations to individual contrasts, a decision matrix was generated based on the function "decide tests" within the "limma" package, where significant up- or downregulations are represented by values of 1 or -1, respectively.

Targeted Resequencing

To identify a potential variant regulating differential *NCALD* expression, complete *NCALD* locus \pm 1 Mb (chr8: 101,505,353–104,404,346) was deep-sequenced from gDNA of family members II-1, III-4, III-8, and IV-3 at Radboud University Medical Center Nijmegen using a 5500xl sequencing instrument (Life Technologies). ~3 Mb genomic DNA from chromosome 8 were captured using a 385K NimbleGen SequenceCapture Array (Roche).

On average, we obtained 2.7 Gb of mappable sequence data/individual. Reads were mapped to the hg19 reference genome with Life Technologies BioScope software 1.3. On average, 94% of bases originated from the target region (mean 544-fold coverage). 99.8% of the targeted region was covered ≥ 20 times. Single-nucleotide variants were subsequently high-stringency called by the DiBayes algorithm. Small insertions and deletions were detected using the Small IndelTool. Variants were annotated using an in-house analysis pipeline.

Zebrafish Experiments

All experiments were performed with the transgenic line *tg(mnx1-GFP)^{ml2TG 40}* and approved by the local animal protection committee (LANUV NRW; reference number 84-02.04.2012.A251).

Zebrafish Injection and Analysis

Morpholinos (MO) were designed against the translational start codons of respective genes (Gene Tools, LLC). *smn*-MO: 5'-CGA CATCTTCTGCACCATTGGC-3'; *ncaldb*-MO: 5'-GGAGCTTGCTG TTTTGTTTTCCCAT-3'; control-MO: 5'-CCTCTTACCTCAGTTA CAATTTATA-3'. For *NCALD* mRNA injections, human *NCALD* cDNA was cloned into pCS2+ mRNA expression vector and transcribed in vitro using mMESSAGE mMACHINE SP6 Transcription Kit (Ambion) according to manufacturer's protocol. Embryos from TL/EK wild-type and TL/EK-*hb9-GFP*⁴⁰ crossings were used to visualize the MN phenotype. Embryos were injected with the respective dose of MOs or mRNA in aqueous solution containing 0.05% PhenolRed and 0.05% Rhodamine-Dextran (Sigma). 6 hr after injection, embryos were sorted according to homogeneity of the rhodamine fluorescence signal.

Immunohistochemistry for Motor Axon Quantification

34 hpf zebrafish were manually dechorionated, fixed in 4% PFA-PBS, and permeabilized by collagenase digest of the whole animal. To visualize the primary motor axons, zebrafish were incubated at 4°C overnight in PBS-T/1%DMSO/10%FCS containing znp-1 antibody (AB2315626, Hybridoma Bank) and stained in PBS-T/ 1%DMSO/10%FCS containing donkey anti-mouse secondary antibody labeled with AlexaFluor488 (Invitrogen) after all-day washing in PBS-T/1%FCS/1%BSA (changing solution hourly) and stored in 80% glycerol/20% PBS in the dark at 4°C or embedded in low-melting agarose microslides for microscopy analysis. The structure of first ten motor axons posterior to the yolk was analyzed, rated as: (1) normal, (2) truncated (truncation ventral from midline), (3) severely truncated (shorter than midline), (4) branched I (branching ventral from midline), (5) branched II (branching at midline), or (6) branched III (branching dorsal from midline).

Western Blot Analysis of Zebrafish

48 hpf dechorionated embryos were gently spun down, sacrificed by incubation on ice, and lysed in RIPA buffer (Sigma) containing protease inhibitors (Complete Mini, Roche). The following primary antibodies were used for overnight incubation: antibeta-actin (zebrafish) (553399, Anaspec), anti-SMN (MANSMA7, Hybridoma Bank; 610646, BD Biosciences), and anti-NCALD (12925-1-AP, Proteintech). Signal detection was performed as described above.

Transmission Electron Microscopy of Zebrafish

48 hpf zebrafish were fixed in 4% PFA for 30 min and postfixed in 0.6% glutaraldehyde for another day. Samples were prepared and embedded in resin as previously described.²⁷ The thickness of semi-thin and ultra-thin sections was 0.5 and 0.1 mm, respectively. For immunogold stainings, pre-stained sections were blocked, incubated with primary antibodies (anti-clathrin [ab273, Abcam], anti-NCALD), washed in PBS, and stained with gold-labeled secondary antibodies (donkey-anti-mouse 6 nm gold [ab39616, Abcam], goat-anti-rabbit 20 nm gold [ab27237, Abcam]). Image acquisition was performed with TEM CM10 (Philips) microscope, Orius SC200W 1 Gatan camera, and the Digital Micrograph software.

Motor Behavior Analysis of Zebrafish

30 zebrafish treated with respective MOs were placed in 10 cm petri dish containing embryo medium. To trigger a swimming response, zebrafish were stimulated with an electrical impulse (60V; delay: 60 ms, duration: 4 ms, frequency: 6 pps [SD9 Stimulator]). Swimming behavior was recorded with 120 frames/s using a high-speed camera (FC-100, Casio). Swimming velocity and distance were analyzed using LoliTrack software (Loligo Systems).

Endocytosis Inhibitor Treatment

Dynasore (dynamin inhibitor) and Pitstop2 (clathrin inhibitor) (Abcam) were dissolved as stock solutions (50 mM) in DMSO. Zebrafish were dechorionated and incubated with the respective inhibitors in the medium starting at 16 hpf at 28°C on a rocking platform (20 rpm) until fixed in 4% PBS-PFA at 34 hpf. Subsequent zebrafish immunohistochemistry was performed as described above.

Electrophysiology

72 hpf zebrafish (control, *smn-*, *ncald-*, and *smn+ncald-*morphants) were anesthetized with 0.02% tricaine (in saline; Sigma) for 1-2 min and rinsed with saline containing (in mM): 134 NaCl, 2.9 KCl, 2.1 CaCl₂, 1.2 MgCl₂, 10 HEPES, 10 glucose adjusted to pH 7.8. Zebrafish were decapitated and pinned under saline in a Sylgard-coated (Dow Corning) recording chamber (~3 mL volume). Skin was removed using a tungsten pin and forceps; preparation was incubated in 3M formamide (in saline; Carl Roth) for 2 min to prevent muscle contractions. After rinsing the preparation, the superficial layer of ventral slow muscle cells was removed by scratching with a tungsten pin to expose deeper fast skeletal muscle cells and remaining superficial slow muscles were removed with a low-resistance pipette (~2 MΩ). The preparation was continuously superfused with saline at a flow rate of ~2 mL/min⁻¹. Experiments were carried out at ~24 $^{\circ}$ C. Muscle cells were visualized with a fixed-stage upright microscope (Zeiss Axio Examiner, Zeiss), using a 40× water immersion objective (Zeiss) with infrared-differential interference contrast and fluorescence optics. Fast muscle cells were identified by their orientation to the spinal cord and ability to generate action potentials.

Caenorhabditis elegans Experiments

Strains used were as follows: LM99 smn-1(ok355)I/hT2(I;III),⁴¹ HA1981 +/hT2(I;III), HA2530 +/hT2(I;III);ncs-1(qa401)X, HA2531 smn-1(ok355)I/hT2(I;III);ncs-1(qa401)X, HA2599 +/hT2(I;III);uls72, and HA2623 smn-1(ok355)I/hT2(I;III);uls72 were maintained at 20°C under standard conditions. +/hT2 strains were used as control for genetic background; RNAi studies were undertaken in a

sensitized background (transgene uIs72) expressing the SID-1 dsRNA channel in neurons.⁴²

C. elegans Pharyngeal Pumping

Pharyngeal grinder movement in any axis was scored as a pumping event. Average pumping rates (\pm SEM) were combined from at least three independent trials ($n \ge 25$ animals in total). For RNAi knockdown, animals were reared for two generations (F2) on either control vector L4440 or C44C1.3/*ncs*-1(*RNAi*) in HT115. *ncs*-1 RNAi clone contains genomic DNA amplified by primers 5'-AAATCGTCTAGCTGTAGTGTCGC-3' and 5'-TTGTGC TCCCTACACTTTGTTTT-3' inserted into L4440. Clone was verified by sequencing.

Mouse Experiments

All mouse experiments were approved by LANUV NRW (reference number 9.93.2.10.31.07.186 and 84-02.04.2014.A 126). The Taiwanese SMA mice (FVB.Cg-Tg(SMN2)2Hung Smn1tm1Hung/J, Stock Number 005058) and heterozygous *Ncald*^{ko/wt} mice (Bl6N(Cg)-*Ncald*^{tm1.1(KOMP)VIcg}/J, Stock Number 018575) were purchased from Jackson Laboratory. The severe SMA (*Smn*^{ko/ko}; *SMN2*^{tg/0}) mice and the corresponding heterozygous *Smn* (HET [*Smn*^{kowt-}; *SMN2*^{tg/0}]) mice were produced as previously described.^{9,43} The breeding scheme and genotypes for SMA-*Ncald*^{ko/wt} and HET-*Ncald*^{ko/wt} mice are similar to SMA+ASOtreated mice (Figure 5A), except that all animals were on congenic C57BL/6N and untreated.

Primers used for mouse genotyping are as follows: mmu *Smn*KOfw: 5'-ATAACACCACCACTCTTACTC-3'; mmu *Smn*KOrev1: 5'-AGCCTGAAGAACGAGATCAGC-3'; mmu *Smn*KOrev2: 5'-TAG CCGTGATGCCATTGTCA-3'; hsa *SMN2fw*: 5'-CGAATCACTTGAG GGCAGGAGTTTG-3'; hsa *SMN2*rev 5'-AACTGGTGGACATGGC TGTTCATTG-3'; mmu *Ncald*KOfw: 5'-CGGTCGCTACCATTAC-3'; mmu *Ncald*KOrev: 5'-GCATGTGTGACAACAG-3'.

A mild SMA mouse model was produced by suboptimal subcutaneous injection of severe SMA mice (50% FVB/N: 50% C57BL6/N) on P1 with 30 μ g of SMN-ASO (IONIS Phamaceuticals) using a MICROLITER syringe (Hamilton). The SMN-ASO was diluted as previously described.¹⁹ SMA-*Ncald*^{ko/wt}+ASO and HET-*Ncald*^{ko/wt}+ASO were produced using the breeding scheme in Figure 5A. Unless stated otherwise, all mouse experiments were performed blinded for genotype and treatment.

Mouse Motoric Tests

Righting reflex test was performed as previously described.⁴⁴ Righting time scores were evaluated as followed: 0–2 s, 1; 3–4 s, 2; 5–6 s, 3; 7–8 s, 4; 9–10 s, 5; \geq 11 s, 6. Muscle strength was assessed in P73 SMN-ASO injected mixed₅₀ background mice by the animal's grasp of a horizontal metal bar mounted to a highprecision force sensor (Grip strength meter, TSE Systems). Muscle force was recorded in pounds and converted to Newton [N].

Quantification of Proprioceptive Inputs

Analysis of proprioceptive input on MN soma was performed as described.²⁹ The spinal cord was dissected from euthanized mice and fixed in 4% PFA overnight. The lumbar L4-L5 region was rinsed in PBS, embedded in tissue freezing medium (Jung) after cryoprotection (first day: 20% sucrose, second day: 30% sucrose), and sliced into 100 μ m sections (cryostat, Leica). Samples were permeabilized, blocked in PBS/4% BSA/1% Triton/PBS for 1 hr, and incubated with anti-CHAT (Choline acetyltransferase, a MN-specific marker) (AB144P, Millipore) and anti-VGLUT1 (Vesicular glutamate transporter 1 or SLC17A, an excitatory neurotransmitter used for proprioceptive inputs) (135303, Synaptic Systems)

antibodies overnight. Samples were washed and incubated with secondary antibodies (donkey anti-rabbit AlexaFluor488, donkey anti-goat AlexaFluor568) and mounted in Mowiol. Images were taken in Z stacks of 30–60 slices of $0.3 \,\mu$ m interval. Proprioceptive input numbers on MN and MN soma size were quantified using the ImageJ software.

Quantification of NMJ Size and Maturity

The transversus abdominis (TVA) muscle was prepared at the indicated time points and fixed in 4% PFA for 20 min. TVA was stained with anti-NF-M (Neurofilament M, NEFM, used as neuronal axon and dendritic marker [Hybridoma-Bank]) and with secondary goat-anti mouse AlexaFluor488 and Bungarotoxin (both Invitrogen, labeled with AlexaFluor555). Surface area of Bungarotoxinpositive post-synapse was measured by ImageJ with threshold set to the method established by Li. NMJ immaturity index was analyzed as described previously:⁴⁵ NMJs exhibiting \geq 3 perforations were evaluated as mature, NMJs with <3 perforations as immature.

FM1-43 Endocytic Uptake at NMJ under Electrical Stimulation

FM1-43 endocytic uptake at NMJ under electrical stimulation was undertaken as recently described.²⁹ Three animals per genotype and stimulation set were used. Imaging was performed as described above. All imaging processes and analyses were performed double-blinded. Images were analyzed with ImageJ using a macro setting and Li threshold method applied to the postsynaptic terminals to delineate the area of interest in the presynaptic site.

Microscopy

Unless indicated otherwise, all microscopic experiments were performed with a fully motorized fluorescence microscope AxioImager M2 (Zeiss) equipped with an ApoTome. All quantitative measurements were performed using Zen software (Zeiss) and ImageJ and evaluated with indicated statistical packages.

Primary Motor Neuron Culture

Spinal cords were dissected from E13.5 mouse embryos.⁹ Neurons were singularized with trypsin (Worthington) and DNase (Sigma), sieved, plated on poly-D-lysine/laminin (Sigma)-coated coverslips, and cultured in neurobasal medium with B27 supplement, 2 mM L-glutamine, $1 \times$ pen-strep (Invitrogen) containing 50 ng/µL BDNF, 50 ng/µL GCNF, and 50 ng/µL CNTF (Peprotech) at 37°C in a humidified incubator with 5% CO₂.

Cell Culture Experiments Quantitative RT-PCR

RNA was extracted from cell lines using RNeasy kit (QIAGEN). 150 ng RNA was reversely transcribed to cDNA (Quantitect Reverse Transcription Kit, QIAGEN). For *NCALD* cDNA measurements, 9 ng cDNA was used for RT-PCR (LightCycler, Roche). RT-PCR was performed in triplicates according to manufacturer's protocol (annealing temperature 68°C, *NCALD* cDNA primers: 5'-GGA ATGCCCAGAGCCCCAGTGT-3'; 5'-GCCCCAACCCCCGAGTCTT ACG-3'). Standard curve-based absolute transcript quantification was performed using Excel (Microsoft). For statistical evaluation, the Student's t test was applied. For quantitative measurements of *SMN* and *PLS3*, previously described protocols were used.²⁴

siRNA-Mediated RNA Knockdown

For all siRNA experiments, NSC34 (CLU140)³² and PC12⁴⁶ cells were transfected with Dharmafect1 (Thermo Scientific) according to manufacturer's protocol. siTOX (Dharmacon) and AllStars Negative Control (QIAGEN) siRNA were used as controls. siRNAs sequences: mmu-*Smn*: 5'-AAGAAGGAAAGTGCTCACA

TA-3'; mmu-*Ncald* 5'-CAGGTGATTCACCCATTATAA-3'; rn-*Smn* 5'-CCCGACCTGTGAAGTAGCTAA-3'; rn-*Ncald* 5'- AGAGACTTC CTAGCAATTTAA-3. After incubation, cells were harvested for protein isolation or imaging. Every experiment was performed at least in triplicates.

Transient Overexpression

Human *NCALD* cDNA was cloned into pcDNA3.1/CT-GFP TOPO using primers NCALD-FWD 5'-ATGGGGAAACAGAACAGCAA G-3' and NCALD-REV 5'-GAACTGGCCGGCACTGCTC-3' (IDT) and manufacturer's protocol (Invitrogen). To overexpress human NCALD-GFP, NSC34 cells were transfected with Dharmafect1 according to manufacturer's protocol.

Western Blot Analysis

Cells were lysed on ice in RIPA buffer (Sigma) containing protease inhibitors (Complete Mini, Roche). The following primary antibodies were used: anti-ACTB (actin, beta), used as control for equal loading (A5316, Sigma), anti-SMN (MANSMA7, Hybridoma Bank; 610646, BD Biosciences), anti-NCALD (12925-1-AP, Proteintech), and anti-CLTC (clathrin heavy chain) (C1860, Sigma). Signal was detected with HRP conjugated-secondary antibodies and Chemiluminescence reagent (Thermo Scientific) according to manufacturer's protocol.

NCALD Co-immunoprecipitation

NSC34 cells transiently transfected with pcDNA/FLAG-His-NCALD or control vector were lysed in the following buffer: 50 mM Tris/HCl, 5% (w/v) glycerol, 270 mM sucrose, 0.5%(v/v) Tween 20, 0.1%(v/v) β -mercaptoethanol (pH 7.5), with protease inhibitor cocktail (Complete Mini, EDTA-free, Roche). Immunoprecipitations were performed in 1 mM EGTA/1 mM EDTA or in the presence of 100 μ M free Ca²⁺. Cell lysates were immunoprecipitated with FLAG-M2 affinity beads (Sigma) under gentle agitation overnight at 4°C. Bound proteins were eluted in laemmli buffer (240 mM Tris-HCl [pH 6.8], 6% SDS, 30% (v/v) glycerol, 0.06% bromophenol blue (w/v), 16%(v/v) β -mercaptoethanol) and analyzed by western blots as described above.

Immunocytochemistry

Cells were cultured on laminin-coated coverslips, washed with PBS, fixed in 4% PFA/4% sucrose (AppliChem), permeabilized in PBS-T (PBS/0.2%Tween20 [AppliChem]), and blocked in blocking solution (PBS-T/5%BSA [Sigma]/5% FCS [Biochrom]). Cells were incubated in blocking solution overnight at 4°C containing the following primary antibodies: α -HB9 (homeobox 9), used as MN-specific marker (1:100); AB2145209 from Hybridoma Bank; α-SV2 (Synaptic vesicle glycoprotein 2), used as synaptic vesicle marker; AB2315387 (SV2-c) from Hybridoma Bank; α-CHAT, AB144P from Millipore; α-Tau (axon-specific marker), sc-390476 from Santa Cruz; and α -NCALD. After washing in PBS, cells were incubated with secondary antibodies labeled with AlexaFluor488, AlexaFluor647, or AlexaFluor568 (Invitrogen) in PBS, optionally with phalloidin-AlexaFluor568 (Invitrogen). Cells were washed and mounted on objects slides with Mowiol (Sigma) for imaging. Endocytosis Assay

Fibroblasts were plated in DMEM (Invitrogen) and starved for 10 min in starvation media (DMEM transparent [HEPES], 2% FCS) prior to fluorescein isothiocyanate (FITC)-Dextran treatment (5 mg/mL, Sigma) for respective time periods at 37°C. Subsequently, cells were washed with ice-cold PBS and fixed in 4% PFA for 10 min. After washing, cells were stained with phalloidin-AlexaFluor568 and DAPI (Invitrogen) and mounted with Mowiol for imaging.

Flow Cytometry Analysis

NSC34 cells were transfected with indicated siRNAs for 48 hr prior to 6 hr starvation and incubation with 5 mg/mL FITC-Dextran

(Sigma) for 20 min at 37° C. Cells were trypsinized (Trypsin, Sigma) on ice and washed with PBS. Uptake of FITC-Dextran was measured with FACS Calibur (BD Biosciences) and analyzed with Cyflogic software (CyFlo Ltd.). Dead cells were excluded by propidium iodide staining (10 µg/mL, Sigma).

Ca²⁺ Current Recordings in NSC34 and PC12

Whole-cell recordings were performed at 24°C. Electrodes (tip resistance 2.5-3 MΩ) were made of borosilicate glass (0.86 mm OD, 1.5 mm ID, Science Products) with a temperature-controlled pipette puller (PIP5, HEKA Elektronik) and filled with solution containing (in mM) 133 CsCl, 1 CaCl₂, 2 MgCl₂, 10 HEPES, and 10 EGTA, adjusted to pH 7.2 and osmolarity of 415 mOsm. During experiments, cells were constantly superfused with saline solution containing (in mM) 84 NaCl, 20 CsCl, 2.5 KCl, 10 CaCl₂, 2 MgCl₂, 10 HEPES, and 30 glucose, adjusted to pH 7.3 and osmolarity of 310 mOsm. To isolate Ca²⁺ currents, a combination of pharmacological blockers and ion substitution was used. Transient voltage-gated Na⁺ currents were blocked by tetrodotoxin (10⁻⁶M TTX, T-550, Alomone). 4-aminopyridine (4AP, 4 \times 10⁻³ M, A78403, Sigma) blocked transient K⁺ currents (I_A) and tetraethylammonium (TEA, 2 × 10-3, Sigma) blocked sustained K⁺ currents ($I_{K(V)}$) and Ca²⁺-activated K⁺ currents $(I_{K(Ca)})$. The pipette solution did not contain potassium. Whole-cell voltage-clamp recordings were made with EPC10 patch-clamp amplifier (HEKA Elektronik) controlled by Patchmaster program (V2x53, HEKA-Elektronik). Electrophysiological signals were low-pass filtered at 2.9 kHz (3pole Bessel filter). Data were sampled at 50 µs intervals (20 kHz). The offset-potential and capacitance were compensated using "automatic mode" of EPC10 and liquid-junction potential between intracellular and extracellular solution of 2.5 mV (calculated with Patcher's Power-Tools plug-in) was compensated. Whole-cell capacitance was determined using EPC10 capacitance compensation (C-slow). To remove uncompensated leakage and capacitive currents, p/6 protocol was used.⁴⁷ Voltage errors due to series resistance (RS) were minimized using RS compensation of EPC10 to 70%-80% with 100 μ s time constant (τ).

Statistical Analysis

Statistical Analysis

If not mentioned otherwise, all statistical analyses were performed using software programs Excel 2013 (Microsoft), GraphPad Prism 6 (GraphPad Software), and Sigma Plot 11 (Systat Software); ANOVA, Mann-Whitney U test, Fisher's exact test, or unpaired two-tailed Student's t tests were applied. All data are represented as mean \pm SEM/SD. Significance of RNA expression and protein levels was tested using a directional Student's t test for uncorrelated samples. For experiments performed in *C. elegans*, Mann-Whitney U test was performed. Significance in the differences of mouse behavioral analyses, NMJ, and muscle fiber surface area size, motor axon length, proprioceptive inputs on MNs, NSC34 neurite length, and width of the synaptic cleft was determined by the use of 1-way ANOVA or directional Student's t test for uncorrelated samples. Survival was analyzed using Kaplan-Meier method by log rank test.

For all studies using mice, animals numbers were calculated prior to experiments by power calculation using the G*Power 3.1.7 software (Power = 0.8 and alpha-error = 0.05). Endpoint criteria for mouse experiments were defined in animal application prior to experiments. Animal samples were processed equally and allocated to experimental groups post-analysis. For all other

experiments, sample size was estimated based on the known variability of the assay.

Values of p<0.05 were considered significant. In all cases, three levels of statistical significance were distinguished: *p <0.05, **p <0.01, and ***p <0.001.

Specific statistical tests, sample size, and p values are indicated in the figure legends.

Statistical Analysis of Electrophysiology

Data were analyzed using Spike2 and statistical analysis was performed in GraphPad Prism 5.05 (GraphPad Software). All calculated values are shown as mean \pm SEM. The EEP frequencies for each cell were measured as mean frequencies over 30 s intervals. Frequencies before and during NMDA application were compared by a paired t test for each group. Kruskal-Wallis test followed by Dunns multiple comparisons was used to compare EPP frequencies in different groups. A significance level of p < 0.05 was accepted for all tests.

Results

Identification of NCALD as a Potential SMA Modifier by Genome-Wide Linkage and Transcriptome-Wide Differential Expression Analysis

In a four-generation Mormon family from Utah, we identified seven individuals carrying homozygous *SMN1* deletions, two affected by type 1 SMA and five fully asymptomatic, except for increased photosensitivity (Figure 1A; see Supplemental Data for full clinical investigation description of Utah family members).

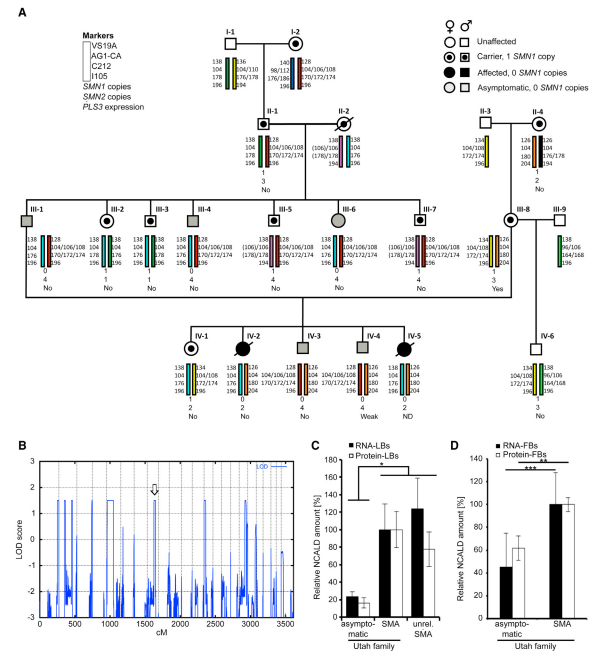
Haplotype analysis of SMA regions showed a co-segregation of three different SMA alleles (Figure 1A). The two type 1 SMA-affected individuals carried no *SMN1* and two *SMN2* copies. By contrast, all five asymptomatic individuals showed homozygous absence of *SMN1* and presence of four *SMN2* copies, resembling a genotype associated with type 3 SMA¹³ (Figure 1A). *SMN2* sequencing excluded any further variants affecting expression. In lymphoblastoid cells (LBs), SMN RNA and protein levels were similar to those in typical type 3 SMA-affected individuals, thus excluding *cis-* and *trans-*acting factors regulating *SMN2*. Increased *PLS3* expression was not found (Figure 1A; GEO: GSE58316). Thus, we concluded that a previously unknown SMA modifier potentially protects these individuals.

To identify the SMA modifier, we combined linkage with transcriptome-wide differential expression analysis. Assuming a dominant mode of inheritance, a parametric linkage analysis with 14 family members revealed 8 positive peaks with a maximum LOD score of 1.5 (Figure 1B). In parallel, a transcriptome-wide differential expression analysis with 12 total RNA samples was performed (GEO: GSE58316) and revealed 17 transcripts significantly differentially regulated in asymptomatic individuals (Table S2). *NCALD* was represented by two independent hybridization probes on the array, both showing a 4- to 5-fold down-regulation in the asymptomatic group versus familial type 1 SMA or an independent type 3 SMA group. Most importantly, *NCALD* was the only transcript localized in one of

the eight linked regions on chromosome 8q22.3 (between rs28144 and rs958381), making it a highly likely candidate. Microarray data were confirmed by qRT-PCR and western blot (Figures 1C and 1D).

To search for the potential genetic mechanisms involved in reduced NCALD expression, targeted resequencing of ~3 Mb genomic DNA encompassing NCALD in five family members was carried out. On average 2,723 variants were called per sample. Based on previous haplotype data, we filtered for heterozygous variants shared between individuals II-1, III-1, III-4, and IV-3 but absent in III-8. This yielded 43 variants (21 previously annotated SNPs), none of which were in the NCALD coding region. Only the SNP rs147264092 in intron 1 (HGVS: NC_000008.10: g.103128181_103128182insCT) with a minor allele frequency (MAF) = 0.11 (1000 Genomes) was located in NCALD UTR (Table S3). Approximately 600 kb upstream of NCALD, we identified a 17-bp deletion (HGVS: NC_000008.10: g.103783522_103783538del17; dbSNP: rs150254064; 1000 Genomes: MAF = 0.04) perfectly segregating with the modifier haplotype (Figure S1) that seemed interesting. The 17-bp deletion is localized adjacent to an H3K27AC block and a super enhancer (ENCODE), which may influence NCALD expression. We hypothesize that the combination of both variants acts on NCALD expression (unpublished data). Both variants were further analyzed in 50 SMN1-deleted individuals, who were chosen because of a discrepant SMA severity according to their SMN2 copy number, and 65 controls. The combination of both variants was found in one individual, who unexpectedly carried only one SMN2 copy. This genotype is regarded as a type 0 SMA with death in utero or immediately after birth.⁴⁸ In contrast, this individual survived 9 months, suggesting a potential protection by a genetic modifier, which could be NCALD. No LBs were available to test expression. The combination of both variants on a haplotype is a very rare event: 0.003 (13/5,008 haplotypes included in the Phase 3 1000 Genomes project, see LDlink). Since homozygous deletions of SMN1 occur with a frequency of 1:6,000 to 1:20,000 depending on ethnicity,49 the combination of homozygous SMN1 deletion and the chromosome 8 modifier haplotype would statistically occur in fewer than 1:8,000,000 people. Further work is in progress to fully understand the impact of these variants on chromatin structure and NCALD expression. However, since understanding gene regulation and the interplay between cis- and trans-regulatory elements is extremely challenging and may not yield solid results, we decided to take the direct approach and analyze the impact of NCALD reduction in four different SMA animal models: C. elegans, zebrafish, and a severe and a mild SMA mouse model.

NCALD is one of 14 neuronal calcium sensor (NCS) proteins in mammals. These proteins are highly conserved across species and primarily involved in neuronal Ca^{2+} signaling.^{50,51} NCALD encodes a ~22 kDa protein that contains two pairs of EF-hand domains and an N-terminal





(A) Pedigree of the Utah family: haplotype analysis of microsatellite markers in the 5q13 SMA region and *SMN1* and *SMN2* copies are indicated. Black filled symbols indicate SMA-affected individuals; gray filled symbols indicate asymptomatic *SMN1*-deleted individuals; and symbols with a dot indicate SMA carriers. Quantification of *PLS3* expression in LBs was done according to Oprea et al.²⁴ Note that weak *PLS3* have no impact on SMA phenotype.²⁴

(B) Genome-wide linkage analysis identified eight regions with positive LOD scores. Open arrow marks 8q22.3 region containing NCALD.

(C) Verification of microarray results (Table S2) of *NCALD* RNA and protein in lymphoblastoid (LB) cells (NCALD levels are relative to NCALD in SMA-affected individuals of the Utah family [set to 100%]). *NCALD* is represented by two independent probes on the expression array, showing a 4- to 5-fold downregulation in the asymptomatic group versus familial type 1 SMA or an independent type 3 SMA group. Three independent experiments including all 17 cell lines (asymptomatic, n = 5; symptomatic, n = 2; independent SMA-III, n = 10) were performed. Error bars indicate SD; * $p \le 0.05$.

(D) Expression analysis of *NCALD* RNA and proteins in fibroblasts (FB) derived from the Utah family (asymptomatic, n = 5; symptomatic, n = 2). Three independent experiments including all seven cell lines were performed. Error bars indicate SD; ** $p \le 0.01$; *** $p \le 0.001$.

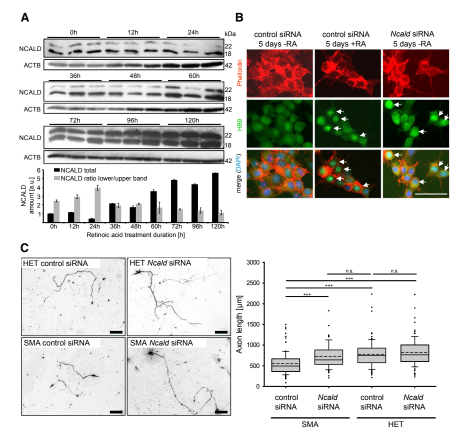


Figure 2. NCALD Downregulation Restores Neurite Outgrowth Defect in SMN-Deficient Neuronal Cells

(A) Western blot of NSC34 cells treated with 1 μ M retinoic acid (RA) for 0–120 hr as a model of MN differentiation and maturation (n = 3 independent experiments). Error bars indicate SEM.

(B) Ncald siRNA-treated NSC34 cells show signs of MN differentiation (HB9-positive staining, marked with white arrows) even in absence of RA (right). As positive control, cells were differentiated with RA and treated with control siRNA (middle). Negative control was treated only with control siRNA (left). Scale bar represents 100 µm. (C) Primary MNs from SMA or HET murine embryos were fixed at 8 DIV and stained with anti-neurofilament M (anti NF-M). Quantitative analysis of axon length of MNs. SMA: n = 7, HET: n = 6, n = 100 per measurement; *** $p \le 0.001$; dashed line indicates mean; straight line indicates median; values covered from 25%-75% and dotted outliers at <5% and >95% CI. Scale bars represent 100 µm.

We found that *Ncald* knockdown in *Smn*-deficient NSC34 cells restored impaired neurite outgrowth to control levels (Figure S2E). Similar results were

myristoyl anchor, which enables switching from cytosolic to membrane-bound forms in a Ca²⁺-dependent manner.^{52,53} A Ca²⁺-dependent mobility shift of both myristoylated and non-myristoylated forms was reported.⁵⁴ NCALD is highly abundant in cerebral neurons, spinal MNs, and in axonal growth cones.⁵⁵ NCALD overexpression inhibits neurite outgrowth.⁵⁶ NCALD is important in phototransduction,⁵⁷ which may explain photosensitivity in asymptomatic individuals. Importantly, NCALD interacts with clathrin and actin, both of which are involved in endocytosis and synaptic vesicle recycling.^{58,59}

NCALD Knockdown Triggers MN Differentiation and Restores Neurite and Axonal Growth in SMA

First, we analyzed NCALD levels during MN differentiation and maturation in NSC34 cells treated with retinoic acid (RA)⁶⁰ to induce differentiation and observed a steady increase in NCALD amount over time under RA treatment (Figure 2A). siRNA-mediated *Ncald* reduction (Figure S2A) induced MN differentiation (indicated by HB9-positive staining) and triggered neurite outgrowth even without RA treatment (Figure 2B). In contrast, NCALD overexpression in RA-treated NSC34 cells impaired neurite outgrowth (Figures S2B and S2C). NCALD is highly abundant in axonal growth cones of spinal MNs.⁵⁵ In addition, we show that it localizes at the presynaptic terminals of NMJs, suggesting a potential role at the NMJ (Figures S2D and S2E). obtained in cultured primary MNs from SMA (*Smn*^{ko/ko}; *SMN2*^{tg/0}) versus HET (*Smn*^{ko/wt};*SMN2*^{tg/0}) embryos, where reduced axon length of SMA MNs⁶¹ was restored by siRNA-mediated *Ncald* knockdown (Figure 2C). These findings indicate that reduced NCALD levels counteract the impaired axonal development of SMN-deficient MNs.

ncald Knockdown Restores Axonal Growth and NMJ Functionality in Zebrafish *smn* Morphants

Human NCALD and its ortholog in zebrafish are 98% identical, suggesting important conserved functions across species. We next investigated the modifying effect of *ncald* in vivo in a *mnx1:eGFP*-expressing zebrafish line⁴⁰ by MO-mediated knockdown of either smn, ncald, or both together. Consistent with previous results, smn depletion resulted in motor axon-specific outgrowth defects, such as truncations and ectopic branches^{24,62} (Figure 3A). Knockdown of ncald led to enhanced motor axons branching, whereas double *smn*+*ncald* knockdown fully rescued the truncated motor axon defect associated with Smn deficiency (Figures 3A, 3C, and S3A). Knockdown efficiency was confirmed by western blot (Figure 3B). We also found that overexpression of human NCALD mRNA in wild-type zebrafish caused truncation and branching of motor axons (Figure S3B), resembling the phenotype of *smn* morphant zebrafish (Figure 3A) similar to NSC34 cells (Figure S2C).

During NMJ maturation, the width of the synaptic cleft is increasing, which is essential in neurotransmission.⁶³ Ultrastructural analysis of the synaptic cleft revealed an

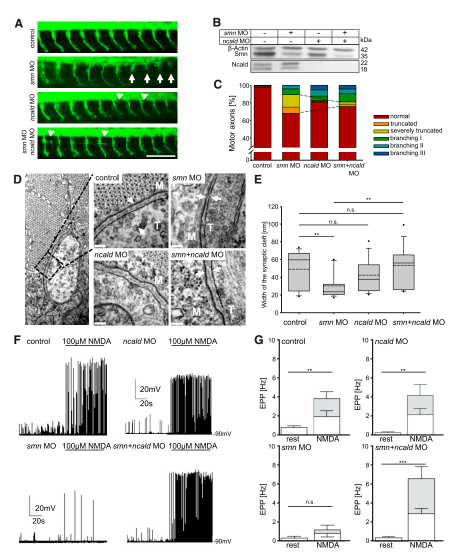


Figure 3. *Ncald* Reduction Corrects the Phenotype in *Smn*-Deficient Zebrafish

(A) First 10 motor axons posterior to the yolk globule of 34 hpf zebrafish embryos injected with respective morpholinos (MO). White arrows mark truncated motor axons. Arrowheads mark extensive branching in *ncald* or *smn+ncald* morphants; green shows Znp1 staining, for motor axons. Scale bar represents 100 μ m. (B) Western blot of lysates of zebrafish embryos injected with indicated MO.

(C) Quantification of motor axon phenotype. Dashed lines mark the rescue of the truncation phenotype (** $p \le 0.01$). *smn+ncald* and *ncald* morphants showed increased branching. n > 500 motor axons per MO injection.

(D) TEM images of NMJs of 48 hpf zebrafish embryos injected with respective MO. White arrows mark synaptic clefts including basal lamina. M indicates muscle fiber, T indicates nerve terminal. Scale bars represent 100 nm.

(E) Quantification of synaptic cleft width of MO-injected 48 hpf fish (n = 15 per treatment). **p \leq 0.01, dashed line indicates mean; straight line indicates median; values covered 24%–75% and dotted outliers at <5% and >95% CI.

(F and G) Whole-cell current clamp recordings EPPs (F) and quantification (G) of mean EPP frequencies in ventral fast muscle cells of control (n = 12), *smn* (n = 10), *ncald* (n = 11), and *smn*+*ncald* (n = 12) morphants under control conditions or NMDA induction. White bar parts reflect the mEPP frequencies, gray bar parts reflect the frequency of the TTX-sensitive large EPPs. **p \leq 0.01; ***p \leq 0.001. Error bars indicate SEM.

impaired NMJ maturation in *smn* morphants (Figures 3D and 3E). The width of the synaptic cleft in *smn* morphants was significantly smaller than in controls or *ncald* morphants; double *smn*+*ncald* knockdown significantly restored synaptic maturation, resulting in a cleft width similar to control embryos (Figures 3D and 3E).

To test the functionality of neuromuscular synapses between caudal primary MNs and ventral fast muscle cells,⁶⁴ we performed whole-cell patch clamp recordings from muscle cells during MN stimulation in control (ctrl), *smn, ncald,* and *smn+ncald* zebrafish morphants. We recorded spontaneous endplate potentials at rest (without stimulation) and during MN stimulation by NMDA (N-methyl-D-aspartate, agonist of NMDA receptors) (Figure S3C). In controls, we recorded at rest small endplate potentials that were primarily not tetrodotoxin (TTX) sensitive (Figures S3D and S3E) and mostly resembled miniature endplate potentials (mEPPs).⁶⁵ During NMDA stimulation, the mEPP frequency did not significantly increase, but large TTX-sensitive endplate potentials and muscle action potentials were induced by MN spikeevoked transmission. In *smn* morphants, a significantly lower spontaneous mEPP frequency and only occasional action potentials during NMDA stimulation were observed (Figure 3F). In the *smn*+*ncald* morphants, the spontaneous mEPP frequency was slightly increased and the frequency of large NMDA-induced EPPs was restored to control levels (Figures 3F and 3G). In line with the electrophysiological data, swimming velocity after electrical stimulation was reduced in *smn* morphants but rescued in *smn*+*ncald* morphants (Figure S3F). Together, these results show that Ncald knockdown rescues neural circuit function at the NMJs of *smn* morphants.

Loss of NCALD Ortholog Suppresses Defects of *C. elegans* SMA Model

C. elegans lacking the *SMN* ortholog *smn-1*, referred to here as *Cesmn-1*, show neuromuscular defects, including decreased pharyngeal pumping rate (Figure S4A).^{26,41} The *C. elegans* ortholog of *NCALD* is encoded by neuronal calcium sensor-1 (*ncs-1*).⁶⁶ Either *ncs-1* knockdown by RNA interference or introduction of the *ncs-1(qa401)*

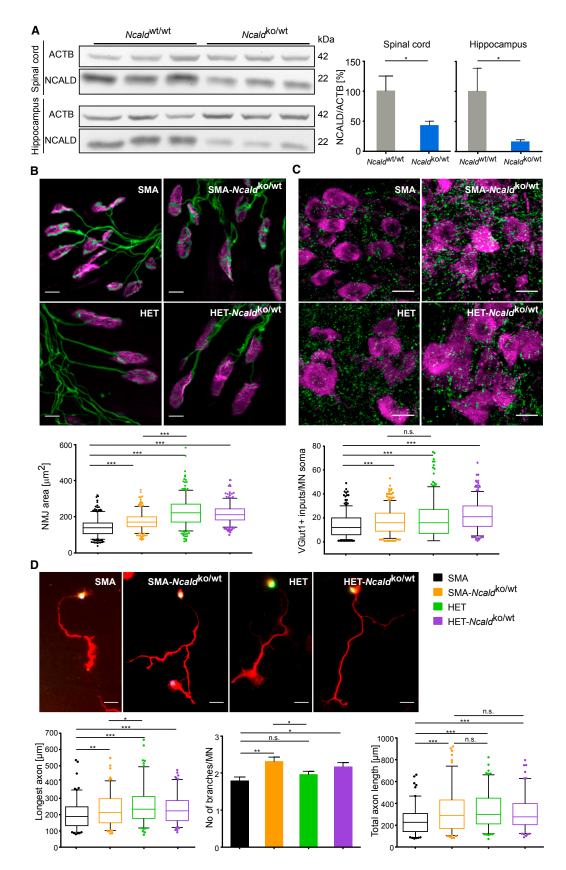


Figure 4. Heterozygous *Ncald* KO Improves Axonal Outgrowth, Proprioceptive Input, and NMJ Size in Severe SMA Mice (A) Western blot and quantification of NCALD and ACTB (loading control) in spinal cord and hippocampus of P10 WT and *Ncald*^{ko/wt} mice. * $p \le 0.05$. Error bars indicate SD.

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loss-of-function allele in *Cesmn-1* animals significantly ameliorated pumping defects (Figures S4B and S4C), confirming that NCALD loss ameliorates the SMN loss-of-function-induced neuromuscular defects across species.

Heterozygous *Ncald* KO Ameliorates Motor Neuron Development in Severe SMA Mice

We took advantage of an *Ncald* knockout mouse (*Ncald*^{ko/ko}) recently generated by the Knockout Mouse Phenotyping Program at the Jackson Laboratory. Heterozygous *Ncald*^{ko/wt} mice are asymptomatic and show >50% reduction of NCALD levels in spinal cord and brain (Figure 4A). Homozygous *Ncald*^{ko/ko} mice are viable and fertile; however, preliminary reported data by the International Mouse Phenotype Consortium (IPMC) (online database) and our data revealed behavioral abnormalities, vision defects, and metabolic impairment. In contrast, heterozygous *Ncald*^{ko/wt} mice showed no gross morphological or behavioral problems even at 18 months of age. Since asymptomatic individuals show reduced but not full loss of NCALD, we used the heterozygous *Ncald*^{ko/wt} animals for all further experiments herein.

The Ncald^{ko/wt} allele was bred into a severe SMA mouse model9 on pure C57BL/6N background. Both SMA and SMA-Ncaldko/wt mice die at a mean age of 13 days and there is no difference in weight progression at this age (Figures S5A and S5B). Severe SMA mice show multi-organ failure^{27,43,67} due to very low SMN levels, which could not be rescued by heterozygous Ncald knockout alone. Nonetheless, we found that other hallmarks of SMA were improved upon heterozygous Ncald knockout: the size of the NMJs in the transversus abdominis muscle (TVA) was increased and the number of proprioceptive inputs on MN soma was elevated in SMA-Ncald^{ko/wt} versus SMA mice (P10) (Figures 4B and 4C). Moreover, SMA-*Ncald*^{ko/wt} mice showed more inputs per MN than SMA mice independent of cell size (Figure S5C). A comparison of axonal development in cultured primary MNs revealed a large impact of NCALD reduction on axonal growth and arborization (Figure 4D), confirming our initial results with siRNAmediated Ncald knockdown (Figure 2C). Therefore, NCALD reduction counteracts impaired axonal development and restores NMJ size in SMN-deficient mice but is not able to improve survival due to severe multiple organ impairment.

Combinatorial Therapy with a Suboptimal Low-Dose SMN-ASO and Reduced *Ncald* Expression Ameliorates SMA Pathogenesis in a Severe SMA Mouse Model

In our study, we combined suboptimal low-dose SMN-ASOs with heterozygous Ncald knockout mice for four reasons: (1) asymptomatic individuals carry four SMN2 copies, similar to typical type 3 SMA-affected individuals, but not two SMN2 copies as our severe SMA mouse model or most type 1 SMA-affected individuals; (2) genetic modifiers efficiently protect against SMA only if a sufficient SMN level is present to suppress inner organ dysfunction;²⁹ (3) NCALD expression is mainly restricted to neuronal tissues, so its beneficial effect is directed to MN, but cannot improve other peripheral organs affected in severe type of SMA; and (4) type 1 SMA-affected individuals, currently treated with SMN-ASOs, show only a moderate SMN elevation and may need additional drugs/molecules supporting MN function. For these reasons, we chose to establish a mild SMA mouse model that shows no impairment in lifespan or peripheral organs, but has a prominent motoneuronal phenotype. Since presymptomatic subcutaneous (s.c.) injection of high-dose SMN-ASO in severely affected SMA mice fully rescues SMA⁶⁸ and low-dose SMN-ASO in C57BL6/N congenic mice increased survival to only 1 month (intermediate phenotype),²⁹ we opted for a different strategy to produce a mild SMA phenotype. We crossed C57BL/6N Ncaldko/wt;Smnko/wt males with FVB/N Smn^{ko/ko};SMN^{tg/tg} females to produce 50% C57BL/ 6N:50% FVB/N (mixed₅₀) offspring (Figure 5A). This breeding strategy was already performed previously and showed increased lifespan and more robustness when compared to pure C57BL6/N or FVB/N mice.²⁷ However, almost as expected, untreated mixed₅₀ SMA and SMA-Ncald^{ko/wt} mice live 16.5 and 17.0 days, respectively, showing that the modifier alone is still unable to counteract the massive loss of SMN (Figure 5B). Therefore, mixed₅₀ offspring were injected s.c. with a single suboptimal dose (30 µg) of SMN-ASO on P1. Elevated SMN levels were obtained in liver, but not in spinal cord or brain (Figure S6A). Survival of SMA+ASO mice was rescued (Figure 5B), but their motoric abilities were visibly impaired as determined by righting reflex and grip strength tests (Figures 5C and 5D). This suggests that slightly elevated SMN levels achieved by systemic SMN-ASO treatment rescued non-neuronal multi-organ impairment,²⁹ but not MN function. In contrast, heterozygous

⁽B) Representative images and quantification of NMJ area (μ m²) in TVA muscle from P10 mice stained with antibodies against NF-M and SV2 (green, for presynaptic terminals) and Bungarotoxin (magenta, for postsynapse). NMJ area was analyzed with ImageJ software (N = 3, n = 100–120 NMJs/mouse). ***p \leq 0.001. Scale bars represent 10 μ m.

⁽C) Representative images and quantification of proprioceptive inputs (VGLUT1, green) on MN soma (CHAT, magenta) in lumbar spinal cord sections from P10 mice. Mean input number within 5 μ m of MN soma was analyzed (N = 3, n = 100–120 MNs/mouse). ***p \leq 0.001. Scale bars represent 25 μ m. Note, color code for genotypes is identical to (D).

⁽D) Representative merged images of 6 DIV MNs isolated from E13.5 embryos and stained with DAPI (blue, for DNA) and antibodies against HB9 (green, for MN) and Tau (red, for axon). The longest axon and axonal branches were quantified with ImageJ (N = 3–5, n = 20–40 axons per mouse). Scale bars represent 25 μ m. Each boxplot covers values from 25%–75% with line at median and dotted outliers at <5% and >95% CI. For each experiment, image analysis was double-blinded. n.s. indicates non-significant; *p \leq 0.05; **p \leq 0.01; ***p \leq 0.001.

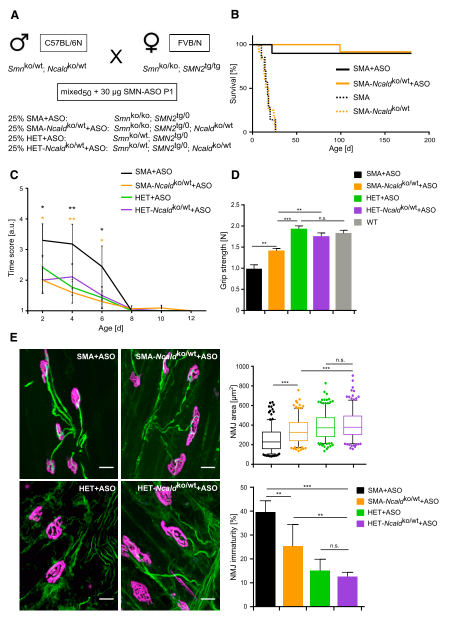


Figure 5. NCALD Reduction Improves Motoric Function, NMJ Size, and NMJ Architecture in SMA+ASO Mice

(A) Breeding scheme to produce mixed₅₀ SMA and HET mice. All mixed₅₀ offspring were injected with 30 μ g SMN-ASO at P1. (B) Kaplan-Meier curves of uninjected mixed₅₀ mice show no differences in survival between SMA (17 days, N = 7) and SMA-*Ncald*^{ko/wt} (16.5 days, N = 12). Injection of 30 μ g SMN-ASO on P1 increases survival to >180 days for both SMA+ASO (N = 10) and SMA-*Ncald*^{ko/wt}+ASO (N = 12) mice.

(C) Righting reflex test shows improvement in SMA-*Ncald*^{ko/wt}+ASO but not SMA+ASO mice during P2–P6 ($n \ge 12$ per genotype). Error bars represent SEM; n.s. indicates non-significant, ** $p \le 0.01$, *** $p \le 0.001$.

(D) Grip strength test performance at P73 reveals enhanced strength for SMA-*Ncald*^{ko/wt}+ASO mice compared to SMA+ASO mice (N \geq 12 per genotype). Error bars indicate SEM. *p \leq 0.05, **p \leq 0.01, ***p \leq 0.001.

(E) Representative images of NMJs of ASOtreated mixed₅₀ mice at P21 stained with the antibody against NF-M (green, for presynaptic terminal) and Bungarotoxin (magenta, for postsynaptic terminal). Scale bars represent 20 μ m. Boxplot shows quantification of NMJ area in μ ^{m2} in TVA muscle which was analyzed and represented as in Figure 4. Bar graph shows percentage of immature NMJs in TVA muscle (mean \pm SD). N = 3 mice per genotype; n = 60– 100 NMJs per mouse. n.s. indicates nonsignificant, *p \leq 0.05, **p \leq 0.01, ***p \leq 0.001.

late voltage-dependent Ca²⁺ currents (I_{Ca}) in MN-like cells. We performed whole-cell patch-clamp recordings and ratiometric Ca²⁺ imaging with fura-2. We recorded I_{Ca} of RA-differen-

Ncald knockout, in addition to low-dose SMN-ASO treatment, significantly improved motoric abilities (Figures 5C and 5D). Analysis of NMJ maturation score on P21⁴⁵ showed that both NMJ size and maturation were markedly restored by *Ncald* reduction as compared to SMA+ASO mice (Figure 5E). Heterozygous *Ncald* knockout did not rescue tail necrosis and slightly impacted weight progression in male mice (Figures S6B–S6D). Our data provide conclusive evidence of the beneficial effect of reduced NCALD on the neuromuscular system and motoric function in SMA+ASO mice.

Low SMN Decreases Ca²⁺ Influx in NSC34 and PC12 Cells

Since NCALD is a neuronal Ca²⁺ sensor and impaired Ca²⁺ homeostasis has been reported in SMA,⁶⁹ we tested whether lowering SMN and NCALD levels could modu-

tiated NSC34 cells that were treated with siRNAs specific to *Smn*, *Ncald*, or *Smn*+*Ncald* and analyzed the I_{Ca} tail currents with a series of increasing voltage pulses. In NSC34 cells, *Smn* depletion significantly reduced the voltage-dependent Ca²⁺ influx, which was not restored by additional NCALD reduction (Figure 6A). Ratiometric Ca²⁺ imaging with fura-2 revealed a reduced voltage-dependent Ca²⁺ influx in SMN-depleted PC12 cells compared to controls (Figure S7A). These data show that low SMN levels impair Ca²⁺ influx, which is not restored by NCALD knockdown and that NCALD depletion rescues synaptic transmission through a different mechanism.

Disturbed Endocytosis and Synaptic Vesicle Recycling Is Ameliorated by NCALD Depletion

We next sought for a common pathway in which both SMA modifiers, NCALD and PLS3, might operate. Since

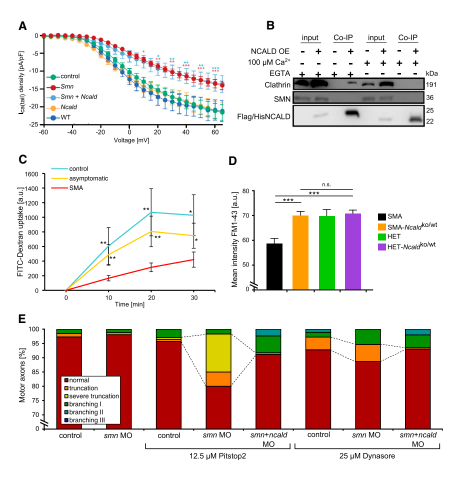


Figure 6. Interconnection between SMN, NCALD, Voltage-Dependent Ca²⁺ Influx, Endocytosis, and SMA

(A) Measurement of *I-V* relations of Ca²⁺ tail currents in differentiated NSC34 cells treated with respective siRNAs and depolarized for 5 ms to 60 mV, in 5 mV increments, at holding potential -80 mV. Currents were not different between wild-type (n = 7), control siRNA (n = 33), and *Ncald* KD (n = 13) and were significantly reduced upon *Smm* KD (n = 15) and *Smn+Ncald* KD (n = 12) at current pulses above -35 mV. *p \leq 0.05; **p \leq 0.01; ***p \leq 0.001. Error bars indicate SEM.

(B) Western blot of co-immunoprecipitation experiment. NSC34 cells were transiently transfected with FLAG-His-NCALD or control vector. Co-immunoprecipitations with FLAG-M2 affinity beads were performed in the presence or absence of Ca^{2+} . NCALD interacts with clathrin only in the absence of Ca^{2+} (addition of EGTA to the cell lysate) but not in the presence. Note the positive clathrin band in the test-Co-IP (fourth lane) in the absence but not in the presence of Ca^{2+} (last lane).

(C) Quantification of endocytosis by FITCdextran uptake in fibroblasts from SMA (n = 10), controls (n = 3), and asymptomatic individuals (n = 5); n = 50 per cell line and time point. Mean \pm SD. *p \leq 0.05, **p \leq 0.01.

(D) Quantification of FM1-43 intensity at presynaptic terminals in TVA muscles

under low-frequency stimulation (5 Hz, 1 s). n = 3 per genotype, $n \approx 100$ per mouse. Mean \pm SEM; n.s. indicates non-significant; *** $p \leq 0.001$.

(E) Quantification of MN axon phenotype of zebrafish embryos treated with sub-phenotypical doses of *smn* MO (2 ng), *ncald* MO (2 ng), and the endocytosis inhibitors Pitstop2 and Dynasore, respectively. Dashed lines highlight the synergistic effect of *smn* MO and Pitstop2 and the effect of Dynasore on axon truncation. Additional *ncald* MO injection ameliorates the truncation defect. ***p ≤ 0.001 . Motor axons per treatment: Pitstop2: n ≥ 100 , Dynasore: n ≥ 150 .

NCALD binds clathrin directly⁵⁸ and PLS3 knockout in yeast impairs endocytosis,^{58,70} we hypothesized that low SMN levels may impair endocytosis, which in turn is rescued by reduced NCALD or increased PLS3 levels. Indeed, we recently reported impaired endocytosis as a disturbed cellular mechanism affected in SMA, which is rescued by elevated PLS3 levels.²⁹ Impaired endocytosis and endocytic trafficking have further been demonstrated in a *C. elegans* SMA model.⁷¹

Co-immunoprecipitation studies in NSC34 cells revealed NCALD interaction with clathrin only in the absence of Ca^{2+} (Figure 6B) or at low Ca^{2+} levels (data not shown). TEM analyses after immunogold staining of wild-type zebrafish sections showed co-localization of Ncald and clathrin in the presynaptic sites of NMJs (Figure S7B).

To study the effect of NCALD on endocytosis, we undertook FITC-dextran internalization assays in various cell culture systems. In primary fibroblast cell lines derived from SMA-affected individuals, endocytosis rates were strongly reduced compared to controls but were restored in fibroblasts of asymptomatic individuals (Figures 6C and S7C). Moreover, *Smn* knockdown in NSC34 cells significantly reduced FITC-dextran uptake, which was rescued by concomitant *Ncald* knockdown. *Ncald* knockdown alone increased the rate of endocytosis by 1.3-fold, demonstrating that low NCALD levels already facilitate endocytosis (Figure S7F).

Moreover, we analyzed endocytic uptake of FM1-43 in mouse NMJs under stimulation at 5 and 20 Hz as described.²⁹ FM1-43 uptake was markedly decreased in SMA mice at 5 Hz stimulation (triggering clathrin-dependent endocytosis), but heterozygous *Ncald* knockout fully restored the levels similar to HET mice (Figures 6D and S7D). Heterozygous *Ncald* knockout had no impact at 20 Hz stimulation (triggering bulk endocytosis), further strengthening the specific role of NCALD in the clathrin-dependent endocytosis at the NMJ (Figure S7E).

Lastly, we investigated in vivo the mutual effect of endocytosis and the Smn-Ncald-clathrin network for SMA using pharmacological inhibition of endocytosis in zebrafish. Using sub-phenotypical concentrations of either *smn* MO (2 ng) or a suboptimal dose of Pitstop2 (12.5 μ M), an inhibitor of clathrin,⁷² showed almost no axon truncation and branching phenotype as compared to higher concentrations of smn MO (4 ng, Figures 3A and 3C) or Pitstop2 (25 µM, Figure S7G). Instead, combination of suboptimal smn MO (2 ng) together with suboptimal Pitstop2 (12.5 µM) resulted in severe motor axons truncation, suggesting a synergistic effect. Notably, this SMA phenotype was strongly ameliorated by additional Ncald reduction (Figures 6E, 3A, and 3C). Moreover, the treatment with Dynasore (25 µM), an inhibitor of the endocytosis-driving GTPase dynamin,⁷³ either alone or in combination with low smn MO, resulted in an SMA-like axonal truncation (Figure 6E). These defects were ameliorated by additional treatment with ncald MO (Figures 6E and S7G). Together, these findings suggest that SMN and clathrin interact genetically to promote endocytosis and MN axonogenesis, whereas NCALD negatively interferes with an SMN-dependent function of clathrin.

Discussion

Here, we describe NCALD as a genetic SMA modifier in humans. In summary, we show the following. (1) Reduced NCALD levels protect individuals from developing SMA, despite lacking SMN1 and carrying only four SMN2 copies, usually causing type 3 SMA.¹³ Thus, unlike PLS3, which alleviates SMA pathology upon overexpression,²⁴ NCALD reduction acts as a genetic suppressor of SMA. (2) NCALD is localized at SMA relevant sites including MN soma and growth cones as well as the presynaptic site of the NMJ. Furthermore, NCALD knockdown is relevant for MN differentiation and restores neurite and axon outgrowth in MNs or MN-like cells. (3) NCALD has a Ca²⁺-dependent interaction with clathrin and is thereby able to modulate endocytosis and likely vesicle recycling at the motor endplate. (4) NCALD knockdown rescues neural circuit function of zebrafish smn morphants by restoring axonal outgrowth defects, endplate potentials, and swimming velocity. (5) ncs-1 knockdown in smn-1-deficient C. elegans restores pumping to normal rates. (6) Heterozygous Ncald knockout in severely or intermediately affected SMA mice causes clear improvements on the structural level, such as NMJ size and architecture, MN outgrowth, and proprioceptive inputs. (7) Heterozygous Ncald knockout in mild SMA mice with no lifespan impairment has beneficial effects on NMJ size and architecture as well as motoric abilities. (8) Finally, across species, the mechanism by which reduced NCALD level improves SMA pathology is restoration of endocytic function, strengthening the existing models holding endocytosis as a main impaired cellular mechanism in SMA.

NCALD Downregulation as a Potential Therapy in Combinatorial Approach

Clinical trials using ASOs to correct *SMN2* splicing are highly promising and close to FDA approval.² However,

for type 1 SMA-affected children with only two SMN2 copies, these approaches are likely insufficient to fully suppress SMA symptoms. It is also unclear to what extent the elevation of SMN after disease onset will be able to protect from SMA and whether combinatorial therapies including SMN-dependent and SMN-independent pathways will be required to achieve full and long-term rescue.⁷⁴ There is increasing evidence-at least in mouse models-that systemic SMN elevation is required to fully counteract SMA; systemic injection of SMN-ASOs or AAV9-SMN led to a robust survival increase in various SMA mouse models in comparison to a central nervous system (CNS)-restricted application.^{68,75} This is in line with the observation that additional non-neuronal organs and tissues are impaired in severe SMA mouse models and partially in type 1 SMA-affected individuals (reviewed in Hamilton and Gillingwater¹⁸ and Shababi et al.⁷⁶).

Recently, we have shown that PLS3 overexpression in combination with low SMN elevation using SMN-ASOs⁶⁸ increased the survival of severely affected SMA mice from 14 to >250 days.²⁹ This might resemble a hypothetic situation in which individuals with type 1 SMA are treated with a molecule or drug that increases SMN levels acting on the endogenous SMN2 copies in combination with an additional molecule or drug acting on the genetic modifier. In contrast to PLS3, the effect of NCALD was less pronounced, which is in line with the observation that asymptomatic individuals protected by reduced NCALD require the presence of four SMN2 copies, while in case of elevated PLS3, three SMN2 copies are sufficient.^{24,25} In addition, the limited effect of NCALD might be due to the restricted expression in neuronal tissues as compared to PLS3, which is ubiquitously present. Moreover, the broader impact of PLS3 on F-actin dynamics that influences various cellular processes at NMJ level^{27,29} may further contribute to the more prominent protection.

Nonetheless, a combinatorial therapy that both elevates SMN and decreases NCALD (e.g., by ASO treatment) may provide a full protection, resulting in asymptomatic individuals. The advantage of NCALD, in comparison to PLS3, is that suppression of gene function is, in general, easier to achieve than its activation.

NCALD Suppression Restores Endocytosis and Synaptic Vesicle Recycling in SMA

To allow rapid and repeated rounds of neurotransmission at the synaptic endplate, synaptic vesicle recycling is essential.⁷⁷ In brief, after Ca²⁺-dependent exocytosis and release of acetylcholine (ACh) into the synaptic cleft, the synaptic membrane has to be retrieved rapidly via endocytosis. Then, retrieved vesicular membranes need to be transformed into synaptic vesicles, which are refilled with ACh. These are eventually transported to the readily releasable pool near active zones.⁷⁸ Despite the robust fail-safe factor in the motor neuron endplate potential, disturbances in the presynaptic vesicle cycle can severely impact neurotransmission. In SMA, impaired neurotransmission,

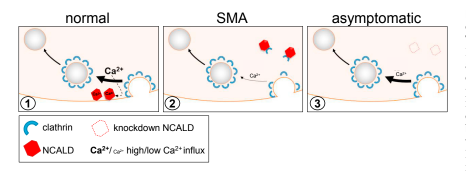


Figure 7. NCALD Acts as a Ca²⁺-Dependent Regulator of Endocytosis in Synaptic Vesicle Recycling

Diagrammatic presentation of the mode of action of NCALD in synaptic vesicle recycling in normal, SMA, and asymptomatic pre-synapse of neuronal cells. From left to right: (1) after neurotransmitter release, clathrin binds to empty vesicle membrane causing membrane bending and vesicle formation. High concentration of local Ca^{2+} which is present after vesicle release⁸⁵ causes NCALD conformational change and thereby a release of clathrin

so that it can perform its function. NCALD may fine-tune recycling speed and help to coordinate proper clathrin coating. (2) In SMA, voltage-dependent Ca^{2+} influx is reduced, decreasing NCALD-clathrin dissociation, thus inhibiting clathrin coating of vesicles. In our model NCALD regulates (increases) the Ca^{2+} dependence of clathrin function. (3) When NCALD level is reduced, the Ca^{2+} dependence is reduced too and even at relatively low intracellular Ca^{2+} levels, clathrin can mediate endocytosis.

disturbed Ca²⁺ homeostasis, decreased synaptic vesicle number, and reduced F-actin caging of reserve pool synaptic vesicles have been reported.^{16,69,79,80} For repeated neurotransmitter release, subsequent endocytosis is important;⁸¹ furthermore, endo- and exocytosis are regulated by the Ca²⁺ dynamics within the presynaptic terminals.⁸²

We found that low SMN levels cause reduction of voltage-activated Ca²⁺ influx, in accordance with recent studies in a zebrafish SMA model and reported mislocalization of calcium channels in SMA.^{83,84} However, unlike SMA pathology, Ca²⁺ influx was not restored by reduced NCALD, suggesting a different counteraction mechanism. Since NCALD interacts with clathrin and actin, two major players in endocytosis, 58,59 we hypothesized that reduced SMN may disturb endocytosis and synaptic vesicle recycling, possibly via decreased Ca²⁺, whereas NCALD knockdown subsequently compensates for SMN loss. We demonstrate in vitro and in ex vivo mouse NMJs that NCALD reduction restores impaired clathrin-dependent endocytosis. Furthermore, chemical endocytosis inhibition in zebrafish caused MN axonogenesis defects that were reversed upon Ncald suppression. Importantly, NCALD binds clathrin only at low Ca²⁺ levels (mimicking unstimulated MNs) but not at high Ca²⁺ levels (mimicking action potentials in MNs). For SMA MNs, with low Ca²⁺ levels even during action potential, we predict that NCALD constantly binds clathrin, thereby inhibiting/reducing its function in synaptic vesicle recycling. However, low NCALD levels, as in asymptomatic individuals, may allow free clathrin to act in endocytosis and synaptic vesicle recycling even at reduced Ca^{2+} levels (Figure 7).

Implication of NCALD in Other Neurodegenerative Disorders

In agreement with this hypothesis, two other proteins connected to endocytosis cause various forms of SMA. Mutations in *UBA1* (MIM: 314370), an E1 Ubiquitin-Activating Enzyme involved in monoubiquitination which serves as a signal for endocytosis and trafficking of cell surface proteins, has been associated to X-linked SMA (SMAX2 [MIM: 301830]).^{86–88} *BICD2* (MIM: 609797), which when mutated causes autosomal-dominant lower-extremity-predominant spinal muscular atrophy-2 (SMALED2 [MIM: 615290]), binds to clathrin heavy chain to promote its transport and augments synaptic vesicle recycling.⁸⁹⁻⁹³ These findings provide additional evidence that disturbances in synaptic vesicle recycling underlie general SMA pathology. Our findings are further strongly supported by data in C. elegans, in which disturbed endocytic trafficking at the synaptic level has been reported, and it has been suggested that an increased resistance against infection may explain the high SMA carrier frequency in the population.⁷¹ Moreover, reduced NCALD amount might be beneficial for other MN or neurodegenerative disorders with impaired endocytosis and Ca²⁺ homeostasis, as was shown for Alzheimer disease (AD [MIM: 104300]), where NCALD is highly upregulated,⁹⁴ or Parkinson disease (PD [MIM: 168600]), hereditary spastic paraplegia (MIM: PS303350), and ALS (MIM: PS105400), where impaired endocytic trafficking was found.⁹⁵ Therefore, it is tempting to speculate that NCALD downregulation might become an efficient strategy against SMA and other neurodegenerative diseases.

Accession Numbers

All microarray data are available in GEO: GSE58316.

Supplemental Data

Supplemental Data include seven figures, three tables, and clinical description of Utah family members and can be found with this article online at http://dx.doi.org/10.1016/j.ajhg.2017.01.005.

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Web Resources

1000 Genomes, http://www.internationalgenome.org/

Bioconductor, http://www.bioconductor.org

ClinicalTrials.gov, http://clinicaltrials.gov

Cyflogic, http://www.cyflogic.com

ENCODE, https://www.encodeproject.org/

GEO, http://www.ncbi.nlm.nih.gov/geo/

ImageJ, http://rsbweb.nih.gov/ij/

International Mouse Phenotyping Consortium, http://www. mousephenotype.org/data/genes/

LDlink, https://dceg.cancer.gov/tools/analysis/ldlink

Patcher's PowerTools plug-in (WaveMetrics), http://www3. mpibpc.mpg.de/groups/neher/index.php?page=software R statistical software, http://www.r-project.org/ RefSeq, http://www.ncbi.nlm.nih.gov/RefSeq

UCSC Genome Browser, http://genome.ucsc.edu

UniProt, http://www.uniprot.org/

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