Rescue of a severe mouse model for Spinal Muscular Atrophy by U7 snRNA-mediated splicing modulation

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In Spinal Muscular Atrophy (SMA), the leading genetic cause of early childhood death, the survival motor neuron 1 gene (SMN1) is deleted or inactivated. The nearly identical SMN2 gene has a silent mutation that impairs the utilisation of exon 7 and the production of functional protein. It has been hypothesised that therapies boosting SMN2 exon 7 inclusion might prevent or cure SMA. Exon 7 inclusion can be stimulated in cell culture by oligonucleotides or intracellularly expressed RNAs, but evidence for an in vivo improvement of SMA symptoms is lacking. Here we unambiguously confirm the above hypothesis by showing that a bifunctional U7 snRNA that stimulates exon 7 inclusion, when introduced by germ-line transgenesis, can efficiently complement the most severe mouse SMA model. These results are significant for the development of a somatic SMA therapy, but may also provide new means to study pathophysiological aspects of this devastating disease.

The hypothesis that boosting the inclusion of exon 7 in the SMN2 gene might provide an effective treatment or cure for SMA is based on the fact that all SMA patients have at least one, often several, SMN2 copies (1,2). This copy number inversely correlates with the severity of the disease (3,4). However, there also appear to be modulating effects from other genes or of non-genetic origins (5-7).

Exon 7 of the SMN2 gene is arguably one of the best studied exons of the human genome. It appears to be weak, as it is skipped, albeit infrequently, even in the functional SMN1 gene. This weakness is due to a suboptimal intron 6 branch point (8), a non-canonical 5’ splice site (ss), intronic splicing silencers (ISS) in both flanking introns and an inhibitory secondary structure towards the end of the exon (9). Positive elements are two exonic splicing enhancers, termed SE1 and SE2 which contain binding sites for the splicing factors ASF/SF2 and hTra2β, respectively. In the SMN2 gene, a silent mutation appears to weaken SE1 (10,11).
and may cause the formation of an additional hnRNP A1-dependent silencing element (12,13).

A stimulation of exon 7 inclusion has been achieved in cell culture models with antisense oligonucleotides that target either of the two ISS, the inhibitory sequences towards the end of exon 7, or, alternatively, mask the intron 7 branch point and 3' ss of exon 8 (9,14,15). In one of these studies, oligonucleotides were also introduced into a SMA mouse model, and splicing correction could be detected in liver and kidney. However, the oligonucleotides did not reach the spinal cord, and hence no therapeutic benefit could be demonstrated (15). Moreover, an elevation of SMN protein could recently be demonstrated after intracerebral injection of an oligonucleotide (16). Two studies, published in 2003, were using bifunctional oligonucleotides targeting the part of exon 7 that differs between SMN1 and SMN2 and additionally containing either an exonic splicing enhancer (ESE) sequence (17) or a peptide composed of alternating serines and arginines mimicking a SR protein (18). We have recently adapted several of these oligonucleotide-based approaches to an U7 snRNA-derived in vivo expression cassette. Additionally, we have tried to improve exon 7 inclusion by using modified U1 and U2 snRNAs targeting the 5' ss and the intron 6 branch point, respectively (8,19). As a conclusion from these studies, we arrived at an optimal exon 7 inclusion strategy based on a bifunctional U7 construct (termed U7-ESE-B; Fig. 1A) which targets the 3' part of exon 7 and carries an ESE sequence that can attract stimulatory splicing factors (19). This construct induced a nearly complete exon 7 inclusion of a SMN2-luciferase reporter in HeLa cells as well as of the endogenous SMN2 gene in SMA type I patient fibroblasts. In these fibroblasts, we also saw a 2- to 3-fold increase in SMN protein and, concomitantly, an increase of SMN-containing nuclear gems to a frequency also seen in certain wild-type cells or very mild forms of SMA (20). In the present study, we have introduced this U7-ESE-B cassette by transgenesis into the most severe SMA mouse model. We observe a clear
suppression of disease-associated symptoms which can, in the most efficient cases, allow a normal weight development, muscle performance and life expectancy. These results fully confirm that exon 7 is the target of choice for SMA therapies, although future studies will have to show whether a correction starting after birth can still be beneficial. In addition, the transgenic approach in SMA mouse models should also be helpful for further investigations into SMA pathophysiology.

**RESULTS**

**Therapeutic benefit of U7-ESE-B in a severe SMA model**

We used the U7-ESE-B construct in a transgenic approach to test the hypothesis that the stimulation of *SMN2* exon 7 inclusion could alleviate SMA symptoms. U7-ESE-B transgenic mice were generated by lentiviral vector (LV)-mediated transgenesis (21). Two founder mice were chosen, based on cytofluorometric analyses of blood samples for the intensity of the green fluorescent protein (GFP) contained in the LV (Fig. 1B), and crossed with carriers of a mouse SMA model (described below). The mean fluorescence intensities and the numbers of integrated LV copies (determined by quantitative real-time PCR of Gag and WPRE sequences) of these and further animals described below are documented in Supplementary Material, Table S1.

In the chosen SMA model, the most severe of its kind, the breeders are heterozygous for mouse *smn* (*smn+/-*) and homozygous for human *SMN2* (*hSMN2+/+*) and therefore have a similar genotype as human SMA carriers (22). The SMA-affected pups emerging from these breedings (*smn-/-; hSMN2+/+*) have an average lifespan of 5.2 ± 0.2 days (23). In their terminal stage, they have the weight and appearance of 2-3 day old normal pups.

When crossing the U7-ESE-B transgenic mice with SMA breeders, we found that both founder animals transmitted the GFP and U7 genes through the germline. GFP mean
fluorescence intensities and integrated gene copy numbers decreased, on average, by a factor of 2 for each generation (Supplementary Material, Table S1), which suggested that the multiple integrations were not clustered. In successive crossings of the first generation offspring with SMA breeders, we obtained mice with the desired genotype: \textit{smn-/-}; \textit{hSMN2+/-}; \textit{U7-ESE-B+} (variable copy numbers); these will henceforth be called SMA/U7 animals. For 15 of these SMA/U7 animals, a survival curve was established (Fig. 2). Copy numbers of the LV carrying the U7-ESE-B cassette varied between 1 and 12.5 in these animals (median 1.7). Six of these animals were censored (three for removal of tissues on day 32, one having died of an unrelated cause on day 275 and two mice remaining alive beyond 300 days). The other SMA/U7 animals died or were euthanised because of severe SMA symptoms between days 18 and 290. The median survival time was 123 days. In comparison, SMA mice (eleven animals, seven censored because they were euthanised on days 3 or 4 based on genotyping) had a median survival time of 6.5 days. This is comparable to the average lifespan reported for such mice (23) (see above). Most importantly, the ~20-fold increase in survival times of SMA/U7 compared to U7 mice is highly significant (P<0.0001).

Two litters from U7-ESE-B bearing mothers with ~15 and ~7 LV copies per diploid genome which had 8 and 9 pups, respectively, were compared in more detail for their weight development during the first two months and for their muscle performance. Each of these litters consisted of 3 SMA/U7 pups, 3 \textit{smn++} pups (termed wild-type) and 2 or 3 \textit{smn+-} pups (termed heterozygous), respectively (see Supplementary Material, Table S1 for more complete genotype description). Weights, general appearance, motility and additional disease-associated symptoms were initially recorded twice daily and later at wider intervals. Fig. 3 shows the weight development of these animals for the first 60 days after birth.

The three SMA/U7 pups (568, 573, 575) born from mother 523 with the higher U7-ESE-B copy number had a nearly normal weight development, compared to their littermates (Fig.
They contained between 5 and 12 LV insertions in their genomes (Supplementary Material, Table S1; Fig. 3C). They never showed any SMA-related disease symptoms (one of them died of an ear infection on day 275). One of these female SMA/U7 animals, 575, was bred with a wild-type male to see if it was able to carry, deliver and feed pups normally. Out of this coupling, seven pups were born which all developed normally beyond the time of weaning. These findings clearly show that the U7-ESE-B cassette can improve and sometimes even fully suppress the severe SMA symptoms.

In the litter of mother 558 with the lower U7-ESE-B copy number, two SMA/U7 pups showed a growth retardation after days 3-5, and their weight began to decrease at days 13 and 17, respectively (Fig. 3B). Based on criteria established in agreement with the animal experimentation authorities, these mice were euthanised on days 18 and 20, respectively. The other SMA/U7 pup from the same litter (580) had a reduced weight development and performed very weakly in a grid holding test (see below). It also displayed necrosis of the tail and of two toes on each hind foot. However, the necrosis stopped after some time, and the skin healed at these sites (Supplementary Material, Fig. S1). Such symptoms have been described previously for milder SMA mouse models (24). Importantly, mouse 580 did not display severe general disease symptoms, was always moving and eating well, until it died on day 290, apparently of a sudden SMA progression.

Grid holding tests (see Materials and Methods) performed on days 35, 42 and 56 after birth indicated that the SMA/U7 animal 580 displaying disease symptoms was able to hold suspended on the grid for 6-30 sec in the first, for 1-4 sec in the second and not at all in the third test series (Fig. 4A, column 7). The symptom-free SMA/U7 animals 573 and 575 showed a completely normal holding time (columns 3 and 4). Only animal 568 had a reduced holding time in all tests (column 2), but seemed to improve with each new series (not shown). Note that a reduced performance was also observed in all three series for one of the
heterozygous animals (581, data not shown but included in the wt/hz values; column 1). Thus, a reduced holding time may sometimes reflect an unwillingness to collaborate rather than a muscle weakness. Importantly, when the data of the three SMA/U7 animals from this litter were pooled, the median corresponded to the maximal length of the measurements, 120 sec (column 5).

The SMA/U7 animals 568, 573 and 575 and their wild-type and heterozygous littermates were subjected to Rotarod running tests on days 88 and 105-107 after birth (summarised in Fig. 4B). It is evident from the graph that the SMA/U7 animal 573 performed considerably more weakly than the other two SMA/U7 animals and most of the wild-type or heterozygous mice from the same litter. However, the wild-type animal 572 also had a strongly reduced performance. The performances of the SMA/U7 animals 568 and 575 were only slightly reduced compared to those of the other wild-type or heterozygous littermates.

Taken together, these results demonstrate that a phenotypic improvement is observable in all SMA/U7 animals, but that it is variable, ranging from a short increase in life span to a virtually full suppression of all symptoms (e.g. in the SMA/U7 mouse 575).

**Molecular and histological characterisation of SMA/U7 mice**

To determine how efficiently the U7-ESE-B snRNA corrects hSMN2 splicing, we analysed total spinal cord RNA by reverse transcription (RT)-PCR. As the primers were not complementary to mouse smn; RT-PCR bands were only obtained from mice containing the human SMN2 gene, but not from smn +/+ mouse lacking the human gene or from the mouse neuroblastoma cell line NB2A (Fig. 5A). To analyse the splicing correction, we used spinal cord RNA from 6-day old wild-type, heterozygous and SMA mice with various numbers of integrated U7-ESE-B vector copies and from a 3-day old SMA mouse lacking U7-ESE-B. As expected, in the spinal cord of the SMA mouse lacking a U7-ESE-B insert, only a small proportion of hSMN2 mRNA contained exon 7 (26%; Fig. 5B). In contrast, all the mice
containing the U7-ESE-B vector - wild-type, heterozygotes and SMA/U7 - had an improved hSMN2 exon 7 inclusion. Although there was no strict correlation, mice with higher U7-ESE-B copy numbers generally showed a stronger splicing correction. An improvement of exon 7 inclusion was also observed in RNA samples from other tissues (data not shown).

We then analysed the level of SMN protein in spinal cord extracts by Western blotting (Fig. 6). All analysed animals were 3 days old, and different amounts of extract were loaded on the gel to facilitate a semi-quantitative assessment. Accordingly, high levels of SMN were detected in wild-type (Fig. 6) and heterozygous animals (data not shown) which expressed mouse smn and hSMN2 that was partly corrected. For the SMA mouse that expressed only uncorrected hSMN2, the intensity of the SMN band in 10 µg of total protein was fainter than that seen in 1.25 µg of wild-type extract. Thus it appears to contain less than one tenth the amount of SMN in its spinal cord than a wild-type mouse. In contrast, the SMA/U7 mouse contained ~1/5 the amount of SMN of the wild-type mouse but clearly more than the SMA animal without U7-ESE-B cassette. Similar results were obtained with other mice (data not shown). In conclusion, the presence of the U7-ESE-B allows for a clear but moderate increase in SMN protein level in the spinal cord.

To assess the preservation of motoneurons, 31 day old animals from a single litter composed of three wild-type, heterozygous and SMA/U7 animals each were analysed. Spinal cord cross sections sampled to reflect most of the body axis were stained with Toluidine blue (Fig. 7A). The number of large neuronal cells per ventral horn (considered to be motoneurons) showed a broad variation (data not shown). Since the sampling covered most of the length of the spinal cords and similar numbers of ventral horn sections (50-61) were counted in all animals, average values were used for comparison. This quantitative analysis revealed no significant difference between SMA/U7 animals and their wild-type and heterozygous littermates (Fig. 7B). We also isolated and stained spinal cords from 3 day-old
SMA animals. However, although motoneurons were clearly visible (Fig. 7A), in our hands a reliable quantitative analysis was not possible due to the relatively poor tissue preservation and small size. It has been reported, though, that affected mice surviving up to day 5 show a loss of 35% of motor neurons in the spinal cord and 40% in the facial nucleus (22).

Although the Western blot analysis had indicated a partial restoration of SMN protein expression in the spinal cord, it was important to know whether this was also true at the level of motoneurons. We therefore stained spinal cord sections from the same animals used for the histological analysis with anti-SMN antibody and used a DAPI counterstain to identify cell nuclei. Neuronal nuclei can be distinguished from those of other (mostly glial) cells based on their larger size and patchier DAPI staining (Fig. 7C). The cytoplasm of motoneurons from heterozygous (Fig. 7C, top row) and wild-type animals (not shown) gets intensely stained for SMN, but the nuclear space remains void of staining except for occasional gems. In the SMA animal (middle row) the cytoplasm of motoneurons showed virtually no staining. Staining of motoneurons was clearly recovered in spinal cord sections from SMA/U7 animals (bottom row), although it was weaker than in heterozygous or wild-type animals. The staining appeared to be somewhat more granular, but it is possible that a higher density of the same granules might suggest a more even distribution in the heterozygous or wild-type animals. Importantly, however, the staining also extended into axonal processes. Thus, these analyses show that SMN production is partly restored in SMA/U7 animals, not only at the total spinal cord level, but also within the motoneurons, the cells primarily affected in SMA.

DISCUSSION

This study unambiguously and fully validates the hypothesis that SMA symptoms may be reduced or even completely suppressed by treatments that improve the utilisation of SMN2 exon 7. The range of phenotypic improvement seen in the SMA/U7 mice analysed here goes
from short life prolongation or normal life span with pronounced SMA symptoms to full
weight development, muscular function and the ability of a female to carry to term and feed a
normal sized litter of pups.

Most of these animals contained one to seven copies of the U7-ESE-B transgene (except
for one animal having 12 copies). There was no strict correlation between the number of
integrated U7-ESE-B copies and the phenotype of the SMA/U7 pups, but animals with higher
copy numbers tended to show fewer and less severe symptoms. Exon 7 inclusion in total
spinal cord RNA was increased from 26 up to 52%, and SMN protein levels were also
increased, albeit only to an estimated 1/5 of the levels seen in wild-type or heterozygous
animals. That such a small increase in SMN levels may have a significant impact on viability
and SMA symptoms is in agreement with findings of Gavrilina et al. who showed that low
SMN cDNA expression in spinal cord can substantially prolong survival (25).

The fact that we saw only partial correction of SMN2 splicing and low levels of SMN
production could be traced to a very weak expression of the U7-ESE-B transgene which was
hardly detectable by primer extension (Supplementary Material, Fig. S2). We have recently
made the striking observation that different mouse U7 gene derivatives that we use for
splicing correction are highly expressed in human, but poorly in mouse, cells (J.M.,
unpublished results). The reasons for this phenomenon are still under investigation, but this
means that a higher expression might be expected in human patients, so that a single U7-ESE-
B transgene per cell might be sufficient for a phenotypic correction. We are also investigating
ways to improve the expression to achieve this goal.

These findings raise hopes for a therapy for SMA. As all patients carry at least one copy
of SMN2, they may all benefit from a treatment able to improve exon 7 inclusion, particularly
in motoneurons. However, there is still a long way to go, and it is presently unclear which
type of therapy - small drugs, oligonucleotides or the type of therapeutic gene used here - will
be first and most successful. Viewing its efficiency, our small U7-ESE-B gene is a good candidate, but its efficient and safe delivery to motoneurons will have to be established. Primary, but not exclusive, options are AAV vectors (26-28), but the serotype, strain and application routes must yet be elaborated.

Besides its importance in opening perspectives for SMA therapy, the transgenic approach that we have chosen also offers distinct possibilities to increase our knowledge about SMA in the mouse model. We have recently developed a doxycycline-inducible system for the regulation of therapeutic U7 cassettes (29). We now plan to introduce this system into SMA mice by transgenesis in order to address important questions, such as when during ontogenesis low levels of SMN are particularly critical, whether there is a point of no return after which a therapy can no longer rescue the motoneurons from dying, or whether the disease will still develop if a correction of SMN2 splicing is discontinued after the first phases of post-natal life. Additionally, mice can now be derived with stable single-copy insertions that should show intermediate SMA phenotypes due to partial SMN2 splicing correction. These could be used to decipher molecular events occurring in the affected motoneurons, e.g. through the use of splicing microarrays. This is particularly important, because recent studies suggest that the distribution of spliceosomal snRNPs and the splicing of many genes are deranged in a tissue-dependent manner, suggesting that SMA may be primarily a disease caused by splicing changes (30,31).

MATERIALS AND METHODS

Molecular cloning and lentiviral vector transgenesis

The previously described U7-ESE-B splicing correction cassette (19) was amplified by PCR with the primers U7-5end-XhoI and U7-3end-EcoRV with Pfu-Ultra polymerase (Stratagene, La Jolla, CA) according to the manufacturer's recommendations. For primer sequences see
**Supplementary Material, Table S2.** The *XhoI/EcoRV* fragment was then transferred into the third generation lentiviral vector pRRL-SIN-cPPT-hPGK-GFP-WPRE (32), between the cPPT and the hPGK promoter, in forward orientation with regard to eGFP. The entire U7 cassette and surrounding elements were verified by sequencing.

Lentiviral vectors were produced as described (33) except that supernatant was harvested twice (at 24 and 36 hrs post-transfection) and that serum-free Episerf medium (Gibco Invitrogen, Basel, Switzerland) was used instead of DMEM with fetal calf serum. Lentiviral vector transgenesis was carried out as described (21).

**Animal strains, breeding and husbandry**

The SMA mouse strain (22) was obtained from Jackson Laboratory (strain name: FVB.Cg-Tg(SMN2)89Ahmb *smn*<sup>1m1Msd</sup>/J, Stock number: 005024). These mice are heterozygous for a LacZ-Neo insertion in exon 2 of their *smn* gene and additionally carry two copies of the human *SMN2* transgene. Crosses between these animals result in progeny that are homozygous for the *smn* mutation and carry 2 copies of the *SMN2* transgene. Such animals have been reported to die between 4-7 days after birth (22,23). In contrast, homozygous *smn* knock-out mice without the *SMN2* transgene show embryonic lethality with an arrest in development before the stage of implantation (34).

The U7 transgene was inserted by lentiviral vector-based transgenesis into oocytes of B6D2F2 mice (work performed in the laboratory of D. Trono at the EPFL, Lausanne, Switzerland) (21). Female transgenic animals were then crossed with SMA carrier males. A second cross with SMA carriers yielded mice of the desired genotype (*smn*/-; *hSMN2*+/+; U7-ESE-B+ [multiple copies]), which are termed SMA/U7.

All mice were kept in the animal facility at the Insel hospital in Berne, Switzerland under IVC conditions in compliance with legal requirements for animal husbandry and with the specific authorisation issued for this project by the cantonal animal protection authority. In
particular, litters with expected SMA genotypes were initially looked after twice daily and weighed once a day. Some older mice no longer used for experiments were moved to a conventional animal facility.

**Cytofluorometric GFP detection in blood samples**

Approx. 50μl tail vein blood was mixed with 1 ml PBS containing 5μM EDTA and centrifuged for 5 min at 500g and 4°C. The pellet was gently resuspended in 4 ml of red blood cell lysis buffer (150mM NH₄Cl, 10mM KHCO₃, 0.1mM EDTA) and incubated for 10 min at room temperature in the dark. Cells were precipitated by centrifugation as above, and the red supernatant containing destroyed red blood cells was removed. Remaining cells were resuspended in 1ml PBS, and GFP expression was measured by cytofluorometry (FACS Calibur, BD Bioscience, Meyrin, Switzerland). Age-matched blood not expressing GFP was used to define a negative population. The geometric mean of fluorescence intensity was determined with the FlowJo software version 7.2.4.

**Genotyping**

Genomic DNA was isolated from tail biopsies, usually taken on the day of birth, by using the Wizard genomic DNA purification Kit (Promega AG, Wallisellen, Switzerland). For the PCR reactions, 20μl of reactants containing 50-200 ng genomic DNA, 400nM of the corresponding primers (see table below), 200μM of each deoxyribonucleotide triphosphate, 4μl of Q-solution and 1 U of Taq Polymerase (Qiagen AG, Hombrechtikon, Switzerland) were incubated for 30 s at 94°C, 1 min at 50°C, and 2 min at 72°C for 40 cycles.

For the smn genotype, one common forward primer (mSmn Fw) and two reverse primers (mSmn Re and Neo border) were used. Intact and mutated smn yield products of 594bp and 1200bp, respectively. In heterozygous animals, both bands are produced. To determine the SMN2 copy number, a common forward primer (Tg89-Grm7-Fw) was used, again together
with two distinct reverse primers (Tg89-Grm7-neg and Tg-89-Grm7-Re) (25). Thus, in heterozygous animals, two products of 280bp and 372bp were observed, representing the alleles with and without integration of SMN2, respectively. A qualitative assay for LV integration made use of the primer pWPTS-Sequencing and GFP10b and yielded an amplicon of 1406bp in the positive case. For gender determination, the primers sry 3’ and sry 5’ yielded a male-specific product of 600bp. For primer sequences see Supplementary Material, Table S2.

Real-time PCR

Starting from 70 ng of mouse genomic DNA, we amplified Gag and WPRE sequences in the lentiviral construct, and Titin, a mouse housekeeping gene (35) (see Supplementary Material, Table S2 for primers and probes). The absolute amount of each gene was determined by referring to a standard curve. This consisted of a 10-fold serial dilution of a plasmid containing one copy of each of the genes, with known copy numbers at each dilution (six concentrations ranging from 10⁶ to 10 copies for each gene). The absolute copy number of transgene integrated in the genome was obtained by dividing the amount of each reporter gene, Gag or WPRE, by half of the Titin amount, since the plasmid contains a single copy of the Titin gene, whereas the genomic DNA contains two copies. Values calculated with the Gag and WPRE assays correlated with a correlation coefficient \( R^2 = 0.9547 \), but WPRE-derived values were on average 20% higher. As it is not possible to say which assay is more correct, all values presented in Supplementary Material, Table S1 and in the main manuscript are averages between the two determinations.

The real-time PCR reactions were run in simplex with the ABI Prism 7000 SDS v1.1 system (Applied Biosystem, Foster City, CA, USA). Mice genomic DNA or serial dilutions of the plasmid were included in a mix containing 2x Absolute QPCR Rox mix buffer (Thermo Scientific, Waltham, MA, USA), 162 nM of specific primers, 800 nM of corresponding
TaqMan probe and water to a 18 μl final volume. The run conditions were as follow: 9600 emulation mode, one 15 min 95°C polymerase activation step, followed by 40 cycles of 2-step qPCR (15 sec of 95°C denaturation, 1 min of 60°C combined annealing/extension).

**Muscle performance tests**

*Grid holding test:* Mice were placed on a grid and allowed to get a firm hold. Then the grid was turned upside down for 2 min, and the time for which the animals were able to hang on to the grid was measured. Three such tests were performed during each session, whereby the animals were allowed to recover for 1 min between the measurements. These tests were carried out three times with one week intervals.

*Rotarod assay:* This assay was carried out in the animal facility of the University of Fribourg, Switzerland with the kind support and advices from J. Schwaller and P. Gregory. After initial training sessions on a RotaRod device (TSE Systems GmbH, Bad Homburg, Germany), the mice were tested with a protocol applying accelerating speed of 10-40 rpm for up to 4 min. Each mouse did six trials per day, three in the morning and three in the afternoon; the break between the individual trials was at least 10 min, and the break between morning and afternoon was 3 hours. The time that the animals were able to keep running was recorded. The data were analysed by using the Prism® software (GraphPad, San Diego, USA).

**RNA analysis**

Total RNA was extracted with homemade Tri-Reagent (29) and resuspended in diethyl pyrocarbonate-treated water. Two μg of total RNA were reverse-transcribed into first-strand cDNA, by using 200 nM oligo-dT, 50 U of Stratascript 6.0 reverse transcriptase (Stratagene), 1x Strata 6.0 buffer (Stratagene), 500 μM of all four dNTPs (Roche Diagnostics AG, Rotkreuz, Switzerland), 10 mM DTT and 8 U RNAsin (Promega) in a total volume of 50 μl.
PCR reactions including radioactive label were carried out with Taq DNA polymerase (Qiagen) in a volume of 25 µl with 5 µl of cDNA, 200 nM each of the SMN-Ex6-Fw (200 nM) and SMN-Ex8-Re primers, a mix containing 200 µM of dATP, dTTP, dGTP and 20 µM dCTP, 5µCi α-32P-dCTP (Hartmann Analytic GmbH, Braunschweig, Germany). Cycling conditions were 45 s at 95°C, 1 min at 50°C and 1 min at 72°C for 27 cycles. The PCR products were separated by electrophoresis on a 6% denaturing polyacrylamide gel. After exposure to phosphor storage screens (Molecular Dynamics, Sunnyvale, CA, USA), the intensity of radioactivity in the bands lacking (130 bp) or containing (184 bp) SMN2 exon 7 was estimated with the AIDA software (version 2.31, Raytest Isotopenmessgeräte GmbH, Straubenhardt, Germany) and values were corrected by normalising them for the number of Cs in the molecule. In one experiment the PCR was carried out in the absence of radioactive dCTP, but the forward primer was 5’ end-labelled with 6-carboxyfluorescein (CF). The PCR product (1.2 µl) was mixed with 12 µl of GS500 LIZ standard (12.5 µl diluted in 1 ml of Hi-Di formamide, Applied Biosystems) and run for fragment analysis on an ABI 3100 sequencer (Applied Biosystems). The relative amounts of the products lacking or containing SMN2 exon 7 were estimated from the areas of the corresponding peaks. The percentage of inclusion was calculated as the value of the included band divided by the sum of both products. The two methods yielded very similar results and the results were therefore averaged for inclusion in Fig. 5 of the paper.

For primer extension, 10 pmol of Pex-U7-Universal (detecting U7-ESE-B) or Pex-U2-WT (detecting endogenous U2 snRNA) primers were 5’-labelled with 10 U of T4 polynucleotide kinase (New England Biolabs, Ipswich, MA, USA) in the presence of 20 pmol of γ-32P-ATP (Hartmann Analytic), and the labelled primers were used without further purification. Five µg of total RNA were slowly hybridised in a thermocycler with a mix of 100 fmol of both primers, and the reverse transcription reaction was performed as described above, except that
the incubation was extended for 2 hours. The products were separated on 15% denaturing polyacrylamide gels and the signals were analysed as described (19).

**Western blot**

A polyclonal anti-SMN antibody was produced by immunising New Zealand white rabbits with a recombinant protein corresponding to the first 180 amino acids of mouse SMN. Antibodies were affinity-purified with the recombinant protein used for immunisation. Bound antibodies were eluted with 0.2 M glycine, pH 2.8, neutralised and stored at 4°C.

For Western blots, tissue samples were lysed in RIPA buffer (137 mM NaCl, 10 mM Phosphate, 2.7 mM KCl, 0.1% Sodium Dodecyl Sulfate, 0.5% Sodium Deoxycholate, 1% Igepal CA-630) supplemented with proteinase inhibitors (Complete EDTA-free, Roche). Extracts were resolved on 10 % high-TEMED SDS–polyacrylamide gels and proteins were transferred onto reinforced nitrocellulose membranes (0.45 μm, Schleicher & Schuell, Bottmingen, Switzerland). SMN was detected with a 1:500 dilution of affinity-purified rabbit anti-mSMN antibody in TBS-tween-milk (20 mM Tris, 135 mM NaCl, 0.1% Tween-20, 5% fat free milk powder), while actin was detected by using a 1:500 dilution of the anti-actin (20-33) antibody (Sigma-Aldrich Chemie GmbH, Buchs SG, Switzerland) in TBS-tween-milk. The primary antibodies were then detected with a species-specific antibody coupled to horseradish peroxidase (Promega), and signals were visualised by the enhanced chemiluminescence method (Amersham Biosciences) and digitalised with the Luminescent Image Analyser LAS-1000 plus (Fujifilm AG, Dielsdorf, Switzerland).

**Histology and immunofluorescence**

**Spinal cord sectioning, staining and neuron counting:** At post-natal day P31, spinal cords from the littermates 634-642 were isolated and cut close to the cervical end. From there on, 5 pieces of 5 mm length were embedded in M-1 Embedding Matrix for frozen sectioning (6
fluid OZ, Thermo Shandon, Pittsburgh) and kept frozen at -20°C until use. From each 5 mm part, 10 to 12 cryosections of 25 μm were cut with a Leitz 1720 Cryostat/Microtome (Leica Microsystems GmbH, Wetzlar, Germany), floating in PBS and then transferred to TBS containing 0.1% Triton-X-100 from where they were taken onto microscope slides (superfrost, Menzel-Gläser, Braunschweig, Germany) and left to dry at least 5 hours or overnight. Then the slides were fixed in 100% ethanol for 10 min and transferred to the Toluidine blue staining solution (1.2g Toluidine blue O and 3g Na2CO3 dissolved in 360ml doubly distilled H2O, 40ml of 70% ethanol and 400mg NaN3) for 6.5 min. Then, the slides were washed twice shortly in 70% ethanol and incubated for 1 min in 100% ethanol before putting them shortly in 100% xylol from where they were taken for embedding in Eukitt.

Before counting, the slides were blinded and assigned random numbers. All slides were then counted by a single person (that was not involved in the blinding) to minimise differences in counting criteria. A virtual line was drawn at the level of the central canal and all clearly visible large cells anterior to this line were considered to be neurons and counted. For each of the five regions of a spinal cord, at least 10 anterior horns were counted.

**Immunohistochemistry:** Cryosections from 31 day old littermates 634 (WT), 635 (HZ) and 642 (SMA-U7), as well as from the 3 day old mouse 504-4-4 (SMA, without therapeutic U7) were used. Floating sections were transferred from PBS into TBS. Then, the slides were incubated with TBS containing 1% Triton-X-100 for 30 min and washed 2x 10 min with TBS followed by a 2-day incubation at 4°C with the monoclonal anti-SMN antibody 2B1 (Sigma, 1:500 diluted in TBS containing 1% BSA, 0.1% NaN3). Then, the sections were washed 3x 10 min in TBS before incubation with the secondary antibody goat-anti-mouse IgG conjugated to Alexa-594 (Invitrogen, 1:1000 diluted in TBS containing 1% BSA) for 2 hours at room temperature in the dark. After washing 3x10 min with TBS, the samples were incubated with DAPI (Invitrogen, 5ng/μl in TBS containing 1% BSA) and washed again 3x10 min before
they were transferred to microscope slides and embedded with Glycergel (Dako Inc., Carpinteria, CA, USA).

Preparations were analysed on an Eclipse 600 microscope equipped with a DXM1200 camera (Nikon, Tokyo, Japan) or on a Laser Scan Microscope SP2, (Leica Microsystems). Image processing was performed with ImageJ 1.34s (National Institutes of Health, Bethesda, MD, USA) and Corel Graphics Suite software (Corel GmbH, Unterschleissheim, Germany).

SUPPLEMENTARY MATERIAL

Supplementary Material is available at HMG Online.

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Conflicts of Interest statement. None declared.

REFERENCES


**Figure legends**

**Figure 1** Structure of U7-ESE-B snRNA-containing lentiviral vector (LV) used for mouse transgenesis. (A) The U7-ESE-B splicing correction cassette contains an antisense sequence directed to the 3’ part of human \textit{SMN2} gene exon 7 and an additional splicing enhancer sequence (19). The modified U7 moiety is represented by its 3’-terminal stem loop and its Sm protein core (pictured as a grey ball). Within exon 7, the C to U transition that distinguishes \textit{SMN1} and \textit{SMN2} and that is thought to disrupt an ESE, the central ESE (SE2) and the UAA stop codon are indicated with white letters. (B) Structure of the LV pRRL-SIN-cPPT-hPGK-GFP-WPRE(32). The U7-ESE-B cassette is inserted upstream of the human phosphoglycerate kinase promoter (PGK) in sense orientation with respect to the green fluorescence protein (GFP) open reading frame. Typical for a third generation vector, the 5' LTR contains U3 sequences from Rous sarcoma substituting for those of HIV-1 (RSV-RU5). The 3'-LTR carries a self-inactivating (SIN) deletion. WPRE, woodchuck hepatitis virus posttranscriptional regulatory element; cPPT, HIV-1 central polyurine tract; \textit{\Psi}, HIV-1 packaging signal.
**Figure 2** The presence of the U7-ESE-B cassette prolongs survival in SMA mice. SMA mice (turquoise) are homozygous for an insertional inactivation of the *smn* gene and homozygous for a human *SMN2* transgene (22). SMA/U7 mice (purple) additionally contain between 1 and 12 copies (median 1.7) of a lentiviral vector carrying the U7-ESE-B cassette. Dashed lines indicate 95% confidence intervals. n, number of mice reaching endpoint (death or euthanasia due to severe SMA symptoms) or being censored, separated by a slash. Times when animals were censored are shown as upward blips. Reasons for censoring were: SMA, euthanasia after positive genotyping; SMA/U7, removal of tissues on day 32 (three animals), death of an unrelated cause (ear infection) on day 275 (one) and survival beyond the reporting period (two). The difference between the median survival times - 123 days for SMA/U7, 6.5 for SMA - is highly significant (P<0.0001)
Figure 3  Weight development of SMA mice containing the U7-ESE-B cassette. Two female mice (smn+/--; hSMN2+/--; containing multiple copies of U7-ESE-B) were bred with SMA carrier males and the weight of their pups was measured at frequent intervals. (A) Mouse 523 (~30 copies of U7-ESE-B per diploid genome) had 3 SMA/U7 pups with 2 hSMN2 copies, 3 wild-type and 2 heterozygous pups. (B) Mouse 558 (~7 U7-ESE-B copies) had 3 SMA/U7 pups with 2 hSMN2 copies, 3 wild-type and 2 heterozygous pups. The SMA/U7 pups 558-1-4 and 558-1-5 had to be euthanised on days 18 and 20, respectively. (C) Numbers of U7-ESE-B LV copies inserted in the genome of the above SMA/U7 mice. The real-time PCR indicated 0.5 copies in mouse 558-1-4, but non-quantitative PCR indicated that it contained at least one transgene copy. Animal identification: three digit numbers represent ear tag numbers introduced at weaning age. Animals killed at an earlier age are identified by the ID of their mother, the litter, and a numbering system for animals of the same litter, based on foot tattooing on the day of birth (for example, 558-1-4 is pup 4 from the first litter of mother 558).
Figure 4  Muscle performance of SMA mice containing the U7-ESE-B cassette. (A) On days 35, 42 and 56, the ability of the remaining mice from litters 523-1 and 558-1 to hang suspended under a metal grid for up to 120 sec was measured (3 tests per day and mouse). n, number of individual test values. (B) On days 88 and 105-107, the mice from litter 523-1 were tested for their ability to run for up to 240 sec on a Rotarod running device with the rotation speed increasing linearly from 10 to 40 rpm (6 tests per day and mouse, 24 values in total). The boxes (grey for SMA/U7 mice, white for wild-type or heterozygous animals) encompass all values in the 10-90 percentile range. A thick horizontal bar indicates the median. Because of the maximum cut-off time, the values do not fall into a normal (Gaussian) distribution.
Increased *SMN2* exon 7 inclusion in SMA mice containing the U7-ESE-B cassette. (A) Gel with controls showing that the reverse transcription (RT-)PCR assay detects transcripts of human *SMN2*, but not mouse *smn*. NB2A, RNA from mouse neuroblastoma cell line NB2A; noh*SMN2*, RNA from spinal cord of a mouse that does not contain the human *SMN2* gene; the last three lanes correspond to the identically labelled lanes in panel B; - RT, identical reaction as for the wt mouse, but with reverse transcriptase omitted; M, 100 bp size marker. The picture is a negative photograph of an ethidium bromide-stained agarose gel. The RT-PCR bands correspond to full-length *SMN2* mRNA (FL) and RNA lacking exon 7 (Δex7). (B) RT-PCR reactions carried out with total spinal cord RNA from various mice. Note that all animals in the black labelled lanes (including wild-type [wt] and heterozygous [hz] ones) contain the U7-ESE-B cassette and hence show a splicing correction compared to the SMA mouse (red labelled lane). The numbers of integrated U7-ESE-B vector copies, as determined by quantitative PCR, are indicated above the lanes. The picture shows a denaturing polyacrylamide gel of reactions carried out in the presence of $^{32}$P-labelled dCTP. The percent inclusion values were averaged from well correlating values of this gel scan and of a capillary electrophoretic analysis of an RT-PCR carried out with a fluorescently labelled downstream primer (see Materials and Methods). All samples were obtained at post-natal day 6, except for the SMA sample which was from a 3 day old animal. M, size marker (end-labelled *Hpa*I digest of pBR322).
**Figure 6** Increased SMN protein levels in SMA mice containing the U7-ESE-B cassette. Western blot detecting SMN and β-actin (loading control). The samples used were spinal cord lysates from 3 day old animals 520-4-9 (wt; 2.5 U7-ESE-B copies; see Supplementary Material, Table S1), 608-2-1 (SMA) and 520-4-2 (SMA/U7; 5.3 U7-ESE-B copies). The amounts of protein loaded on the gel are indicated below the lanes. The anti SMN antibody detects both human and mouse SMN. H, HeLa protein extract loaded as marker.
Figure 7 Preservation of motoneurons and SMN protein localisation in SMA mice containing the U7-ESE-B cassette. (A) Toluidine blue staining of spinal cord cross sections of the indicated mice. Mice 635 hz (heterozygote for smn), 638 wt (wild-type for smn) and 642 SMA/U7 were all from the same litter and 31 days of age; the SMA mouse 503-3-4 (from a separate litter) was 3 days old. (B) Counts of large neurons in spinal cord ventral horns of all nine animals (634-642) from the same litter. Five to seven sections from five different spinal cord areas separated about 5 mm from each other were used to count large neuronal cells in at least 50 ventral horns per animal. The average values from each animal were calculated, and the mean and standard deviation are shown for each group of animals (3 animals per group). For the three SMA/U7 animals, the numbers of integrated U7-ESE-B vector copies as determined by quantitative PCR were between 1.2 and 1.9 (see Supplementary Material, Table S1). (C) Immunofluorescence analysis of SMN protein in spinal cord sections. The panels show overviews (left) and close-ups (right) with SMN immunofluorescence in red (single confocal planes) and DAPI staining of nuclei in blue (stack overlays). Samples shown are from heterozygous (hz) and SMA/U7 animals and from the 3 day old SMA pup also used in (A). Staining patterns of wild-type animals were very similar to the ones shown for the heterozygous mouse. For the SMA
animal, the areas occupied by some of the motoneuron nuclei are indicated by white interrupted lines to illustrate that the surrounding cytoplasmic area shows virtually no SMN staining. The red background staining comes from the secondary antibody which primarily binds to blood capillaries; in contrast no spill over was observed between the DAPI staining and the red channel.