

Home-based Functional Electrical Stimulation for long-term denervated human muscle: History, basics, results and perspectives of the Vienna Rehabilitation Strategy

Helmut Kern (1,2), Ugo Carraro (1,3)

(1) Ludwig Boltzmann Institute of Electrical Stimulation and Physical Rehabilitation, Vienna;
(2) Department of Physical Medicine and Rehabilitation, Wilhelminenspital, Vienna, Austria;
(3) CIR-Myo Translation Myology Lab, Department of Biomedical Sciences, University of Padova, Italy

Abstract

We will here discuss the following points related to Home-based Functional Electrical Stimulation (h-b FES) as treatment for patients with permanently denervated muscles in their legs: 1. Upper (UMN) and lower motor neuron (LMN) damage to the lower spinal cord; 2. Muscle atrophy/hypertrophy versus processes of degeneration, regeneration, and recovery; 3. Recovery of twitch- and tetanic-contraction by h-b FES; 4. Clinical effects of h-b FES using the protocol of the “Vienna School”; 5. Limitations and perspectives. Arguments in favor of using the Vienna protocol include: 1. Increased muscle size in both legs; 2. Improved tetanic force production after 3-5 months of percutaneous stimulation using long stimulus pulses (> 100 msec) of high amplitude (> 80 mA), tolerated only in patients with no pain sensibility; 3. Histological and electron microscopic evidence that two years of h-b FES return muscle fibers to a state typical of two weeks denervated muscles with respect to atrophy, disrupted myofibrillar structure, and disorganized Excitation-Contraction Coupling (E-CC) structures; 4. The excitability never recovers to that typical of normal or reinnervated muscles where pulses less than 1 msec in duration and 25 mA in intensity excite axons and thereby muscle fibres. It is important to motivate these patients for chronic stimulation throughout life, preferably standing up against the load of the body weight rather than sitting. Only younger and low weight patients can expect to be able to stand-up and do some steps more or less independently. Some patients like to maintain the h-b FES training for decades. Limitations of the procedure are obvious, in part related to the use of multiple, large surface electrodes and the amount of time patients are willing to use for such muscle training.

Key Words: SCI, FES, skeletal muscle permanent long-term denervation, recovery of function, 2D and 3D Color TAC, biopsy, histology, electron microscopy

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The series of e-published issues of the European Journal of Translational Myology (EJTM) are starting in 2014 with the Specials on “The long-term denervated muscle” to resound “The denervated muscle”, a book edited in 1962 by one of our virtual mentors, Ernest Gutmann.¹

To explain why you are now reading this chapter, we need a tremendous amount of details that we cannot include in this review. Indeed, looking to his roots, one of us rediscovered a few months ago (emptying the office for retirement) that his M.D. Thesis was on muscle denervation.

Ugo Carraro: Pioneering studies

Prof. Carraro would like to remember that in the early 1960s he was a young student at the School of Medicine of the University of Padova, just admitted in 1964 to Internship of the Institute of General Pathology directed by Prof. Massimiliano Aloisi. When in Padua our full Professor almost every day took a tea cup with the fellows, discussing muscle research and his hope to develop *in vitro* muscle mimics, despite the difficulties to obtain motoneuron-myotubes cultures. Thus, he started lab training doing histology and discussing of the muscle and of its dependence from the motor

neuron. How to study this topic, if not by denervation experiments? Four years later he defended his M.D. Thesis on: "Impairments of the functions of substructural membranes of the denervated muscle (Alterazioni delle funzioni delle membrane substrutturali nel muscolo denervato)."²

Now, after 45 years he is trying to convince experts in aging that sparse, but incremental denervation is one of the many mechanisms that worsen muscle performances and quality of life of seniors, and that a long-term high-level physical activity may defer the unavoidable decay of aging.^{3,4} Mosole et al., indeed, comparing muscle biopsies from sedentary and very physically active seniors observed reduced numbers of denervated fibers and higher percentages of trophic and slow-type groups of reinnervated fibers in the active group. The observations suggest that long-term physical activity promotes reinnervation of muscle fibers undergoing age-related denervation.

Here he may only list the topics he would like to describe in a future book. He will need to start with his mentors (Aloisi, Zatti and Margreth), and describe the importance for his ability to design and perform independent research of his younger or older colleagues Catani, Mussini, Cantini, Salviati and Schiaffino. The explanation of why in Padua there was and there is such a strong tradition of Myology will end the first chapter. It will be a funny story related to fever and burning of toxins in the muscle ...

He will explain why he moved from General Pathology to Muscle Biology and Physiopathology, from Basic to Applied Myology, organizing the Interdepartmental Research Center of Myology of the University of Padova, in which clinical colleagues and biomedical scientists are almost equally present, from organizing the PaduaMuscleDays Meetings and editing the journal Basic and Applied Myology (BAM) to the European Journal of Translational Myology.

He will mention the inter-relationships among his students (Donatella Biral, Donatella Morale, Giorgio Vescovo, Corrado Rizzi, Gianluca Rigatelli, Marco Sandri, Marzena Podhorska-Okolov, Katia Rossini, Massimo Donà, Nicoletta Adami, Sandra Zampieri and Simone Mosole), the visits and lab periods spent in international laboratories (in particular those of John Gergely and Alfred Goldberg in Boston), the Italian and International friends with which he has published papers (Anna Jakubiec-Puka, Claudio Franceschi, Giorgio Arpesella, Mike V. Dodson, Stanley Salmons, Winfried Mayr, Simona Boncompagni, Feliciano Protasi, Antonio Musarò, Giorgio Fanò, Vincenzo Vindigni, Franco Bassetto, Francesco Mazzoleni, Dan Graupe, Amber Pond, Marina Marini, Fabio Francini, Paolo Gargiulo, Thordur Helgason, Tiziana Pietrangelo, Nejc Sarabon, and last but not least Helmut Kern) and of course the many others he met during International Conferences.

His "first" Meeting, as a young fellow of myology was organized in Switzerland by Marcus C. Schaub; but how to forget the International Conferences where he and his students had the chance to know Bruce M. Carlson, John Faulkner, Zipora Yablonka-Reuveni, Eric Monnet, Miranda Grounds, Winfried Mayr and many other Vienna friends) or the Conference he organized in Terme Euganee, Padua: where we met Juan Carlos Chachques, a young Surgeon from Argentina working in Paris with Alain Carpentier, Carlo Reggiani, now full professor of Physiology in Padua University, Terje Lomo, Dirk Pette, Salvatore Di Mauro, Clara Franzini-Armstrong, Tessa Gordon, Victor Dubowitz, Terry Partridge, Ryoichi Matsuda, Stanley Salmons, Jonathan C. Jarvis, Dario Coletti, Werner Lindenthaler and many others. Of Gerta Vrbova he will remember that she was one of the first invited speakers he personally met in 1979 in the Margreth's Lab, as a young fellow who presented to her his first independent publication on "*selective maintenance of neurotrophically regulated proteins in long-term denervated hemidiaphragm*".⁵ Finally, he will identify the main research topics he worked on during 45 years of research activity. He started in 1966 to prepare the M.D. Thesis, but he is not yet ready "retire" from Myology: as a Senior Scholar of the University of Padova, he think that he has a lot to translate to clinical colleagues.

The first topic of his long career was: Contractile protein isoforms identified by several electrophoretic methods^{6,7} and their transitions as tools to study modulation and pathology of muscle fiber units and motoneurons. He will remember his first publication on "*Neural control on the activity of the calcium-transport system in sarcoplasmic reticulum of rat skeletal muscle*" by Margreth, Salviati, Carraro in Nature 1973⁸ and six years later "denervation-induced isomyosin transitions" by Carraro, Catani, Biral. Exp Neurol 1979⁵ and by Carraro et al. 1985.⁹ Some years afterwards, inspired by Terje Lomo and Stefano Schiaffino,¹⁰⁻¹⁹ he independently did corroborating experiments of continuous electrical stimulation of denervated muscle, achieving a yet unexplained and infrequently cited high increase of slow muscle fibers properties in denervated fast muscle.²⁰ A system analysis with flow charts may summarizes all the interactions among old and recent topics, Carraro's mentors - students - Padua colleagues with Italian and International Scientists/Clinicians involved in animal and human muscle biology, pathology, therapy and rehabilitation, but he believe he will need months if not years to complete his book project.

He has to describe: Muscle damage and regeneration via myoblast's proliferation, differentiation and fusion,^{21,22} including exercise-induced muscle fiber apoptosis in normal and dystrophic animal and human muscles.^{23,24} He studied: Isomyosins in hypertension and heart failure²⁵ and introduced the concept of

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“Demand Dynamic Cardio-myoplasty,” first in a sheep model with Giorgio Arpesella²⁶ and then in patients with Gianluca Rigatelli.²⁷ Corroborating evidence of effectiveness of the intermittent stimulation strategy was collected on other sheep models.^{28,29}

Finally, the strong leadership of Helmut Kern convinced Engineers in Vienna and then myologists in Italy (Carraro’s team in Padua and Feliciano Protasi in Chieti) to implement two pilot trials supported by the EU Project RISE, the first a cross-sectional study³⁰⁻³⁶ and then a longitudinal-study³⁷⁻⁴⁰ to demonstrate that a home-based strategy of Functional Electrical Stimulation (h-b FES) recovers muscle mass and functions of permanently denervated human muscle.

With EU-support, Helmut Kern and his European collaborators, Carraro’s team included, are now translating this strategy to the more frequent cases of muscle deterioration due to aging and cancer.^{3,41-43}

Further, in Padua he is extending the EU RISE results to partially reinnervating muscle, developing dedicated monitoring strategies.^{39,44,45} To objectivize results of these researches, he is proud to have revitalized the clinical use of ultrasound muscle approaches, adding dynamic analyses of contractile properties in clinical evaluation of denervated and reinnervating muscles. Further, he suggested to Paolo Gargiulo and Helmut Kern to add false color to “Monitoring of muscle and bone recovery in spinal cord injury using three-dimensional imaging and segmentation techniques”, to allow doctors and their patients to read much easy-to-interpret Computer Tomography analyses of their deteriorating or recovering muscles.^{38,39,46}

That is why after almost 50 years, he is still fond of the effects of denervation and of the modulation by electrical stimulation of skeletal muscle fibers, of their adaptation/damage/apoptosis/regeneration potentials by reciprocal interactions with inflammatory cells and nerve, hoping to identify further clues worth to be translated into clinically relevant therapy and rehabilitation strategies.

Helmut Kern: Pioneering research

In 1990 dr. Helmut Kern achieved his Habilitation for M.D. with a thesis that has been published in German in the *Oesterreichische Zeitschrift fuer Physikalische Medizin* 1995; 5: Heft 1, Supplementum.⁴⁷

The thesis is now reprinted in the special issues “The long-term denervated muscle”.⁴⁸ The English abstract is provided in the following paragraph.

Functional Electrical Stimulation on Paraplegic Patients.

We report on clinical and physiological effects of 8 months Functional Electrical Stimulation (FES) of quadriceps femoris muscle on 16 paraplegic patients. Each patient had muscle biopsies, CT-muscle diameter measurements, knee extension strength testing carried

out before and after 8 months FES training. Skin perfusion was documented through infrared telethermography and xenon clearance, muscle perfusion was recorded through thallium scintigraphy. After 8 months FES training baseline skin perfusion showed 86 % increase, muscle perfusion was augmented by 87 %. Muscle fiber diameters showed an average increase of 59 % after 8 months FES training. Muscles in patients with spastic paresis as well as in patients with denervation showed an increase in aerob and anaerob muscle enzymes up to the normal range. Even without axonal neurotropic substances FES was able to demonstrate fiber hypertrophy, enzyme adaptation and intracellular structural benefits in denervated muscles. The increment in muscle area as visible on CT-scans of quadriceps femoris was 30 % in spastic paraplegia and 10 % in denervated patients respectively. FES induced changes were less in areas not directly underneath the surface electrodes. We strongly recommend the use of Kern’s current for FES in denervated muscles to induce tetanic muscle contractions as we formed a very critical opinion of conventional exponential current. In patients with conus-cauda-lesions FES must be integrated into modern rehabilitation to prevent extreme muscle degeneration and decubitus ulcers. Using FES we are able to improve metabolism and induce positive trophic changes in our patients’ lower extremities. In spastic paraplegics the functions „rising and walking“ achieved through FES are much better training than FES ergometers. Larger muscle masses are activated and an increased heart rate is measured, therefore the impact on cardiovascular fitness and metabolism is much greater. This effectively addresses and prevents all problems which result from inactivity in paraplegic patients.

The 325 references are a remarkable collection of the pioneering work on FES in paraplegics that ended up with the first world implant of a device performed in Vienna in 1983.⁴⁹

Since then, an enormous amount of new work has been necessary to establish a clinically accepted strategy for recovery of contractile function of long-term denervated muscle, but the work in the 1970’s and 1980’s has provided a firm and accurate basis for the current understanding of the recovery process in human muscles.

Collaboration of Austrian and Italian researchers

In 1998 Helmut Kern went once again in Terme Euganee to cycle on the Euganei Hills. A late morning he went to the Padua Institute of General Pathology with his Habilitation Thesis to meet Ugo Carraro and to express his strong willingness to collaborate in a scientific study of a series of Conus Cauda sufferers he was training with h-b FES since several years. The reply of Ugo was outspoken: “*Helmut harvest a muscle*

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biopsy and we will show to skeptics that the astonishing functional improvements in muscle contractility you achieved with your elegant training strategy will be supported by evidence of improved muscle fiber size and ultrastructural features". The first biopsy from muscles treated in this way is described in two articles that report the characteristics of the muscle fibers from the Quadriceps of a person after 26 months of denervation and h-b FES Training.^{30,32}

From the second article, submitted years before acceptance (and only after a cross-sectional study published in a prestigious journal provided stronger evidence of the effectiveness of the h-b FES for denervated muscles³²) we here republish the clinical description of this first, very successful case.

"V. Z., a 47-year-old man, had suffered a traumatic cauda equina lesion at T12. One year later, his quadriceps femoris muscles were severely wasted on both sides. Voluntary movement, sensation, and reflexes were all absent, consistent with total denervation. After a further 6 months, findings at neurological examination were unchanged. Absence of volitional activity on needle electromyography (EMG) and of evoked activity using surface EMG with transcranial and lumbosacral magnetic stimulation confirmed permanent and complete loss of motor functions of spinal nerves L1 to L4. Direct electrical stimulation, which in a normal muscle would elicit a response with a chronaxie of 0.1–0.7 ms, required a chronaxie of more than 20 ms, constituting further evidence of complete loss of innervation. A computerized tomography (CT) scan of the thighs revealed marked atrophy of muscle tissue with replacement by fat; the cross-sectional areas of the quadriceps muscles were 36.0 cm² (right) and 36.1 cm² (left, Fig. 1A), representing 58.9% (right) and 59.1% (left) of the corresponding areas in a typical healthy individual. No detectable knee extension torque could be elicited by stimulation under isometric conditions with the subject sitting with the knee flexed at 90°. Eighteen months after his injury, V. Z. commenced a training program which, after appropriate instruction, he was able to carry out at home. Two pairs of large electrodes, each having an area of 200 cm², were strapped to the anterior surface of the thighs in proximal and distal positions. Twitch contractions were elicited by biphasic rectangular current pulses having duration of 120 ms and amplitude of 200 mA, delivering impulse energy of 1.92 Joules, to recruit fibers throughout the quadriceps femoris muscles. The long duration of the impulses needed for stimulation precluded the use of frequencies that would elicit tetanic contractions; training was therefore initiated with single twitches at 2 Hz and delivered for 15 min per day, 5 days per week. After 4 months, excitability of the muscle fibers had recovered

sufficiently for pulses of shorter duration to be used. At this stage, the protocol was augmented with an additional tetanic pattern consisting of pulses of 40 ms delivered at 20 Hz for 2s on, 2 s off for 15 min daily, 5 days per week. The total amount of stimulation was then 30 min daily for each muscle. The additional tetanic stimulation pattern produced more rapid and more forceful contractions, resulting in a progressive increase in knee extension torque. After 26 months of stimulation V. Z.'s thighs came to resemble those of a healthy sedentary subject; although the external appearance was not entirely normal, it was certainly more acceptable cosmetically to the patient. CT scan showed that the cross-sectional areas of the quadriceps muscles at the same level had increased on the right side from 36.0 to 57.9 cm² and on the left side from 36.1 to 52.4 cm² (Fig. 1B); these figures represent 94.7% (right) and 85.7% (left) of the areas typical of a healthy subject. Muscle density, expressed in Hounsfield Units, had risen from 11.0 to 26.4 on the right side and from 10.7 to 24.1 on the left. Stimulation of the quadriceps muscles elicited a knee extension torque of 12.0 Nm on the right and 10.5 Nm on the left. Despite the marked restoration of muscle cross-sectional area, this was less than 10% that of a normal subject. Nevertheless, this stimulation-induced torque enabled V. Z. to extend the knee from a sitting position and to maintain a standing posture without the support of the upper extremities.

Biopsies were taken from the right and left vastus lateralis muscles and frozen sections were stained with hematoxylin and eosin and with a monoclonal antibody (NCL-MHCd; Novocastra Laboratories Ltd, Newcastle upon Tyne, United Kingdom) to the embryonic myosin heavy chain isoform (MHCemb). The sections consisted mainly of large round myofibers with a mean diameter of 37.2 +/- 24.8 μ m (right) and 40.5 +/- 24.9 μ m (left). There was very little fat or fibrous connective tissue. Small myofibers (<10 μ m diameter) were also present. Some appeared to be severely atrophic (Fig. 1C, arrowheads); others, intensely basophilic with several large internal nuclei, we interpret as undergoing regeneration (Fig. 1C, arrows). The latter stained positively with anti-MHCemb (Fig. 1D, green coloration), providing evidence of their recent formation. The antibody also reacted with some larger myofibers (> 30 μ m diameter) with subsarcolemmal myonuclei; we have seen similar fibers in permanently denervated rat muscles in which regeneration had been induced by myotoxin treatment. MHCemb-positive myofibers constituted 8.7% (right) and 2.3% (left) of the identifiable muscle fibers present in the biopsies. Paralysis and denervation were demonstrated clinically in this patient at 12 months and 18 months post-injury and again after 26 months of stimulation. We conclude that the injury was stable and that no recovery could have occurred spontaneously during the period of treatment. Nonetheless, the intensive

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regime of electrical stimulation was associated with an increase in excitability, size, and force-generating capacity of the denervated quadriceps muscles, and this was sufficient to allow knee extension to be induced by electrical stimulation. There was histologic evidence of an accompanying reduction in fat and connective tissue, of growth in diameter of surviving myofibers, and also of regenerative phenomena resulting in the formation of new myofibers.

Previous studies on denervated muscles in both animals and humans have shown that electrical stimulation can induce a small increase in muscle mass and hypertrophy of the denervated muscle fibers. The present case is unusual in the extent of the changes produced in the long-term denervated and stimulated

rationale to plan research aimed to recover long-lasting denervated muscle.

Permanent denervation of the limbs muscles due to LMN lesion may take place when trauma to spinal cord, roots and peripheral nerves occurs. Examples are spinal cord lesion with concomitant root damage, brachial plexus palsy and sciatic nerve injury. When proximal denervation occurs, it may be necessary more than a year for reinnervation, during which the process of severe atrophy and fibrosis of the affected muscle tissue may impair synaptic reorganization.

One of the effects of SCI is the rapid loss of contractile force and mass of the affected muscles. Atrophy of leg muscles is particularly severe when the injury destroys the LMN and, hence, the contacts between motor neurons and muscle fibers. In this case, within weeks atrophying and fibrillating muscles become unable to sustain tension during tetanic contractions induced by electrical stimulation. Within months, the denervated leg muscles are no longer excitable by standard commercial electrical stimulators because they have undergone severe disorganization of contractile elements (myofibrils) and of the excitation-contraction coupling apparatuses.³¹ Finally, after years of denervation, muscle fibers are replaced with adipose and fibrous tissues.^{30,31,37,38,58}

These severe functional and structural changes of denervated muscle tissue are not detectable in patients suffering with complete upper motor neuron (UMN) lesions even 20 years after SCI at thoracic level.⁵⁹ On the other hand, larger trauma of the lumbar and ischiatic regions, complicated by ischemic and infection necrosis of the spinal cord, may extend the damage to large segments of the medulla and of the nerve roots. In these latter cases, the diagnostic problems are related to completeness of the LMN denervation, while the absence of sensation of the legs and of the pelvic sphincters grants completeness of the transverse spinal cord lesion (ASIA grade A of SCI).

To avoid problems in interpreting clinical results related to residual LMN innervation or reinnervation, we firstly designed and implemented a cross-sectional study³¹ followed by a longitudinal 2-year long prospective study^{37,38} that recruited 25 paraplegic patients specifically selected because of complete LMN denervation of the quadriceps muscles. In the longitudinal study, the same group of patients was assessed before and after two years of h-b FES by clinical, functional, imaging and muscle biopsy analyses.^{37,38,60} To avoid the potential criticisms related to a possible residual innervation or reinnervation, dedicated diagnostic protocols were designed and implemented to test "completeness" of LMN denervation of right and left quadriceps muscles before and during the two years of the study.^{38,61} By means of these tests, in particular Test electrical stimulation by bidirectional rectangular impulses of 1 ms, 40Hz, 100mA of amplitude testing for contraction of the

thigh muscle, the complete and permanent denervation of the quadriceps before the beginning of h-b FES, and after two years of training was fully granted, since the stimulated muscle improved their excitability recovering tetanic contractility, but never responded to the settings which are able to elicit contraction of innervated muscle.³⁸ If no electrical stimulators providing the high level stimulation parameters are available, electrical stimulation with bidirectional rectangular impulses of about 1 ms, a frequency of 40Hz and an intensity of 100mA can be used for the first evaluation of the paralyzed muscle. These parameters can be delivered by most of the commercially available devices and are sufficient for a first diagnosis if the stimulated muscle shows signs of denervation. All together, the above described behaviors leave a significant time window for intervention to avoid degeneration of the LMN denervated muscle by home based electrical stimulation.

From the first biopsy to the end of the European Project RISE: Use of electrical stimulation to restore standing in paraplegics with long-term denervated degenerated muscles (Contract no. QLG5-CT-2001-02191)

From early 2000 to August 2004 more than 130 biopsies of Conus Cauda Patients were analyzed in Padua (by morphometry and immunostaining) and in Chieti (electron microscopy). Further, muscle biopsies from spastic paraplegics (i.e., those with lesion of the upper motoneuron) were also analyzed to described the differential behaviors of truly disconnected muscle fibers to those severely atrophic (but never degenerated) due to severe unloading.⁵⁹

Aim of the EU Commission Shared Cost Project RISE (Contract no. QLG5-CT-2001-02191) was to confirm previous results of a cross-sectional study³¹ by a longitudinal prospective study in 25 paraplegic patients specifically selected because of complete LMN denervation of the quadriceps muscles. The overall conclusions, taken from Kern et al. 2010 NNR article³⁸, of all these studies may be summarized as follows:

"Atrophy of skeletal muscle groups is particularly severe when SCI involves all the lower motor neurons (LMNs). After such a complete injury, the peripheral endings of motor neurons quickly degenerate whereas LMN denervated muscles undergo progressive decay, which can be roughly divided in the following chronological steps: (a) in days denervated muscle starts to spontaneously activate action potentials (fibrillations); (b) in weeks, muscles become unable to sustain tension during tetanic contractions induced by electrical stimulation; (c) within months, muscles are unexcitable with standard commercial electrical stimulators,[62-66] undergoing ultrastructural severe

disorganization of the E-C Coupling and of the contractile apparatuses; and (d) after years, the myofibers are replaced by adipocytes and collagen.

To counteract the progressive changes that transform muscle into an unexcitable tissue, over the past 20 years we have developed a novel therapy concept for paraplegic patients with bilateral and complete LMN denervation of the lower extremity due to complete lesions of the conus and cauda equina (CC). This new training strategy became possible because of the development of a new generation of stimulation equipment specifically designed for home-based functional electrical stimulation (h-b FES). These new stimulators and the large surface electrodes necessary to cover the denervated muscles were developed by the Center of Biomedical Engineering and Physics at the Medical University of Vienna and by the Wilhelminenspital, Vienna (Austria), to reverse long-standing and severe atrophy by delivering high-intensity and long-duration impulses that can directly elicit contraction of denervated skeletal fibers in the absence of nerve endings. Our data, indeed, show that h-b FES can be an effective home therapy to counteract muscle atrophy and degeneration after complete LMN denervation due to CC lesions. The h-b FES device stimulates muscle fibers in the absence of nerve endings and after prolonged denervation, enabling (a) recovery of muscle mass and fiber size, (b) recovery of tetanic contractility, and (c) restoration of muscle fiber ultrastructure.

Up to now, the muscles of affected extremities in these paraplegic patients are commonly not treated with FES because it is widely accepted that long-term and completely denervated muscles cannot be effectively stimulated. On the other hand, studies in animal models and humans indicate that (a) severe atrophy does not occur in rats for at least 3 to 4 months; (c) in rabbit, the degeneration of muscle tissue does not appear during the first year of denervation; and (c) in humans, muscle tissue degeneration starts from the third year onward. Our recent findings that the long-term denervated rat muscle maintains L-type Ca^{2+} current and gene expression of the related proteins longer than functional contractile machinery provide the molecular, structural, and functional rationale of rehabilitation training for permanently denervated muscles, consistent with clinical observations. This leaves a window of opportunity to initiate muscle stimulation and avoid muscle degeneration and infiltration.

Our light microscopy results suggest a window for intervention in patients up to 2 years after injury, because fibers maintain at least 30% of their initial size and the extracellular matrix is still evolving. EM analyses, on the other hand, indicate that the structure of the sarcotubular system (reputed to deliver action potential to the fiber interior) and myofibrils decays quite quickly, suggesting that it is best to start h-b FES

training as soon as possible after SCI, possibly not later than six months. The poor excitability/contractility of human long-term denervated fibers is likely attributable to ultrastructural changes that affect the EC coupling apparatus and contractile elements and precede severe atrophy and degeneration. The reorganization of T-tubules and Ca^{2+} release units and myofibrils that follows h-b FES likely plays a role in the recovered ability of LMN denervated muscles to be stimulated and to respond with tetanic contractions.

Because the progression of recovery in h-b FES-trained LMN denervated muscle is inherently slow, in part due to exercise training for only 30 minutes per muscle group, 5 times a week, patients were clinically evaluated every 12 weeks by physiatrists, who progressively modified the stimulation parameters and training protocol according to the patient's improvements. During the first few months of h-b FES training, the initially poor excitability of the denervated muscle was improved by twitch-contraction training. Three to 6 months later, electrical stimulation induced tetanic contractions against loads that were progressively increased, accompanied by a significant increase in the mass of the quadriceps muscles (24% at the midterm evaluation) and by improvement in limb appearance and muscle cushioning. None of the subjects that reached 1 year h-b FES training ($n = 20$) declined in terms of their muscle properties, and 20% reached the ability to perform stand-up exercise assisted by electrical stimulation of quadriceps muscles.

At 2 years, 90% ($n = 20$) of h-b FES trained subjects recovered or increased tetanic contractions, and 25% stood during electrical stimulation in parallel bars. Minimal functional improvements were associated with long time elapses between SCI and initiation of h-b FES and possibly lower compliance with training. In single case reports, low compliance substantially decreased the effects of training, yet in the same subjects the mass of thigh muscles increased when the patient resumed h-b FES.

The likelihood that the lower extremities of these patients were completely denervated before initiation of h-b FES training and remained denervated during and after the 2 years of training was indicated by several assessments (e.g., test electrical stimulation, needle electromyography, transcranial and lumbosacral magnetic stimulation). In particular, the threshold of excitability of the quadriceps muscles never increased to a level that allowed them to respond to standard commercial electrical stimulators (impulse duration about 0.5-2 ms), which elicit a muscle contraction through the nerve. The severity of postdenervation atrophy (and the extent of h-b FES-induced recovery) was similar in the left and right quadriceps muscles of the same patient. In incomplete denervation (or some re-innervation), we would have

expected greater variability. Finally, patients did not describe pain during surface stimulation with high current (1000-3000 times higher energy [2.4 J at 120 ms and 200 mA] than that delivered by standard commercial stimulators [0.8 mJ at 0.7 ms and 50 mA]), implying complete sensory loss.

In conclusion, our findings strongly support the RISE rehabilitation protocol as a method to improve the mass and contractility of LMN denervated muscles, although we found a limited "measurable" knee torque changes in h-b FES trained muscles. These benefits could be extended to patients with similar lesions, especially to determine whether h-b FES can reduce secondary complications related to disuse and impaired blood perfusion (reduction in bone density, risk of bone fracture, decubitus ulcers, and pulmonary thromboembolism).

On the other hand, the Authors share the following suggestions of Gerta Vrbova, which was so kind to attract our attention on the intrinsic limiting factors that will never allow long term denervated muscle to reach by Electrical Stimulation (as it is feasible in clinical settings) the stage of a fully normal muscle. Indeed, our main evidence for muscle denervation even after years of h-b FES is the fact that the trained muscles never attain the ability to respond to the much lower currents that stimulate curarized or the denervated muscle fibers early after degeneration of the peripheral nerve stump.

"While there is no doubt that impulse activity has a decisive role in determining muscle properties [67] it cannot entirely replace the effect of innervation on denervated muscle. Whether this is due to a trophic influence of nerve on muscle or other factors has not yet been resolved.

There are several possible reasons why electrical stimulation cannot entirely mimic the effect of innervation on skeletal muscle: 1. Denervated muscles are stimulated in a manner that causes synchronous contraction of all muscle fibers in the stimulated muscle. This differs greatly from the activity that the nerve is imposing onto the muscle it innervates. During nerve induced movement different motor units are activated asynchronously, and never at the same time [68]. Thus the recruitment order of different muscle fibers is completely different from electrically induced muscle stimulation. 2. The synchronous activity of denervated muscles cannot mimic that which occurs during natural movement and as a consequence the mechanical conditions of different muscle fibers within the stimulated muscle will be far from normal. The amount of load during contraction affects slow muscle fibers more than fast ones; indeed they degenerate if they contract in the absence of load [69]. 3. The simple interpretation of the effect of whole muscle stimulation is therefore limited for synchronous stimulation of all

muscle fibers in denervated muscles is very different from nerve induced activity during normal movement.

4. Apart from the superbly organized recruitment order of motor units during normal movement that seems to be necessary for the integrity of the different types of muscle fibers there could be an additional trophic effect of the nerve on muscle but there is little evidence for such an influence that is independent on muscle activity or the mechanical conditions.

We are aware, indeed, that the clinical results may appear poor or very poor to "normal people", but, please, reader consider them from the point of view of a disabled person at risk of serious complications. The increase in mass (cushioning effect) and the anti-gravitational pumping of leg blood are muscle "functions" that are fully lost after denervation, but are substantially recovered during long-term daily electrical stimulation.

Devices and Vienna Stimulation Strategy for h-b FES of large denervated human muscles in SCI

To counteract the progressive changes that transform muscle into an unexcitable tissue unable to generate force with standard commercial stimulators (from six months onward), in the past 20 years Clinicians and Engineers developed in Vienna novel rehabilitation concepts for paraplegic patients with bilateral and complete LMN denervation of the lower extremity due to complete lesion of the *Conus Cauda*.⁷⁰ This new rehabilitation protocol became possible due to the development and optimization of new stimulation equipments for FES. The devices have been specifically designed to reverse longstanding and severe atrophy of LMN denervated muscles by delivering high-intensity and long-duration impulses that can directly elicit contraction of denervated skeletal fibers in absence of nerve endings. These new stimulators and the large surface electrodes needed to cover the denervated muscles were developed by the Center of Biomedical Engineering and Physics at the Medical University of Vienna, Austria.⁷¹⁻⁷³ In parallel, specific clinical assessments and training settings were developed at the Wilhelminenspital Wien, Austria.^{61,74,75} The rehabilitation progressive training strategy for LMN denervated muscles (see the Figure 1) are validated by the clinical results, strongly supported by those obtained from light and electron microscopy muscle biopsies' analyses performed in Padua and Chieti Universities (Italy), respectively, as described by Kern et al. in the longitudinal prospective study.^{38,59} Patients were provided with stimulators and electrodes in order to perform stimulation at home for five days per week. The large (180 cm²) electrodes (Schuhfried GmbH, Mödling, Austria) made of conductive polyurethane, were placed on the skin surface using a wet sponge cloth (early training) and fixed via elastic textile cuffs. As soon as the skin was accustomed to



FES training protocols for the functional recovery of permanent complete denervated human muscles

Stefan Löffler¹, Winfried Mayr², Michaela Mödlin³, Samantha Burggraf¹, Helmut Kern^{1,3}

¹ Ludwig Boltzmann Institute of Electrical Stimulation and Physical Rehabilitation, Vienna, Austria

² Center for Medical Physics and Biomedical Engineering, Medical University of Vienna, Austria

³ Institute of Physical Medicine and Rehabilitation, Wilhelminenspital, Vienna, Austria

Table 1. FES training of short term denervated (< 2 years) human muscles (adapted from Kern et al. Neurorehabil Neural Repair. 2010 Oct;24(8):709-21).

Timeline (months)	Stimulation parameters	Training parameters
0-2	120ms ID / 500ms IP; 4s SD / 2s SP	5x3min 5d/week
3-4	120ms ID / 500ms IP; 5s SD / 2s SP 40ms ID / 10ms IP; 3s SD / 3s SP	5x3min 5d/week 3x3min 5d/week
5-6	120ms ID / 400-500ms IP; 5s SD / 1s SP 40ms ID / 10ms IP; 3s SD / 3s SP	5x4min 5d/week 3x3min 5d/week
6-8	120ms ID / 400ms IP; 5s SD / 1s SP 40ms ID / 10ms IP; 3s SD / 3s SP	5x4min 5d/week 3-4x3min 5d/week + ankle weight 2x/week
8-	120ms ID / 400ms IP; 5s SD / 1s SP 40ms ID / 10ms IP; continues + switch	5x4min 5d/week stand up – sit down exercise stand up – stepping – sit down exercise
16-	120ms ID / 400ms IP; 5s SD / 1s SP 40ms ID / 10ms IP; continues + switch	5x4min 5d/week walking exercise

ID, impulse duration (biphasic, rectangular or biphasic); IP, impulse pause; SD, stimulation duration; SP, stimulation pause

The recommended parameters and time intervals are suggestions, based on the EU project FASE and our clinical experience. They should be adapted to personal needs of patients in respect to the time span of denervation and condition of muscle and function.

Table 2. FES training of longer denervated (1% to 4 years) human muscles (adapted from Kern et al. Neurorehabil Neural Repair. 2010 Oct;24(8):709-21).

Timeline (months)	Stimulation parameters	Training parameters
0-2	150ms ID / 500ms IP; 4s SD / 2s SP (maybe no contraction visible)	3x3min 5d/week (week 0-4) 4x3min 5d/week (week 5-6) 5x3min 5d/week (week 7-8)
3-4	120ms ID / 500ms IP; 4s SD / 1s SP 40-80ms ID / 500ms IP; 4s SD / 1s SP	5x3min 5d/week 5x3min 5d/week
4-6	120ms ID / 400-500ms IP; 5s SD / 2s SP 40-50ms ID / 10ms IP; 2s SD / 2s SP	5x4min 5d/week 3x3min 5d/week
6-8	120ms ID / 400ms IP; 5s SD / 1s SP 40ms ID / 10ms IP; 3s SD / 3s SP	5x4min 5d/week 3-4x3min 5d/week
8-11	120ms ID / 400ms IP; 5s SD / 1s SP 40ms ID / 10ms IP; 3s SD / 3s SP	5x4min 5d/week 3-4x3min 5d/week + ankle weight 2x/week
12-	120ms ID / 400ms IP; 5s SD / 1s SP 40ms ID / 10ms IP; continues + switch	5x4min 5d/week stand up – sit down exercise
16-	120ms ID / 400ms IP; 5s SD / 1s SP 40ms ID / 10ms IP; continues + switch	5x4min 5d/week stand up – stepping – sit down exercise

ID, impulse duration (biphasic, rectangular or biphasic); IP, impulse pause; SD, stimulation duration; SP, stimulation pause

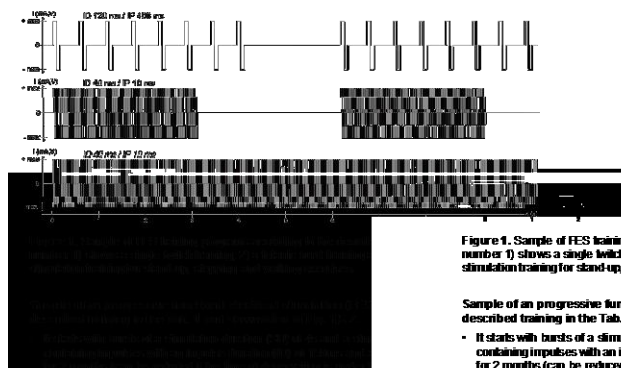


Figure 1. Sample of FES training (number 1) shows a single twitch stimulation training for stand up.

Sample of an progressive functional training in the Tab. 1:

- It starts with bursts of a stimuli containing impulses with an in for 2 months (can be reduced IP, 5s SD and 1s SP after 2 months to activate poor denervated muscle fibers, (see Fig. 1.14, 3)
- The next training phase implements tetanic bursts of 3s SD and 3s SP with impulses of 40ms ID and 10ms IP after 2 months of stimulation - beside the single twitch program - to increase muscle fiber diameter, muscle mass, density and force with leg extensions (after 2.5 months) with and without additional weights on the subjects ankle. (see Fig. 1.24, 4+5)
- If a good condition is achieved (depending not only from the training also from the time span of denervation) the force training can be replaced up to 2-4x/week with stand up, stepping and walking exercises performed with continuous stimulation (controlled by an external switch) with 40ms ID and 10ms IP. (see Fig. 1.34, 6+7)

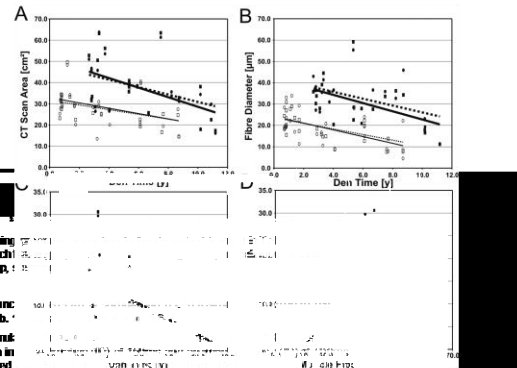


Figure 2. Kern et al. functional outcomes show results before: refer to right and left A, muscle cross-section (quadriceps area) (standing area) B, size of vastus lateralis C, maximum tetanic D, correlation between

Neurorehabil Neural Repair. 2010 Oct;24(8):709-21 Morphological and after 2 years of FES. Thin and thick lines and empty and filled squares/circles and after FES, respectively. Continuous and dotted lines and squares and circles legs, respectively.
Total area by computed tomography scan
28.2 ± 8.1 cm² vs. 38.1 ± 12.7 cm², p=0.001
25.8 ± 8.4 cm² vs. 30.7 ± 9.9 cm², p=0.001
skeletal muscle fibers (mean diameter: 16.6 ± 14.3 μm vs. 29.1 ± 23.3 μm, p=0.001)
torque under stimulation (mean torque 0.8 ± 1.3 Nm vs. 10.3 ± 8.1 Nm, p=0.001)
in mean fiber diameter and stimulated tetanic torque (r=0.67)



Figure 3. Training with single twitch stimulation on the ankle as described in Tab. 1 (0-2 months)



Figure 4. Training with tetanic stimulation in standing position as described in Tab. 1 (2-8 months)

EU Project FASE - Functional Assessment of Stimulation and Exercise in long-term denervated muscles (FASER)
QL-GS-CT-2001-02151

Ludwig Boltzmann Institute of Electrical Stimulation and Physical Rehabilitation | Montleirgasse 37, 1160 Vienna, Austria

Figure 1. Vienna progressive h-b FES strategy for long-term complete denervation of quadriceps muscles

the necessary high current density, gel was used under the polyurethane electrodes to achieve minimal transition impedance. A special design feature was a non conductive bulge along the entire edge of the

electrode that prevents potential skin burns that presumably can occur where a conductive edge gets in electrical contact with skin surface and causes local current density hot-spots (Mayr W, 2007; Patent -

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Surface Electrode, EP2021068, WO/2007/131248). The electrodes were flexible enough to maintain evenly distributed pressure to the uneven and moving skin, thus providing homogeneous current distribution throughout the entire contact area.

Stimulation needle electromyography (SNMEG) was used to study the electrophysiological properties of single muscle fibers. We measured the muscle fiber conduction velocity (MFCV) and the shortest interstimulus interval (ISI) still eliciting a response to the second stimulus delivered to the fiber.⁷⁶ MFCV recorded in the denervated patients before and after h-b FES therapy showed a significant increase in conduction velocity (fastest and mean CV) and reduced refractory periods (shortest ISI). This suggests that electrical stimulation training is effective to improve the electrical properties of the muscle fibre and SNEMG could serve as an additional measurement technique to specify the status of the denervated muscle. The training strategy consisted of two combined stimulation programs.^{31,38} All applicable rules concerning the ethical use of human volunteers were followed during the course of this research (Approval of Ethical Committee, Vienna, Austria: EK-02-068-0702). For a multilingual translation of the Work Packages, ethical and safety issues link to: <http://www.bio.unipd.it/bam/bam18-2&3.html>

At the beginning of the treatment, biphasic stimulation impulses of very long-duration (120-150 ms, 60-75 ms per phase) at high intensity (up to $\pm 80V$ and up to ± 250 mA) were applied (Training Program 1). Then the subjects underwent clinical assessment and stimulated knee torque measurement every 12 weeks by physiatrists, who progressively modified the stimulation. The routine daily training consisted of combined twitch and tetanic stimulation patterns (Training Programs 2, 3 and 4) in consecutive sessions lasting up to 30 min for each group of muscles (gluteus, thigh and lower leg muscles on both sides). After tetanic contractility was achieved and the subject achieved full extension of the leg, the ankle was progressively loaded (Figure 1). Finally, the more compliant patients became able to stand and perform step-in-place and (if young and light) walking exercise.^{37,38}

Perspectives

In collaboration with his international partners, Dr. Kern is now extending the benefits of h-b FES to those subjects, which for different reasons, from the mild but unrelenting process of aging to the devastating fast progression of muscle atrophy in cancer patients, suffer the consequences of muscle weakness. Further, a multi-disciplinary research team of the Interdepartmental Research Center of Myology of the University of Padua is applying the Vienna principles to the apparently easier cases of peripheral incomplete denervation of arms and legs.

Examples in literature of the effectiveness of life-long high-level physical activity in postponing effects of aging,^{3,4,77} and of physical approaches in peripheral and central neural repair, seems to open new perspectives to an approach, home based Functional Electrical Stimulation in paraplegics by implanted electrodes and neuromodulators, that has been abandoned twenty or more years ago despite the successes of heart pacing and mini-implants for deaf, two very successful cases of Functional Electrical Stimulation of human tissues. H-b FES is worth to be reassessed under strict scientific rules, balancing its costs against the needs and rights of patients to see alleviated their burdens.

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Corresponding Author

Ugo Carraro, CIR-Myo Translational Myology Lab, Department of Biomedical Sciences, University of Padova, Italy
E-mail: ugo.carraro@unipd.it

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