Basic Principles of Managing Post-Polio Syndrome
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Once again Dr. Perlman generously shared her time and energy with local polio survivors to discuss post-polio research and answer our questions. The two articles based upon this presentation are printed separately. This report is based on a study in Norway looking at Treatment Options for PPS, plus Dr. Perlman’s answers to questions from the audience; our April 2012 newsletter will feature a summary of the results from Pain research in Sweden and answers to more questions. Although the post-polio news in 2010 was limited, many post-polio researchers around the world continue to enlarge our knowledge.

Update on Current and Emerging Treatment Options
For Post-Polio Syndrome
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Post-polio syndrome (PPS) refers to the clinical deterioration experienced by many polio
survivors several decades after their acute illness. The symptoms are new muscle weakness, decreased muscle endurance, fatigue, muscle pain, joint pain, cold intolerance, and this typical clinical entity is reported from different parts of the world. The pathophysiology behind PPS is not fully understood, but a combination of distal degeneration of enlarged motor units caused by increased metabolic demands and the normal aging process, in addition to inflammatory mechanisms, are thought to be involved. There is no diagnostic test for PPS, and the diagnosis is based on a proper clinical workup where all other possible explanations for the new symptoms are ruled out.

The basic principle of management of PPS lies in physical activity, individually tailored training programs, and lifestyle modification.

- Muscle weakness and muscle pain may be helped with specific training programs, in which training in warm water seems to be particularly helpful.
- Properly fitted orthoses can improve the biomechanical movement pattern and be energy-saving.
- Fatigue can be relieved with lifestyle changes, assistive devices, and training programs.
- Respiratory insufficiency can be controlled with noninvasive respiratory aids including biphasic positive pressure ventilators.
- Pharmacologic agents like prednisone, amantadine, pyridostigmine, and coenzyme Q10 are of no benefit in PPS.
- Intravenous immunoglobulin (IVIG) has been tried in three studies, all having positive results. IVIG could probably be a therapeutic alternative, but the potential benefit is modest, and some important questions are still unanswered, in particular to which patients this treatment is useful, the dose, and the therapeutic interval.

“All polio survivors are not the same and will not necessarily respond to the same treatment,” said Dr. Perlman. “It is important to control pain and fatigue in people with any type of illness. Once your pain is under control and your energy levels are coming back, you can do almost anything.”
Answers to Questions from the Audience

How do I find out which specialist my primary physician should refer me to for fatigue, exercise, incontinence, etc. in post-polio?

This primary physician is clearly feeling overwhelmed by a lot of symptoms. The best type of physician to see is one who can actually deal with rehabilitation of all these areas and that is a physiatrist, a physical medicine and rehabilitation physician (PM&R) specialist.

Brain fatigue – what is it?

Brain fatigue, as described in surveys of polio survivors, is a sense of not being able to think clearly, having trouble with multi-tasking, thinking, and following things through without getting distracted. If you have ever had the flu, that’s brain fatigue. Polio survivors can get that exact feeling. It is felt to be related to the immune system chemicals that have been found to be elevated in the spinal fluid of polio survivors. Some of them are the same immune system chemicals that are elevated in multiple sclerosis (MS) and cause fatigue in MS. For people with brain fatigue, the immune system needs to be an area of research.

Does brain fatigue contribute to physical fatigue?

Yes, if your brain is fatigued you don’t feel like doing anything.

Is it possible to strengthen my breathing muscles, as I have to sleep with oxygen? My oxygen level while sleeping is between 85 to 90%.

I think the only way to strengthen breathing muscles is to work with a pulmonary rehab person who will actually target some of the muscles that are involved in expanding the chest, especially when you are asleep and it is involuntary. During the day when you are awake, you can breathe deeply, but at night it is really out of your control. For somebody clearly with oxygen problems and breathing muscle problems, Dr. Perlman would strongly recommend a round of pulmonary rehab.

Is a surgery such as removal of parathyroid gland contraindicated? (This is someone who uses a BiPAP at night, suggesting there is some respiratory or neck involvement.)

Whenever you have a surgery in a region, whether it is a shoulder or neck, where there are muscles involved, you are going to lose a grade of muscle strength. It might come back, but
post-operatively you may find that the neck is weaker just because they had to manipulate the muscles to get to the parathyroid gland.

**What is resistive exercise?**

Resistive exercise is when you add some resistance, such as a weight or something you are pushing against. You can get the same mileage in half as many repetitions, with less fatigue. That includes ankle weights, wrist weights, using an exercise machine that has a resistive plate that you push, elastic rubber bands, etc.

Antigravity exercise does help strengthen bones. It is one of the things they recommend to people with osteoporosis. Polio survivors can have thinner bones in their polio limbs. So strengthening exercises in those limbs, even two or three repetitions, not only will help the muscles, but will also help the bones – any place where the force of the exercise is felt.

**When exercising, do you recommend a day in between exercises?**

Most exercise for strengthening and conditioning can be done can be done every other day. A rest day in between is good protection from overdoing.

Exercise for pain, which is usually stretching, or exercise for balance does need to be done every day. It is hoped that stretching and balance exercises will not be fatiguing.

**Would Pilates be an appropriate type of exercise for strength and conditioning?**

Pilates works on the core so in that way it does strengthen and condition core muscles. If you have a more specific problem with shoulders, upper arms, or quads you may have to ask for something special with the Pilates therapist. One thing about Pilates is that it tends to be non-fatiguing so it seems to be well tolerated by neuromuscular patients.

**Is there exercise for torn rotator cuffs?**

What you need is advice from your orthopedist about what is safe exercise to do for the amount of rotator cuff tear you have, in order to not make the tear bigger and to promote healing.
Are there any hopeful studies showing stem cell treatment for post-polio? Is post-polio not a good candidate for stem cell treatment?

If we want to replace nerve cells in a shoulder that has no deltoid muscle, no posterior shoulder muscles (they come from the same part of the cervical cord), potentially this could be treated like a spinal cord injury. You have a specific place to go, and you can target it regionally.

Does post-polio research relate to other diseases or disabilities?

Absolutely. Post-polio research relates to any disease of the spinal cord and any disease of motor nerves, for sure. If there is an immune component, then it connects itself with all the immune mediated neurological diseases such as Guillain-Barre, multiple sclerosis, and there are many others. Dr. Perlman knows people who have done studies on ALS, and have added a small parallel group with post-polio. Discoveries in post-polio can help research in other diseases and vice versa. That needs to be emphasized when talking with research groups.

Is there a need for funding of post-polio research?

Yes. The National Institutes of Health (NIH) have funded a lot of prior post-polio research. Many of the studies on www.clinicaltrials.gov are funded by the NIH but their budget has gone down and there is less money for research.

In order to get money from the NIH you have to demonstrate that you have an ironclad, rock solid research project that is going to hit the basic clients of post-polio and combine it with something that has some treatment related potential. They are not interested in general reviews. They want to see things that get to the heart of post-polio and similar diseases and will also open a door to treatment. Certainly there are excellent researchers working.

Those $25,000 grants by Post-Polio Health International fund extremely important research. That is enough money to hire a technician to do research in your lab on a particular project. Even those small amounts of money, as opposed to the mega grants, can be helpful.

Why are polio and post-polio syndrome not addressed in medical school?

Changes in curriculum are very slow; polio was never part of the original curriculum. Post-polio is still not well enough defined as an immune problem so that it could be presented in the same unit as Guillian-Barre. That is one reason it is not in the main curriculum.

On the other hand, if a school has a neuro-rehab unit and students hear a lecture on it, polio and
post-polio syndrome absolutely should be mentioned. If the medical students hear a lecture by a physiatrist about rehab, which hopefully they are hearing about, that lecturer could include post-polio.

Remember, “All polio survivors are not the same and will not necessarily respond to the same treatment,” said Dr. Perlman.

**Conclusion**

As PPS research in many areas continues worldwide, we look forward to learning more from Dr. Perlman in the future.

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