Skin Punch Biopsy May Predict, Diagnose Neuropathy Early

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WASHINGTON — Analysis of skin punch biopsy specimens may help predict diabetic neuropathy in at-risk patients, diagnose the condition earlier, and assess treatment effectiveness, using the Neuropathy Disability Score. ENFD correlated well with assessments of corneal sensitivity and neuropathy severity.

Epidermal denervation indicates small fiber neuropathy in a newly diagnosed diabetic with foot symptoms.

Skin punch biopsy specimens contain small unmyelinated nerve fibers that are damaged in early diabetes. Normative data have shown that the degree of involvement is related to the degree of glucose dysmetabolism, said Dr. Polydefkis, codirector of the cutaneous nerve laboratory at Johns Hopkins University, Baltimore. Analysis of skin biopsy specimens can help exclude other potential causes of painful feet such as radiculopathy, Morton’s neuroma, tarsal tunnel syndrome, and intrinsic foot disease, he said.

A 3-mm diameter, circular skin punch biopsy specimen (about half the size of a pencil eraser) is sliced into about 60 sections, 4 of which are randomly selected for analysis to reduce sample bias since there may be differences in nerve density in different parts of the specimen. Biopsies should be taken at the ankle or other distal sites in patients with early neuropathic symptoms but at the thigh or other proximal sites in those with advanced neuropathy, he said.

In a study of 73 patients with an unknown cause of peripheral neuropathy, Dr. Polydefkis and his associates found that epidermal nerve fiber density (ENFD) in skin punch biopsy specimens was a good marker of early neuropathy. Of the 73 patients, 25 were diagnosed with impaired glucose tolerance (IGT) and 16 had diabetes. The ENFD in biopsy specimens of patients with diabetes or IGT was significantly reduced, compared with healthy control patients. In specimens from the distal leg, but not the distal or proximal thigh, diabetic patients had significantly lower ENFD than did patients with IGT. Yet nerve conduction studies yielded normal results on average in both groups.

Dr. Polydefkis and his associates concluded that the measurement of ENFD in skin punch biopsies is a more sensitive marker for detecting neuropathy early than are nerve conduction studies that test the function of large myelinated nerve fibers (Neurology 2003;61:108-11). At the end of an average follow-up of 4.4 years, repeat testing in 29 patients showed that decline in ENFD was greatest in patients with diabetes, followed by patients with IGT and patients with idiopathic neuropathy. ENFD was essentially stable in 10 healthy control patients. “This longitudinal data provide some of the best evidence that the association between IGT and neuropathy is, in fact, causal,” Dr. Polydefkis said.

Information on nerve morphology also can be extracted from skin punch biopsy specimens. In one study, investigators performed nerve conduction studies, quantitative sensory testing, and skin punch biopsies in the proximal thigh and distal leg at baseline and after a mean of 19 months of follow-up in 15 patients with foot pain but few or no symptoms of neuropathy (6 with diabetes, 1 with AIDS, 1 with paclitaxel toxicity, 7 with an idiopathic nature) and 15 age-matched, healthy control patients. At baseline, patients had significantly lower ENFD in the distal leg than did controls. At follow-up, the ENFD had declined in both biopsy locations but only by a significant amount in the distal leg, compared with the controls (Neurology 2003;61:631-6).

The results of that study provide “evidence that the skin biopsy can be used to predict development of neuropathy,” Dr. Polydefkis said. Clinically meaningful changes in ENFD are on the order of a loss of 2-3 nerve fibers per millimeter. Diabetic patients who present with painful peripheral neuropathy appear to have ENFD losses in the range of 1 fiber/mm per year, while losses for those with established diabetes and peripheral neuropathy may be slightly higher, he said.

Skin punch biopsies also may help investigators to understand how nerve regeneration occurs after an injury in diabetic patients, Dr. Polydefkis said. In a study of patients who applied capsaicin topically to their distal thighs—causing denervation of the epidermis—he and his colleagues found the rate of regeneration depended in part on the patient’s baseline ENFD. Diabetic patients had a significantly lower regeneration rate than did healthy control patients even after adjustment for baseline differences; the rate was even lower among diabetic patients with neuropathy than in those without it. After 100 days, neither group of diabetic patients regenerated their ENFD to baseline levels (Brain 2004;127:1606-17). (See photos.)"