Late-Onset GAD More Common Than Thought

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PARIS — Generalized anxiety disorder occurs as a late-onset condition more often than is recognized, according to investigators who presented the first large treatment trial in elderly patients during the annual congress of the European College of Neuropsychopharmacology.

Dr. Francesca Baldini reported that the average age of onset was 56 years in 273 outpatients with generalized anxiety disorder (GAD) who were aged at least 65 years and who enrolled in the placebo-controlled trial. The study found pregabalin (Lyrica) to be at least as effective as in previous trials with younger patients.

“In reality, there is a late onset of generalized anxiety disorder in the elderly, as in previous trials with younger patients,” said Dr. Baldini, medical director of worldwide neuroscience at Pfizer Inc., in a presentation of the new data at a session on psychiatric disorders in older people.

“That is not in the literature,” added the lead author, Dr. Stuart A. Montgomery, a professor emeritus of psychiatry at the Imperial College London.

Commenting from the audience at Dr. Montgomery's invitation, an investigator of pregabalin in adult patients responding to the characterization of GAD as an early-onset disorder as a misconception. Most patients develop the full syndrome after the age of 30, according to Dr. Hans-Ulrich Wittchen, professor of clinical psychology and epidemiology at the Technische Universität Dresden (Germany).

He urged careful questioning of newly diagnosed older GAD patients who report they first had symptoms as children. "It may be that there are anxiety disorders preceding that (new) development... but if you miss the GAD symptoms, then this is another disorder," he said. "This is not an early anxiety disorder.""

In a presentation, Dr. Baldini said that at least half of all newly diagnosed cases of GAD occur after age 30 years and a third after age 40. Five different studies from the United States and Europe have shown prevalence rates ranging from 1.9% to 7.2% in the elderly, she said.

All patients in the pregabalin study had Hamilton Rating Scale for Anxiety (HAM-A) scores of 20 or more, with a baseline average of 26.6 for 177 patients randomized to pregabalin and 26.1 for 96 patients on placebo. Patients with major depression and other anxiety or substance abuse disorders were excluded. The participants had three to four episodes of GAD on average, with the current episode lasting 15 months. The average age was 72 years, as about three-quarters of the population was 75 years of age or older. About three-fourths of those enrolled were female.

A flexible-dose regimen started at the active drug group on 150 mg per day of pregabalin, which clinicians could titrate up to 600 mg by week 6 of the 8-week trial. The average dose used was 270 mg.

Dr. Baldini suggested that the flexible dosing probably was responsible for the main difference in outcomes relative to the earlier adult trials. Whereas younger patients on pregabalin had a significantly better response at day 21, elderly patients on pregabalin began to show significant improvement compared with the control group during week 1.

By the elderly trial's end, 64.2% of the pregabalin cohort had responded versus 50% of the control group. Total HAM-A scores fell by more than 12 points in the pregabalin group, versus more than 10 points with placebo in a last-observation-carried-forward (LOCF) analysis.

The effect was seen in HAM-A psychiatric subscales: In the subscale for somatic symptoms, psychiatric symptoms fell by 7.8 points with pregabalin versus 6.3 points with placebo.

Elderly patients on pregabalin began to show significant improvement during week 2.

DR. BALDINI noted that the drug's benefits were significant in subgroups of patients with severe anxiety and with subsyndromal depression.

About a quarter of the patients—28%—of those on placebo and 25% of the pregabalin group—dropped out of the study. Only 7% on placebo and 4% on pregabalin did so for lack of efficacy. Adverse events caused discontinuation by 11% of the pregabalin group and 9% on placebo.

The most common adverse events with pregabalin were dizziness (20.3%), somnolence (13%), headache (10.2%) nausea (9%), and infection (5.6%).

An anticonvulsant with anxiolytic and anxiolytic properties, pregabalin is approved in the United States for treatment of neuropathic pain and seizures. It is also under study for fibromyalgia.

Occlusional Therapy Improves Daily Functioning in Dementia

Occlusional therapy improves the daily functioning of patients with mild to moderate dementia, results of a randomized, controlled trial show.

We believe that, in the long term, occupational therapy will result in less dependence on social and health care resources and less need for institutionalization," reported Maud J.L. Dopheide, assistant professor of Health Care at University Medical Center Nijmegen, the Netherlands, and her associates (BMJ 2006 Nov 17 [Epub ahead of print]).

In the study, dementia patients were randomized to an occupational therapy treatment group (68) or to a control group (67). Study participants had to be aged 65 years or older, diagnosed with mild or moderate dementia, and cared for on a week or more by a primary caregiver.

At 6 weeks, those in the intervention group had statistically significant improvements over the controls on three tests. Patients in the therapy group had a mean score that was 1 point higher on the 8-point motor and prehensile test and 1.9 points higher on the 44-point performance interview. Caregivers in the therapy group had a mean score 16.2 points higher on the 135-point compliance questionnaire.

One limitation of the study was that it was not double blinded.

—Jonathan Gardner