Basal ganglia calcification and psychosis in Down's syndrome

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Summary
A case of basal ganglia calcification (diagnosed in vivo) and schizophreniform psychosis occurring in a young adult with Down's syndrome is reported. A stress-vulnerability model is suggested. Because of the relatively high prevalence of basal ganglia calcification in Down's syndrome, this population appears well suited for systematic study of the neuropsychiatric aspects associated with this neurological condition.

KEY WORD: schizophrenia.

Introduction
There is growing interest in the association between calcification of the basal ganglia and schizophreniform psychoses (Lowenthal and Bruyn, 1968; Francis, 1979). Systematic study of the relationship of these disorders has been limited by the rarity of basal ganglia calcification (BGC). Two recent surveys (Koller, Cochrans, and Klawans, 1978; Murphy, 1979) demonstrated BGC in less than 0.5% of all computed tomographic (CT) scans reviewed. However, Malamud (1964) found BGC in 7% of individuals with Down's syndrome (DS), suggestive that this population may be useful for prospective study of the behavioural correlates of BGC. While the clinical significance of BGC in DS is not known, a case was recently reported (Jakab, 1978) demonstrating the coexistence of idiopathic BGC and psychosis in an 18-year-old female with DS. In this paper, a second consecutive young adult with DS admitted to our hospital (cf. Jakab, 1978) with psychosis and BGC demonstrated by computed axial tomography is described.

Case report
A 32-year-old woman with moderate mental retardation (WAIS IQ: 43) and trisomy G Down's syndrome was admitted for evaluation and treatment of a psychotic episode characterized by social withdrawal, paranoid ideation, and auditory hallucinations. She had been well at home and attending a daily activity programme until 6 months before admission, at which time she was allegedly raped by her sister's common-law husband. Shortly afterwards, she became progressively more withdrawn, ceased performing her household chores, stopped caring for herself, became incontinent of urine and stool, and would sit for hours staring into space. She began to speak out loud to herself in several distinctly different 'voices' and reported hearing commentary and derogatory auditory hallucinations. She became convinced that a man was watching her at all times and that this man planned to kill her. There was no previous personal or family history of neurological or psychiatric disorders.

On admission, she was an obese, short, lethargic woman with obvious features of DS. She appeared much older than her stated age and had faeces smeared over her abdomen. Her affect was anxious and she reported fearing that someone would harm her and to be hearing voices telling her that she was bad and going to die. She oriented to person and place, but not time, and could write her name. Physical examination was remarkable for perioral dermatitis, brachycephaly, epicanthal folds, Brusfield spots, gingivitis, protruding tongue, prominent abdomen, and short extremities.

Cardiovascular examination was unremarkable. Neurological evaluation revealed hyperactive deep tendon reflexes, bilateral ankle clonus, and bilateral palommental reflexes. Posture was stooped and gait was slow and wide-based, with diminished arm swing. No tremor, cogwheeling, paresis, or dyskinetic movements were noted. Cranial nerve and sensory examinations were unremarkable.

Laboratory investigations included complete blood count, SMA-12 chemistry screen, calcium and phosphorus levels, and thyroid function studies (all within normal limits). Chest X-ray and electrocardiogram were unremarkable. Electroencephalogram (EEG)
was mildly and diffusely abnormal with generalized slowing. Computer tomographic (CT) head scan demonstrated no evidence of cortical atrophy or ventricular dilatation; however, dense, bilateral basal ganglia calcification was found (Fig. 1).

![Computerized axial tomogram of the brain showing dense, bilateral basal ganglia calcification.](image)

**FIG. 1.** Computerized axial tomogram of the brain showing dense, bilateral basal ganglia calcification.

Treatment was begun with thioridazine, 200 mg at bedtime, and supportive, individualized activity therapy. The medication was tolerated without worsening of her mild extrapyramidal syndrome. During 2 weeks of inpatient treatment, her activity level and self-care improved, anxiety diminished, and she reported relief from her auditory hallucinations. She regained bowel and bladder continence. At time of discharge, she was alert, fully oriented, neatly dressed and groomed, and eager to return home. She was discharged under the care of her community mental health center and resumed her regular activities. Brief follow-up 6 months after discharged revealed no recurrence of psychosis.

**Discussion**

Basal ganglia calcification occurs in a wide range of clinical conditions, including hypoparathyroidism and pseudo-hypoparathyroidism (Babbitt et al., 1969), idiopathic familial degenerative diseases (Francis, 1979), and following cranial irradiation or intrathecal chemotherapy (Murphy, 1979). While rare in normal adults, the incidence of BGC increases with age, suggestive of age-related vascular changes (Murphy, 1979). Murofushi (1974) has postulated that the basal ganglia deposits in DS are due to cerebral circulatory insufficiency. Perhaps a critical factor in development of BGC is some impairment of the blood-brain barrier, whether from the toxic effects of irradiation or chemotherapy, ageing, micro-vasculature abnormality, or metabolic abnormality. However, in the present case, there was no evidence of any underlying process aetiologically related to the development of BGC.

Psychotic episodes occur in about 10% of individuals with DS, roughly one half the rate expected in the moderately to profoundly mentally retarded population overall (Gibson, 1978). However, such high rates of psychosis were determined in institutionalized samples and diagnoses were not based on explicit, operationalized criteria currently employed for classification of schizophreniform disorders. When prevalence of psychosis is determined in community-based samples of mentally retarded persons, and diagnoses made according to DSM-III criteria, prevalence of psychosis may be proportionally lower (cf. Eaton and Menolascino, 1982).

In Down's syndrome, psychosis is more likely to occur in individuals with severe or profound mental retardation, presenile dementia, or seizure disorder, all conditions associated with more severe neuropathological abnormality (Gibson, 1978). In the present case, an early form of the presenile dementia associated with DS (Ellis, McCullough and Corley, 1974; Thase, 1982) was considered in the differential diagnosis because of the patient's deterioration, apathy, development of incontinence, encephalopathic EEG, and the presence of several released reflexes. While this diagnosis could not be ruled out, the CT scan demonstrated no evidence of cortical atrophy and the patient demonstrated considerable symptomatic and social recovery with antipsychotic treatment.

In reporting the original case, Jakab (1978) did not suggest any causal connection between the presence of BGC and the development of psychosis. The occurrence of a second consecutive case with similar presentation and symptomatology is statistically unlikely and hence raises the index of suspicion about a possible relationship. In both the present case and the case described by Jakab (1978), the behavioural decompensation shortly followed a severe social stressor, and therefore BGC may represent a vulnerability factor rather than a primary aetiology. Although the association may be fortuitous, it is interesting that the two conditions have similar prevalence in DS, at least in institutionalized samples [BGC: 7% (Malamud, 1964); psychosis: 10% (Gibson, 1978)]. To date, no studies have been published utilizing CT scanning in the assessment of a DS population, and thus a possible association cannot yet be examined either prospectively or retrospectively. While further research is necessary to determine if BGC represents a specific risk factor for schizophreniform psychosis in DS, investigators interested in the neuropsychiatric manifestations of BGC should consider DS as a high risk population for systematic study.
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References


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