Fahr Syndrome – an Important Piece of a Puzzle in the Differential Diagnosis of Many Diseases

Krzysztof Jaworski1, Maria Styczyńska2, Monika Mandecka2, Jerzy Walecki3, Dariusz A. Kosior2

1 Department of Cardiology and Hypertension, Central Clinical Hospital of The Ministry of Interior, Warsaw, Poland
2 Mossakowski Medical Research Centre, Polish Academy of Sciences, Warsaw, Poland
3 Department of Radiology, Postgraduate Medical School, Warsaw, Poland

Author’s address: Dariusz A. Kosior, Department of Cardiology and Hypertension, Central Clinical Hospital of the Ministry of Interior, 137 Wolska Str., 02-507 Warsaw, Poland, e-mail: dkosior13@gmail.com

Summary

Fahr syndrome is a rare neurodegenerative disorder characterized by symmetrical, bilateral calcifications in the basal ganglia, nucleus gyrus and cerebral cortex. The continuous advancement as well as widespread use of brain imaging have contributed to the increasing detection rates of such changes. Nevertheless, their etiology is understood only partially and the methods of causative treatment are limited. Due to various symptoms, Fahr syndrome may resemble diseases from the field of neurology, psychiatry, cardiology and even urology. This article provides an up-to-date review of the literature concerning Fahr syndrome in terms of clinical practice.

MeSH Keywords: Basal Ganglia Diseases • Calcinosis • Tomography, X-Ray Computed

Background

In 1930, a German pathologist, Karl Theodor Fahr, described a case of a man with symmetrical calcifications of the basal ganglia and cerebral cortex [1]. The syndrome, named after this author, manifests with a wide range of neurological as well as psychiatric symptoms. The pathogenesis and clinical presentation of Fahr syndrome, known also as bilateral striatopallidodentate calcinosis (BSPDC) or Chavany-Brunhes syndrome, are partially understood, but there is still a lot to be discovered [2].

Etiology

Fahr syndrome is characterized histologically by foci of symmetrical, non-atheromatic compounds embedded in a protein-polysaccharide complex that are located within the globus pallidus, striatum, dentate nucleus, basal ganglia as well as within white and gray matter of the brain and cerebellum [2]. The deposits consist of calcium, zinc, iron, aluminum, magnesium, silicon, copper and phosphorus [3]. There are significant differences in the distribution of calcifications between individual patients. This may indicate the different mechanisms of their formation. The location within the brain structures as well as contact with blood vessel are the most important factors determining the chemical composition of the deposits. On this basis, two types of mineralization were described, non-vascular and perivascular [4].

Some reports describe the inheritance of Fahr syndrome, mainly in an autosomal dominant way. So far, four genes have been proved to be related to primary familial brain calcification; namely, SLC20A2, PDGFRB, PDGFB and XPR1 [5–12]. However, in the majority of patients, the syndrome does not have a genetic background. Bilateral basal ganglia calcifications can be observed in disorders of calcium and phosphorus metabolism, especially in hypoparathyroidism. However, the frequency of their occurrence is low [13,14]. Kulczycki et al. distinguished the following three types of biochemical abnormalities of calcium-phosphate homeostasis:

1. True hypoparathyroidism (low calcium and high phosphate serum concentration, increased cAMP and phosphate excretion in urine after stimulation with parathyroid hormone);
2. Pseudohypoparathyroidism (low calcium and high phosphate serum concentration, no increase in the excretion...
of cAMP and phosphate in urine after stimulation with parathyroid hormone); 3. Calcifications co-existing with normal serum calcium and phosphate levels, a significant increase in the excretion of cAMP after stimulation with parathyroid hormone and its relatively low phosphaturic effect [15,16].

According to Pronicka et al., resistance to parathyroid hormone may play a key role in some individuals with Fahr’s syndrome. It is manifested by a decreased phosphaturic effect of this hormone, while cAMP excretion in urine remains within a normal range. Interestingly, this response can be improved by beta-blockers, suggesting that Fahr’s syndrome may be a result of a defect of adrenergic receptors and their impaired relationship with parathyroid hormone receptors [17]. Another biochemical abnormality observed in Fahr’s syndrome is an increased alkaline phosphatase activity in the basal ganglia of the brain, while its level in serum remains normal. These changes may promote a precipitation of insoluble calcium phosphate salts in the nervous tissue [18]. In up to one-third of patients with Fahr’s syndrome, a close relationship between calcifications and impaired cerebral blood flow can be demonstrated [19]. Such patients present cerebral circulatory disturbances in the form of transient ischemic attacks or strokes. Perivascular positioning of calcifications may suggest a aberrant function of adventitia that could predispose to blood-brain barrier damage and deposition of calcium compounds. A major argument in favor of a vascular theory was given by the studies using brain flow scintigraphy that provided evidence of an impaired blood flow in calcification sites [20]. Moreover, inflammatory conditions, e.g. meningitis or encephalitis, are taken into account when considering the pathogenesis of Fahr’s syndrome [21]. Coexistence of Fahr’s syndrome with lupus erythematosus, monoclonal gammapathy and brain tumors, such as astrocytoma or pineal body gangliocytoma, was described in some case reports [22–25]. A significant role in the discussed processes may be attributed to homocarnosine, a specific antioxidant dipeptide found in the central nervous system. It is believed that the inhibition of calcification formation within subcortical structures is a task of homocarnosine [2].

The presented data do not encompass all clinical situations leading to the formation of intracranial calcification. Calcifications occur more often in elderly patients and may be symptoms of many diseases such as toxoplasmosis, cytomegalovirus, syphilis, tuberculosis, torulopsis, cystercerosis, tuberous sclerosis and others [26,27]. It is worth mentioning that calcifications are quite commonly found in clinical practice and may be encountered as physiological calcifications in the pineal gland or choroidplexuses as well as on the walls of blood vessels.

**Clinical Presentation**

Fahr’s syndrome has an insidious, slow and progressive course. The symptoms usually occur between the fourth and sixth decade of life, but they have been also reported in children as well as in young adults [28]. The clinical presentation varies depending on the time of onset. Thus, three forms of the syndrome were distinguished:

1. Early childhood onset – characterized by inhibition of mental development and early mortality;
2. Early onset (approximately at the age of 30) – characterized by psychiatric symptoms;
3. Late onset (approximately at the age of 50) – characterized by progressive dementia and movement disorders [29–32].

The most common symptoms of Fahr’s syndrome in adults include parkinsonism, dystonia, ataxia, chorea and extrapyramidal syndromes. Some patients may present with seizures and pyramidal disorders [33,34]. Psychiatric symptomatology typically includes cognitive impairment, schizophrenia-like psychosis, hallucinations, delusions, anxiety, mania, depression, hypoaesthesia as well as personality changes [35–38]. Headache, vertigo, stroke-like events, orthostatic hypotension, dysarthria, paresis, overactive bladder and syncope of unknown etiology have also been reported [39–41]. In children, Fahr’s syndrome may take the form of severe encephalopathy with dwarfism, microcephaly and optic nerve atrophy [28]. The patients who are under our observation complained of memory impairment, dyskinesia, recurrent loss of consciousness and urinary incontinence. It should be emphasized that only a small percentage of people with intracranial calcifications are symptomatic. In most cases, it is an accidental discovery without any clinical implications.

**Diagnosis**

Although the first radiographic descriptions appeared more than 50 years ago, a broad availability of computed tomography (CT) was a cornerstone for the detection of intracranial calcifications and increased the number of diagnosed Fahr’s syndrome cases [42]. Currently, CT is the most valuable method that surpasses magnetic resonance imaging (MRI) (Figure 1A, 1B) [43,44]. Toscano et. al. suggested that transcranial sonography can be a useful tool in imaging basal ganglia calcinoses [45]. Differential diagnosis is hampered significantly by the lack of precise criteria. In that respect, it is important to detect calcification foci greater than 800 mm² in surface area, regardless of their number [46]. Electroencephalography has a low diagnostic value, because all kinds of alterations in the central electrophysiological activity are possible [47]. Thus, neurological and psychological examinations as well as CT imaging remain the basic techniques for the diagnosis of Fahr’s syndrome.

The following diagnostic criteria were established:
1. Bilateral calcifications of the basal ganglia or other areas of brain on neuroimaging (may not apply to patients from families with Fahr’s syndrome);
2. Progressive neurological dysfunction and/or psychiatric symptoms;
3. Onset between the ages of 40–50;
4. Absence of biochemical abnormalities and somatic states suggestive of a metabolic or mitochondrial disease;
5. Absence of toxic, infectious or traumatic causes of intracranial calcifications;
6. Positive family history of Fahr’s syndrome and/or proved genetic background.
If a patient meets the last criterion, the diagnosis can be made without the presence of one of the first two criteria. In cases of a negative or unknown family history, other five criteria must be fulfilled [48–51]. Not all intracranial calcifications point towards Fahr’s syndrome. Calcium deposits with a thin, linear or cloudy pattern have a high-specificity. On the other hand symmetric micronodular or asymmetric unilateral calcifications are not characteristic of this disease [51].

Treatment

In most cases, treatment is symptomatic and includes antipsychotics, antidepressants, antiepileptic and pro-cognitive drugs. A distinctive feature of parkinsonian syndromes with concomitant calcifications is an outstanding resistance to the therapy with levodopa. This disorder more likely results from insensitivity of postsynaptic striatal structures rather than presynaptic damage observed in the primary parkinsonism [52].

Conclusions

Fahr’s syndrome has a rare radiological presentation of bilateral calcifications in the basal ganglia, nucleus gyrus and cerebral cortex. It may coexist with many symptoms resembling the diseases from the field of neurology, psychiatry, cardiology as well as urology. From a practical point of view, the most important issue is to differentiate the symptomatic and asymptomatic cases of Fahr’s syndrome and to detect abnormalities of calcium metabolism, primarily hypoparathyroidism, because their correction is the only causative treatment that can significantly improve symptoms and prognosis.

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