Case study

2	<mark>Familial idiopathic basal ganglia calcification</mark> (Fahr's syndrome): Initial clinical
3	neuropsychiatric presentation without corresponding neurological deficit
4	
5	Abstract
6	Familial idiopathic basal ganglia calcification (FIBGC) or Fahr's syndrome is a rare disorder
7	with various clinical presentations which can mimic in particular psychiatric illness. The
8	following case is characterized by the typical basal ganglia calcifications and presentation of
9	neuropsychiatric symptoms indicating the first clinical presentation in the absence of a
10	neurological deficit. As previously reported, the extent of calcification did not predict
11	neurological impairment, however, predicted severe psychosis.
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13	Keywords: Basal ganglia, calcification, Fahr's syndrome, neuropsychiatric deficits, sychosis -
14	Extrapyramidal symptoms
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16	Background
17	FIBGC or Fahr's disease or syndrome is a rare, neurological disorder characterized by
18	abnormal calcified deposits in the basal ganglia and cerebral cortex and typically affects
19	individuals in the 3 rd and 4 th decades of their lives [1].
20	Etiologically, this syndrome has been most commonly associated with endocrine disorders,
21	mitochondrial myopathies, dermatological abnormalities, and infectious diseases. The
22	understanding of the molecular genetics of this disorder remains limited.
23	Clinically, a range of symptoms including neurological symptoms such as extrapyramidal
24	symptoms, parkinsonism, chorea, or tremors to neuropsychiatric deficits of concentration and
25	memory have been described.

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27 Case Report

Mr. A. is a 57-year-old male patient with pituitary germinoma, s/p resection and radiation 28 29 therapy resulting in pituitary insufficiency and desmopressin substitution and no previous psychiatric history. The patient presented with loss of consciousness and myoclonic seizures, 30 31 followed by an altered mental state with hallucinations and delusions, as well as agitation and aggressive behavior. Police and ambulance were notified and the patient required sedation in 32 order to be taken to the emergency room. The initial presentation of symptoms occurred two 33 34 months before, the family reported episodes of unresponsiveness, disorientation, inability to use the computer and play video games. 35 36 In the emergency room, the patient continued to report hallucinations and delusions, 37 grimacing faces, angels and the Holy Ghost. He was concerned that his marriage was not going well, wary his wife could be wearing a mask. Furthermore, he was disoriented, anxious 38 and psychomotor retarded. 39 40 Initial laboratory findings revealed a discrete hyponatremia (125 mmol/l) from overadministration of desmopressin and a mildly increased creatinine kinase (320 U/l). The 41 complete blood count and electrolytes were within normal limits, including calcium, 42 phosphorus, and magnesium. Liver function tests including the alkaline phosphatase were 43 normal. Endocrinologically, parathormon was within normal limits. The cerebral spinal fluid 44 45 vielded no abnormalities. A computed tomography scan of the brain revealed profound calcifications of the basal 46

ganglia bilaterally (figure 1.). An MRI confirmed the post-ischemic changes in the right
superior parietal lobes and obstruction of the right carotid artery and collateralization of the
middle cerebral artery. An encephalographic study revealed discrete general changes and a
mild focus in the right temporal lobe. Neurologically, a documented right amaurosis and

discrete post-ischemic hemiparesis of the left lower extremity were present and no movementdisorder discovered.

The patient was admitted for further management and work-up. The hyponatremia was
corrected and over the following days, the neuropsychiatric symptoms remitted. The patient
was able to return home and follow-up was arranged.

56 Molecular genetic testing was not deemed necessary, as no family history existed.

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58 **Review of the literature**

The diagnostic criteria of FIBGC or Fahr's syndrome include 1- bilateral calcification of the 59 basal ganglia, 2- progressive neurologic dysfunction, 3- the absence of biochemical 60 61 abnormalities, an infectious, traumatic or toxic cause, and 4- a significant family history[1]. 62 The presentation of Fahr's syndrome varies and the diagnosis remains challenging. In adults, both loss of consciousness and seizures have been reported in patients with hypothyroid 63 hypocalcaemia [2]. Tetany occurs, which is difficult to distinguish from occasional 64 65 myoclonus caused by epileptic disorder. In addition, spasticity, gait disorder, speech impairment, dementia, parkinsonism, chorea, tremors, dystonia, myoclonus, and coma 66 manifest, as well as papilledema due to intracranial hypertension, CSF-pleocytosis, 67 paroxysmal choreoathetosis, even of the dystonic-choreoathetotic type [1]. 68 The prevalence of neurological symptomatology in Fahr's syndrome ranges from one third to 69 70 one half of patients [3,4]. The location and extent of lesions have an effect on the 71 manifestation, in particular in patients with dementia or patients with extrapyramidal symptoms, more extensive lesions cause more severe symptomatology [5]. 72 On the genetic basis, FIBGC or Fahr's disease is most commonly transmitted as an autosomal 73 74 dominant trait, however, is also transmitted as an autosomal recessive trait, or occurs

rs sporadically. A locus at 14q (IBGC1) is commonly involved, a second locus on chromosome

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76 8, and a third on chromosome 2. On the molecular level, on chromosome 8, a loss of function

77 mutation in the gene encoding type III sodium dependent phosphate transporter 2 (SLC20A2)

⁷⁸ has been thought to form the genetic basis for the pathophysiology of this disease. In

79 instances, in which no identifiable mutation or deletion in SLC20A2 can be found, PDGFRB

80 might be another sequence worthwhile evaluating [1].

81 The neuropsychiatric symptoms range from mild cognitive impairment to changes in

82 personality and behavior, to dementia and psychosis [1]. In rare presentations, frontotemporal

dementia, neurofibrillary tangles and calcification of the Fahr's type have been described,

84 however, neither extrapyramidal symptoms nor metabolic disorder occurred. In unusual types

85 of pre-senile dementia, imaging revealed calcareous depositions of Fahr's type and

86 Alzheimer's as well as frontotemporal dementia were ruled out. Severe compromised

87 attention and memory were reported in a patient with intact basic and higher motor function.

88 Thus, in these cases, neurological symptoms were not present [6].

Fahr's syndrome also presented with frontal lobe symptomatology; initially, uncontrollable 89 90 bursts of laughter and crying were noted and later dysarthria, as well as progressive changes in personality and behavior [7]. In another patient with disturbed selective attention and 91 cognitive flexibility, verbal perseverations, and declarative memory deficits, reduced glucose 92 uptake in PET scan was not only confined to the putamen and globus pallidus, but extended 93 to the bilateral temporal and parietal cortices, corresponding to the neuropsychological 94 deficits observed. Functional imaging revealed that the changes preceded cerebral atrophy in 95 Fahr's syndrome and reflected deficits in functional circuits involving the basal ganglia and 96 the frontal, parietal, and temporal lobes [8]. 97

98 The current understanding indicates that the extent of calcification does not predict

99 neurological impairment, however, predicts the neuropsychiatric disorders.

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Diagnostically, recommended imaging includes cranial CT or MRI and plain radiography of
the skull. Further investigations of interest include blood and urine testing for hematologic
and biochemical indices.

Calcification of the basal ganglia is an incidental finding in about 0.3%-1.5% of brain CT
scans, especially in elderly individuals. Microscopic calcifications have been observed in the
globus pallidus and dentate nucleus in up to 70% of autopsy series. However, calcifications
confined to this area usually do not cause clinical symptomatology [1].

107 Endocrine disorders, in particular parathyroid disturbances such as hypo- and

108 hyperparathyroidism have been most commonly associated with Fahr's syndrome and vitamin

D, crucial for the calcium metabolism and its homeostasis, has significant implications [1].

110 To date, no curative approach exists for Fahr's syndrome; as a consequence, management

111 strategies mainly focus on symptomatic relief and elimination of causative factors. Limited

112 evidence suggests that early diagnosis and management can reverse the calcification process

113 leading to complete recovery of mental functions. Various treatments have been administered

to Fahr's patients in an attempt to achieve stabilization and remission. These approaches base

on pathophysiological theories resulting in the proposal of small scale clinical experiences.

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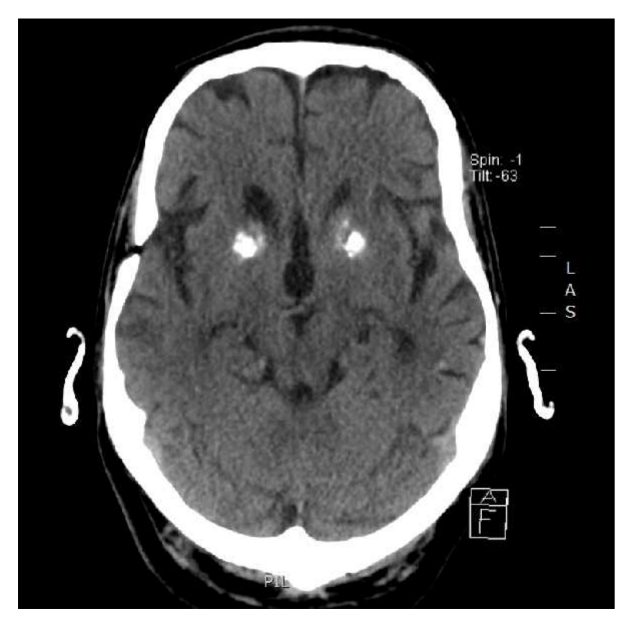
117 Conclusion

In summary, this case of FIBGC or Fahr's syndrome initially presenting with psychosis in the absence of neurological deficits, in particular movement disorder, and dementia, which, to date, has not yet been reported in the literature and adds to the evidence that the typical calcification predicts neuropsychiatric symptomatology in contrast to neurological deficits. Since other etiologies contributing to the presentation have been ruled out and the typical calcifications were present, this case illustrates the necessity for a heightened awareness of

- 124 potential Fahr's syndrome and the obligatory requirement of cranial imaging in order to
- 125 confirm the correct diagnosis.
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- 127 Conflict or interest: None
- 128 Patient Consent
- 129 The patient consented to the publication of this case
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132	References		
133	1.	Saleem S, Aslam HM, Anwar M, Anwar S, Saleem M, Saleem A, Rehmani	
134		MA.Fahr'ssyndrome: literaturereview of currentevidence. Orphanet J Rare Dis.	
135		2013;8:156.	
136	2.	Arias Mayorga J, González Martín T, Escorial Miguel C, Marañón Cabello A.	
137		[Intracranialcalcifications in the differential diagnosis of epilepticdisease]. Rev Clin	
138		Esp. 1991;189(9):425.	
139	3.	Kazis AD. Contribution of CT scan to the diagnosis of Fahr'ssyndrome. ActaNeurol	
140		Scand. 1985;71(3):206.	
141	4.	König P. Psychopathologicalalterations in cases of symmetricalbasalgangliasclerosis.	
142		Biol Psychiatry. 1989;25(4):459.	
143	5.	Taxer F, Haller R, König P. [Clinicalearlysymptoms and CTfindings in	
144		Fahrsyndrome].Nervenarzt. 1986;57(10):583.	
145	6.	Shibayama H, Kobayashi H, Iwase S, Nakagawa M, Marui Y, Kayukawa Y, Iwata H,	
146		Takeuchi T. Unusualcases of preseniledementia with Fahr'ssyndrome. Jpn J Psychiatry	
147		Neurol.1986;40(1):85.	
148	7.	Lam JS, Fong SY, Yiu GC, Wing YK. Fahr'sdisease: a differential diagnosis of frontal	
149		lobesyndrome. Hong Kong Med J. 2007;13(1):75.	
150	8.	Hempel A ¹ , Henze M, Berghoff C, Garcia N, Ody R, Schröder J. PETfindings and	
151		neuropsychologicaldeficits in a case of Fahr's disease. Psychiatry Res. 2001;108(2):133.	
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- 157 Figure 1. Cranial computed tomography. Profound calcifications of the basal ganglia
- bilaterally as typically seen in FIBGC or Fahr's syndrome.



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