Autoimmune polyendocrine syndrome type 1 (APS-1), which is also termed autoimmune polyendocrinopathy-candidiasis-ectodermal dystrophy (APECED), is a rare disorder and is inherited in an autosomal recessive fashion, caused by mutations in the autoimmune regulator gene (AIRE) and characterized by the failure of several endocrine glands as well as nonendocrine organs. APS type 1 appears to occur worldwide. Approximately 500 patients have been reported so far. APS type 1 has been noted to occur more frequently among homogeneous populations, such as those in Finland, northern Italy, Norway and Sardinia, as well as among Iranian Jews. The major components of APS type 1 are chronic mucocutaneous candidiasis, hypoparathyroidism and Addison’s disease. These are collectively called Whitaker’s triad, which is pathognomonic for APS type 1. The spectrum of associated minor clinical diseases include primary hypothyroidism, type 1 diabetes, nail dystrophy and dental enamel hypoplasia. In general, the first manifestation usually occurs in the childhood and the complete evolution of the three main diseases takes place in the first 20 year of age, whereas other accompanying diseases continue to appear until at least the fifth decade. In a majority of cases, candidiasis is the first clinical manifestation to appear, usually before the age of 5 year, followed by hypoparathyroidism, usually before the age of 10 year and later by Addison’s disease that occur usually before 15 year of age. Overall, the three main components of APS type 1 occur in a fairly precise chronological order, but they are present together in only about one third to one half of the cases. It has been reported that the earlier the first component appears, the more likely it is that multiple components will develop; conversely, patients who have late manifestations of the disease are likely to have fewer components. Therefore, patients with APS-1 should be followed-up on a regular basis because the majority of the above conditions develop later in the course of the disease. Classical diagnosis is made by presence of at least 2 major components or only one component if a sibling has already been diagnosed. Diagnostic criterion of having at least two elements of this triad would leave many cases missed. In some cases the minor components dominate with none of the triad present. If APS type 1 is suspected, genetic analysis of the AIRE gene may be helpful to confirm diagnosis, especially in atypical clinical presentations. Autoantibodies especially IFNα- and IFN-û are the diagnostic tool for APS type 1, especially in cases where mutational analysis is complicated (for example, large deletions, duplications, or mutations in regulatory or intronic regions).

Case Report
A 15 year old boy, hailing from Boalmari, Faridpur, was a known case of hypothyroidism for last 11 years with irregular intake of thyroxine. He was nondiabetic, normotensive and got admitted on 3rd April, 2013 in FMCH with repeated convulsion for last one month. Each of the attack started from his right hand and then became generalized and comprised a tonic and clonic phase. There
were 5-6 episodes of convulsion per day and each episode persisted for 5-10 minutes followed by loss of consciousness for about 30 minutes to 1 hour. Convulsion was associated with tongue bite and urinary incontinence and not associated with fever, headache and vomiting. He had no history of neck surgery, head injury or syncopeal attack. He developed recurrent skin infection in different parts of his body since childhood. He had a cataract surgery of his right eye five months back and developed blurring of vision in his left eye for few months. On examination he was ill looking with stunted growth and his body build was below average, mildly anaemic, pulse-76 beats/minute, blood pressure-90/60 mm of Hg. There was no postural drop of blood pressure. Trousseau’s sign was positive. His thyroid gland was not enlarged. Fundoscopy reveals normal (40.34 gm/dl) and random blood sugar (92mg/dl) were normal. CT scan of brain showed diffuse nearly symmetrical calcification of both dentate nuclei, both basal ganglia region, both thalami, both corona radiate, sub-cortical region of both frontal, parietal, temporal and occipital lobes. His complete blood count, serum creatinine, serum electrolytes, U.S.G of whole abdomen, E.C.G and chest x-ray P/A view findings were normal.

Fig. 1: Chronic mucocutaneous candidiasis

in right side and left side of fundus could not be visible due to early stage of cataract. Fungal infection was present in oral cavity and different parts of body like cheek and trunk. Nail dystrophy and dental enamel hypoplasia were present in almost all of the nails and teeth respectively. His bowel and bladder habit was normal.

Fig. 2: Dental enamel hypoplasia

His serum TSH level (75 µIU/ml) was high, serum PTH (<1pg/ml) and calcium (4.6 mg/dl) were low, serum albumin

Fig 3: Nail dystrophy

(40.34 gm/dl) and random blood sugar (92mg/dl) were normal. CT scan of brain showed diffuse nearly symmetrical calcification of both dentate nuclei, both basal ganglia region, both thalami, both corona radiate, sub-cortical region of both frontal, parietal, temporal and occipital lobes. His complete blood count, serum creatinine, serum electrolytes, U.S.G of whole abdomen, E.C.G and chest x-ray P/A view findings were normal.

Fig. 4: CT scan of brain showed diffuse nearly symmetrical calcification of both dentate nuclei, both basal ganglia region, both thalami, both corona radiate, sub-cortical region of both frontal, parietal, temporal and occipital lobes.
Discussion
The reported patient presented with hypothyroidism for last 11 years, chronic mucocutaneous candidiasis since childhood and features of hypoparathyroidism (convulsion for last one month, positive Trousseau’s sign and calcification in CT scan of brain). He had also nail dystrophy and dental enamel hypoplasia. Chronic mucocutaneous candidiasis (CMC) is the most common manifestation in APS-1. It is present in almost 100% of the patients. In most cases, CMC is the first of the major components of APS-1 to appear, often occurring before age 5, and its severity is variable. It preferably affects the oral mucosa causing a mild form of intermittent angular cheilitis. More severe cases include inflammation of most of the oral mucosa, hyperplastic CMC with thick white or grey plaques of yeast and hyperkeratosis, and atrophic form with thin mucosa and leuoplakic areas.10, 11 Our patient presented with chronic mucocutaneous candidiasis in oral cavity and different parts of body like cheek and trunk since childhood. CMC was present in a case report made by Kibirige and Kambugu.12

Another cardinal component, hypoparathyroidism, is usually the first endocrine disorder to develop during the course of APS I and it has been reported in 70%-93% of the cases.13 It varies according to gender, affecting 98% of female patients, but only 71% of male patients. Interestingly, when adrenal insufficiency is the first endocrinopathy, susceptibility to hypoparathyroidism appears to be reduced.14 Our patient had hypoparathyroidism as he presented with repeated attack of convulsion and positive Trousseau’s sign. We also found very low parathyroid hormone and calcium level in blood and CT scan showed diffuse calcification in brain. Hypoparathyroidism was also present in case report made by Kibirige and Kambugu.12

Our patient was a known case hypothyroidism for last 11 years. In APS type 1 hypothyroidism is relatively uncommon, affecting no more than 30% of the APS-1 patients. It develops more often following puberty and by middle age, usually before the age of 30.10 Hypothyroidism was present in case report made by Ming-Chen et al.15

Among the minor clinical disease the reported patient also presented with nail dystrophy and dental enamel hypoplasia. Primary adrenal insufficiency or Addison’s disease is present in 60%-100% cases, with peak incidence at around 12 years of age. It is a life threatening condition that should be rapidly recognized and treated. Symptoms are fatigue, weight loss, salt craving, hypotension, abdominal pain and increased pigmentation of the skin.13 Our patient had no feature of Addison’s disease. Addison’s disease was also absent in case report made by Kibirige and Kambugu.12

The presentation of our case was typical. He had two major components of classical triad and diagnosed as a case of APS type 1. Autoimmune polyendocrine syndrome type 1 (APS-1), is an extremely rare and frequently debilitating disorder of childhood. It is an autosomal recessive condition and a monogenic disorder linked to a defect of the AIRE gene located on chromosome 21q22.3.16 A clinical diagnosis of APS-1 classically requires the presence of two of the three cardinal components: chronic mucocutaneous candidiasis, hypoparathyroidism and Addison’s disease.6 Early immunogenetic testing for the different mutations of the AIRE gene and human leukocyte antigen (HLA) typing are essential for identification of patients at risk. This is because APS-1 occurs sporadically or in siblings11. HLA-A28 has been demonstrated to occur more frequently in patients with APS-1 than in normal controls. Generally, HLA-A3 is mostly observed in patients with APS-1.17 However, these tests are not readily available in most resource-limited settings.

Conclusion
Early recognition of multi-organ autoimmune disease is the best way to minimize associated morbidity and mortality. The aim of this case report is to grow awareness among health related personale about APS-1. A thorough history and physical examination should always be performed and a high index of suspicion should be maintained.

Conflict of Interest : None

References
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