3. No child had any chronic respiratory disease.
4. Sample collection of the cases was done following admission to the hospital and initiation of the investigation procedure. The first dose of empirical antibiotic therapy was already administered.
5. We admit the limitation of missing immunization history against Haemophilus influenzae B.
6. Follow-up of the patients as regarding the treatment course was not carried out in this study.

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Rubinstein-Taybi Syndrome with Psychosis

Rubinstein-Taybi syndrome is characterized by a broad thumb and bulbous hallux, short stature, intellectual disability and distinctive facial features [1]. It is a rare neuro-developmental disorder with a reported prevalence of 1 in 1,25,000 births [2]. Psychosis in RTS is highly infrequent with only a few scattered case reports [3]. A comprehensive literature search yielded only one case report of non-affective psychosis [4].

A 15-year-old girl was admitted to our department with spells of irritability and aggression for last 20 days. These episodes were accompanied by abnormal behavior like singing aloud and pacing. She appeared fearful, and was clinging to her mother. Upon detailed evaluation, there were no well-formed delusions and no clear-cut affective component could be distinguished. Therefore, a diagnosis of non-affective psychosis (Hallucinatory psychosis; ICD F28) was made. Behavioral problems were rated on the Brief Psychiatric Rating scale for Children (BPRS-C) on admission and 6 weeks later on follow-up.

Clinical examination showed short stature, with a height of 129 cm (below 50\textsuperscript{th} percentile). The thumbs were broad and flattened, as were the terminal phalanges of the other digits. The great toes were short and bulbous. There was microcephaly and typical facies, with a low hairline, hypertelorism, bushy eyebrows, broad nose and open mouth. Thoraco-lumbar scoliosis was noted. Muscle tone was low globally. Multiple keloids were present over the left scapular region and popliteal regions of both knees. Findings were consistent with a diagnosis of Rubinstein-Taybi syndrome.

Investigations revealed normocytic hypochromic anemia. MRI spine showed thoraco-lumbar scoliosis and decreased vertebral height. Intelligence Quotient on Binet-Kamat test gave a score of 57, indicating mild intellectual disability. Cyogenetic analysis by Giemsa showed a normal karyotype (46, XX). The patient was started on Quetiapine and recorded a reduction of more than 50\% in BPRS-C scores at 6 weeks on a dose of 50 mg, indicating significant response to therapy.

The association of psychosis with Rubinstein-Taybi syndrome is rare and only a handful of cases have been reported in literature. A novel study from Japan determined that variation in the promoter region of the same CREB gene may modify gene expression and contribute to schizophrenic psychosis [5]. The rare co-occurrence of psychosis in this syndrome thus opens up a narrow window of opportunity to identify the common genetic changes that result in this combined phenotypic manifestation. This, in turn, may generate fresh insight into the genetic markers of childhood psychosis.

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