Syndrome of Megalencephaly, Polydactyly, and Polymicrogyria Lacking Frank Hydrocephalus, with Associated MR Imaging Findings

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Megalencephaly, polymicrogyria, polydactyly, and hydrocephalus (MPPH) syndrome was first described in 2004, with only 9 cases reported so far; hence, the prevalence has not yet been determined.\textsuperscript{1,2} Hydrocephalus has been considered a mandatory component of this syndrome.\textsuperscript{1-4} We report a patient with macrocephaly, polymicrogyria, and polydactyly, but without overt hydrocephalus. We also report an elongated pituitary infundibulum along with a regressing cystic cavum septum pellucidum (CSP); cystic CSP has been described previously in MPPH, but without reports of regression.\textsuperscript{2,5,6} We discuss how to distinguish MPPH from other syndromes presenting with polymicrogyria or megalencephaly.

Case Report

A 1-day-old boy was born to a healthy nonconsanguineous family at 39-weeks’ gestation. Findings of prenatal sonography had suggested mildly dilated lateral and third ventricles, but with an otherwise unremarkable prenatal follow-up. After initially low Apgar scores but subsequent improvement with bag-mask ventilation, the patient was admitted to the neonatal intensive care unit with normal breathing. On physical examination, he was macrocephalic (head circumference >3.5 SDs above normal) with normal body length and weight, but with bilateral sixth digits along the ulnar aspects of the hands on plain films (Fig 1A). He had low-set ears, a high arching palate without cleft, and esotropia. No hypotonia/hypertonia was present, and deep tendon reflexes were normal. Results of blood biochemistry, cultures, and investigations for metabolic disease were unremarkable. Findings of cytogenetic analysis were unrevealing. Kidney and cardiac ultrasound findings were unremarkable.

Cranial sonography was then performed, which suggested subtle lateral ventricular dilation. Thereafter, MR imaging demonstrated diffuse polymicrogyria bilaterally, favoring the posterior frontal lobes and perisylvian regions (Fig 1B, -C). Additional findings included an elongated pituitary infundibulum without other sellar abnormalities.

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SUMMARY: Megalencephaly, polymicrogyria, polydactyly, and hydrocephalus (MPPH) syndrome has been recently recognized and is very rare. Each case reported so far has demonstrated hydrocephalus to varying degrees. We report an infant with MPPH syndrome, but lacking frank hydrocephalus. The additional finding of an abnormally elongated pituitary infundibulum has not been described in this syndrome and, along with the presence of a regressing cystic cavum septum pellucidum, suggests that chronic underlying hydrocephalus may have been present.

Discussion

MPPH syndrome was originally described in 2004 in 5 patients with megalencephaly, polymicrogyria, polydactyly, and hydrocephalus, along with severe psychomotor retardation, blindness, hypotonia, epilepsy, and facial dysmorphism.\textsuperscript{1} Subsequent reports have described phenotypic variation with facial dysmorphism of varying degrees (present in our patient); syndactyly, callosal, and midbrain abnormalities have also been described but were not present in our patient, who also had a later onset of epilepsy.\textsuperscript{1-4} Hence, he may have a milder phenotypic variant, given his later onset of seizures and appropriate psychomotor development to this point.

The finding of colpocephaly without overt hydrocephalus raises the question as to whether frank hydrocephalus is mandatory in MPPH. In the small number of patients reported with MPPH, hydrocephalus has been consistently described, along with macrocephaly, polymicrogyria, and polydactyly. The lack of definite hydrocephalus in our patient also suggests that he represents a milder variant. However, the infundibular elongation raises the question of chronic underlying hydrocephalus, possibly extending from an enlarged suprasellar cist-
tern; in such a situation, the ventricles may not dilate due to the elasticity of the infant skull with extracerebral communication of the hydrocephalus. However, it is difficult to discern whether overt hydrocephalus will ultimately develop without years of follow-up.

We noted 2 MR imaging findings in our patient that, to our knowledge, have not been described previously in MPPH syndrome: pituitary infundibular elongation and decreasing size of a cystic CSP. The elongation of the infundibulum is non-specific and again could relate to chronic low-level increased intracranial pressure. Regarding the cystic CSP, some reports in patients without MPPH have described intermittent hydrocephalus from its enlargement, with improvement or resolution following treatment.6 In MPPH syndrome, a cystic CSP has been described in all except 1 patient, a higher occurrence rate than that in the general infant population of 10%–40%.2,5,7,8 Hence, the cystic CSP in our patient could have related to chronic low-level hydrocephalus, with a decreased size from extraventricular decompression outside the ventricles, because the subarachnoid spaces along the convexities were more prominent on follow-up MR imaging. However, we cannot exclude the possibility that the cystic CSP is a developmental defect associated with MPPH.

Notably, given the polymicrogyria, other causes of seizures should also be investigated on MR imaging because the differential would include the symmetric perisylvian polymicrogyria syndromes, megalencephaly-polymicrogyria-mega-corpus callosum (MPMCC) syndrome, and macrocephaly-cutis-marmorata telangiectatica congenita syndrome (MCMTC). The presence of polydactyly and megalencephaly in our patient excluded the isolated symmetric perisylvian syndromes.9 Regarding MPMCC, a thickened corpus callosum is an essential feature, but was not present in our patient.10 Regarding MCMTC, which can have polydactyly, the absence of cutaneous hemangioma and the presence of polymicrogyria excluded this syndrome.11

No genetic alterations in MPPH have been found, and accordingly, our patient had no chromosomal abnormality. However, we did not perform a high-resolution cytogenetic examination, a potential limitation of this report. The presence of congenital metabolic disorders was largely excluded by negative findings on an extensive work-up via laboratory screening.

References

Fig 1. A–C, A neonate with macrocephaly and a sixth digit on plain film (arrow, A), who underwent MR imaging demonstrating diffuse polymicrogyria on T2WI (arrows, B and C). A prominent cystic CSP bows the cyst walls laterally (asterisk, C). D, The temporal horns are not overtly dilated on T2WI images (not shown), though the pituitary infundibulum is elongated (arrow), without dilated anterior third ventricular recesses, arguing against frank hydrocephalus (dotted arrows). E, At 8 months of age, the cystic CSP becomes much smaller (asterisk), but with enlarged subarachnoid spaces over the cerebral convexities. The arrows denote polymicrogyria in the midst of progressing myelination. F, The infundibulum is even more difficult to visualize, whereas the brain stem appears slightly small.


