Case Report

Hyperglycaemia-related complications at the time of diagnosis can cause permanent neurological disability in children with neonatal diabetes

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Abstract

Background  Children with neonatal diabetes often present with diabetic ketoacidosis and hence are at risk of cerebral oedema and subsequent long-term neurological deficits. These complications are difficult to identify because neurological features can also occur as a result of the specific genetic aetiology causing neonatal diabetes.

Case reports  We report two cases of neonatal diabetes where ketoacidosis-related cerebral oedema was the major cause of their permanent neurological disability. Case 1 (male, 18 years, compound heterozygous ABCC8 mutation) and case 2 (female, 29 years, heterozygous KCNJ11 mutation) presented with severe diabetic ketoacidosis at 6 and 16 weeks of age. Both had reduced consciousness, seizures and required intensive care for cerebral oedema. They subsequently developed spastic tetraplegia. Neurological examination in adulthood confirmed spastic tetraplegia and severe disability. Case 1 is wheelchair-bound and needs assistance for transfers, washing and dressing, whereas case 2 requires institutional care for all activities of daily living. Both cases have first-degree relatives with the same mutation with diabetes, who did not have ketoacidosis at diagnosis and do not have neurological disability.

Discussion  Ketoacidosis-related cerebral oedema at diagnosis in neonatal diabetes can cause long-term severe neurological disability. This will give additional neurological features to those directly caused by the genetic aetiology of the neonatal diabetes. Our cases highlight the need for increased awareness of neonatal diabetes and earlier and better initial treatment of the severe hyperglycaemia and ketoacidosis often seen at diagnosis of these children.

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What’s new?

- The cases in this report show that children with neonatal diabetes can sustain irreversible neurological damage as a result of complications of ketoacidosis at diagnosis and not just because of the neurological phenotype caused by genetic mutations.
- Both cases presented with diabetic ketoacidosis, required intensive care and probably developed cerebral oedema, leading to severe permanent spastic tetraplegia.
- The long-term neurological disability means they need intensive institutional or home support and so has a profound impact on their lives.
- Our cases highlight the need for increased awareness of neonatal diabetes and earlier and better initial treatment of the severe hyperglycaemia and ketoacidosis often seen at diagnosis of these children.

the major cause of their permanent neurological disability rather than the underlying genetic aetiology.

Case identification and assessment

Case identification

Our cases were identified from 125 people with neonatal diabetes (diabetes diagnosed at age <6 months, gestational age >32 weeks) referred from the UK to Exeter Molecular Genetic laboratory and confirmed to have monogenic aetiology for their diabetes before December 2015 (Table S1). There were three cases in whom persistent neurological features were severe and unexpected for the given mutation when compared with others in the cohort with the same mutation or mutations in the same gene and/or compared with the published literature. Initial identification of these cases was from the clinical information provided by the referring physicians based on patients’ health records and subsequently validated on clinical examination. The clinical details at the time of diagnosis of diabetes were reviewed from the hospital clinical records. This was available for two cases but not for the third case as the early clinical notes had been destroyed. The third case is therefore not reported.

Neurological and functional assessment

We performed detailed neurological examination on these cases and assessed their disability using the WHO Disability Assessment Schedule 2.0 (WHODAS 2.0) [11]. Scores in six different domains are calculated to give values 0 (no disability in daily living) to 100 (severe disability) and can be completed by carers or family members. Family members with the same mutation who did not present with severe diabetic ketoacidosis were assessed as controls for comparison.

Case report

DIABETIC Medicine

Case 1

Early history

Case 1 was born at term and had increased hunger and frequent wet nappies from 3 weeks of age (Fig. 1a). He was diagnosed with diabetes at 6 weeks when he presented with severe dehydration and diabetic ketoacidosis (pH 7.16, blood glucose 73.2 mmol/l). He had reduced consciousness, opisthotonus and partial seizures and subsequently aspirated. He was diagnosed as having cerebral oedema, required intensive care for 6 days and was hospitalized for 25 days. At 6 months he had motor developmental delay, bilateral spasticity and elevated muscle tone. A genetic diagnosis of a compound heterozygous ABCC8 p.P45L/p.G1401R mutation was made at the age of 8 years and he has been successfully treated with glibenclamide since then. He currently takes glibenclamide 5 mg twice a day and has an HbA1c of 34 mmol/mol (5.3%).

Current functional status

He is now 18 years old and attends a school for the physically disabled. He uses an electric wheelchair to mobilise and needs assistance for transfers, washing and dressing. On examination, there was marked weakness, increased tone with spasticity, brisk reflexes and contractures of all limbs. His language skills have developed normally. His total WHODAS 2.0 score is 57 (96th centile; Fig. 1b).

Family member control

The boy’s younger sister with diabetes has the same genetic mutation but was diagnosed by capillary blood glucose testing by her parents at 7 days of age before she had symptoms or diabetic ketoacidosis. She has normal development milestones and neurological examination results were normal (Fig. 1a).

Case 2

Early history

Case 2 was born at term. She had polyuria and thrush at 16 weeks of age (Fig. 1a). She was diagnosed with diabetes when she presented with reduced consciousness, generalized tonic-clonic seizures and diabetic ketoacidosis (pH 6.95, glucose 78 mmol/l). She was clinically treated for cerebral oedema and had recurrent seizures requiring intensive care for 2 weeks. One month after discharge she developed...
spastic tetraplegia (Fig. 1a). A genetic diagnosis of a heterozygous KCNJ11 p.S3C mutation was made at the age of 19 years. She was transferred from insulin to glibenclamide at the age of 27 years and currently takes glibenclamide 35 mg twice a day and has an HbA1c of 69 mmol/mol (8.5%).

Current functional status

She is now aged 29 years and lives in residential care. She requires assistance with all activities of daily living. She can say single words and nod her head for communication. Examination revealed spasticity and contractures of all limbs. Her total WHODAS 2.0 score is 78 (99th centile; Fig. 1b).

Family member control

The girl’s mother was diagnosed with diabetes aged 4 years after the detection of glycosuria on a routine urine test without diabetic ketoacidosis. The mother’s genetic diagnosis was established at 38 years of age after her daughter’s diagnosis. She has normal developmental milestones and examination results were normal (Fig. 1a).

Discussion

These cases highlight that neurological disability in neonatal diabetes can occur as a result of diabetic ketoacidosis-related complications at diagnosis in addition to being a direct result of underlying genetic aetiology.

Underlying genetic aetiology is not the major cause of the neurological disabilities seen in these cases. Both cases were extremely unwell at diagnosis with cerebral oedema, as shown by their biochemistry, clinical presentation and the need for intensive care. The family members with the same mutation but without diabetic ketoacidosis/cerebral oedema at presentation did not show any permanent neurological deficit. There were no other cases with KCNJ11 (n = 44, excluding the third case that we have not reported here) and ABCC8 (n = 16) neonatal diabetes who had similar neurological deficits in our cohort. Children with KCNJ11/ABCC8
neonatal diabetes commonly have hypotonia and coordination difficulties but not the spasticity and increased tone in all four limbs seen in our cases [12].

Cerebral oedema related to diabetic ketoacidosis is the likely mechanism for the neurological deficits in these cases. It is known that cerebral oedema attributable to diabetic ketoacidosis leads to permanent neurological features, particularly motor disability, in up to 35% of surviving children [2–4,13,14]. Both cases fit the clinical criteria for cerebral oedema [15] and spastic tetraplegia can result from diabetic ketoacidosis with cerebral oedema or other intracranial complications of diabetic ketoacidosis [13,14,16–18]. A case with KCNJ11 neonatal diabetes and neurological deficits attributable to cerebral oedema has been previously reported [9]. Although we do not have brain imaging studies for the cases at the time of diabetic ketoacidosis supporting the presence of cerebral oedema, this is not required to make a diagnosis, especially as 40% of cases with clinical cerebral oedema do not show acute abnormalities on their initial computed tomography examinations [15]; therefore, it has been proposed that cerebral oedema in the presence of diabetic ketoacidosis is a clinical, not a radiological, diagnosis [15]. The severity of disability in these cases may be attributable to late recognition of symptoms leading to severe ketoacidosis, the difficulty of neonatal fluid management and the vulnerability of the developing brain.

These two cases show that not all neurological deficits in people with neonatal diabetes are genetic in origin. This is particularly relevant to genetic aetiologies where there is a variable phenotype, such as KCNJ11 and ABCC8 neonatal diabetes in which ~20% of people have marked neurological features [8]; therefore, neurological features attributed to the genetic change may not be the correct explanation for a person’s condition. This information is also important for clinicians when communicating the diagnosis to patient and family members, particularly the likelihood of developing neurological disability in other family members who have inherited the same mutation.

In conclusion, long-term neurological morbidity in people with neonatal diabetes can occur independently of the underlying genetic mutation because of severe ketoacidosis-related complications at the time of diagnosis. Our cases highlight the need for increased awareness of neonatal diabetes and earlier and better initial treatment of the severe hyperglycaemia and ketoacidosis often seen at diagnosis of these children.

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Competing interests

None declared.

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References

1 Edge JA, Ford-Adams ME, Dunger DB. Causes of death in children with insulin dependent diabetes 1990–96. Arch Dis Child 1999; 81: 318–323.

2 Lawrence SE, Cummings EA, Gaboury I, Daneman D. Population-based study of incidence and risk factors for cerebral edema in pediatric diabetic ketoacidosis. J Pediatr 2005; 146: 688–692.

3 Edge JA, Hawkins MM, Winter DL, Dunger DB. The risk and outcome of cerebral oedema developing during diabetic ketoacidosis. Arch Dis Child 2001; 85: 16–22.

4 Glaser N, Barnett P, McCasdin I, Nelson D, Trainer J, Louise J et al. Risk factors for cerebral edema in children with diabetic ketoacidosis. The Pediatric Emergency Medicine Collaborative Research Committee of the American Academy of Pediatrics. N Engl J Med 2001; 344: 264–269.

5 Usher-Smith JA, Thompson MJ, Sharp SJ, Walter FM. Factors associated with the presence of diabetic ketoacidosis at diagnosis of diabetes in children and young adults: a systematic review. BMJ 2011; 343: d4092.

6 Jefferies C, Cutfield SW, Derraik JG, Bhagvandas J, Albert BB, Hofman PL et al. 15-year incidence of diabetic ketoacidosis at onset of type 1 diabetes in children from a regional setting (Auckland, New Zealand). Sci Rep 2015; 5: 10358.

7 Oyarzabal Irioyen M, García Cuartero B, Barrio Castellanos R, Torres Lacruz M, Gómez Gila AL, González Casado I et al. Ketoacidosis at onset of type 1 diabetes mellitus in pediatric age in Spain and review of the literature. Pediatr Endocrinol Rev 2012; 9: 669–671.

8 Hattersley AT, Ashcroft FM. Activating mutations in Kir6.2 and neonatal diabetes: new clinical syndromes, new scientific insights, and new therapy. Diabetes 2005; 54: 2503–2513.

9 Gloyn AL, Diatloff-Zito C, Edghill EL, Bellanné-Chantelot C, Nivot S, Coutant R et al. KCNJ11 activating mutations are associated with developmental delay, epilepsy and neonatal diabetes syndrome and other neurological features. Eur J Hum Genet 2006; 14: 824–830.

10 De Franco E, Flanagan SE, Houghton JA, Lango Allen H, Mackay DJ, Temple IK et al. The effect of early, comprehensive genomic testing on clinical care in neonatal diabetes: an international cohort study. Lancet 2015; 386: 957–963.

11 Ustun TB, Chatterji S, Kostanjsek N, Rehm J, Kennedy C, Epping-Jordan J et al. Developing the World Health Organization Disability Assessment Schedule 2.0. Bull World Health Organ. 2010; 88: 815–823.

12 Beltrand J, Elie C, Busiah K, Fournier E, Boddart N, Bahr-Buisson N et al. Sulfonylurea therapy benefits neurological and...
psichomotor functions in patients with neonatal diabetes owing to potassium channel mutations. *Diabetes Care* 2015; 38: 2033–2041.

13 Rosenbloom AL. Intracerebral crises during treatment of diabetic ketoacidosis. *Diabetes Care* 1990; 13: 22–33.

14 Marcin JP, Glaser N, Barnett P, McCaslin I, Nelson D, Trainor J et al. Factors associated with adverse outcomes in children with diabetic ketoacidosis-related cerebral edema. *J Pediatr* 2002; 141: 793–797.

15 Muir AB, Quisling RG, Yang MC, Rosenbloom AL. Cerebral edema in childhood diabetic ketoacidosis: natural history, radiographic findings, and early identification. *Diabetes Care* 2004; 27: 1541–1546.

16 Christodoulidou M, Selmi F. Severe diabetic ketoacidosis leading to cardiac failure, pulmonary oedema and spinal cord oedema resulting in tetraplegia. *BMJ Case Rep* 2012; 2012: bcr2012006769.

17 Dixon AN, Jude EB, Banerjee AK, Bain SC. Simultaneous pulmonary and cerebral oedema, and multiple CNS infarctions as complications of diabetic ketoacidosis: a case report. *Diabet Med.* 2006; 23: 571–573.

18 Shrier DA, Shibata DK, Wang HZ, Numaguchi Y, Powers JM. Central brain herniation secondary to juvenile diabetic ketoacidosis. *Am J Neuroradiol* 1999; 20: 1885–1888.

**Supporting Information**

Additional Supporting Information may be found in the online version of this article:

Table S1. Clinical characteristics and genetic aetiology of all 125 patients.