CASE REPORT

Transplant Neonatal Diabetes due to a Mutation in KCNJ11 in a Child with Klinefelter Syndrome

Amanda R. Dahl, Radhika Dhamija, Alaa Al Nofal, Siobhan T. Pittock, W. Frederick Schwenk, Seema Kumar

Introduction

Klinefelter syndrome (KS) is characterized by a 47XXY genotype and is the most frequent chromosomal aneuploidy in males occurring in about 1 in 660 males (1). Epidemiological studies have demonstrated increased risk of type 1 diabetes and type 2 diabetes in adults with Klinefelter syndrome. There is only one previous report of neonatal diabetes in a patient with Klinefelter syndrome.

Abstract

Klinefelter syndrome is the most frequent chromosomal aneuploidy in males occurring in about 1 in 660 males. Epidemiological studies have demonstrated increased risk of type 1 diabetes and type 2 diabetes in adults with Klinefelter syndrome. There is only one previous report of neonatal diabetes in a patient with Klinefelter syndrome. We report transient neonatal diabetes due to a pathogenic heterozygous variant in KCNJ11 in a male infant with Klinefelter syndrome. A 78-day old male infant was noted to have sustained hyperglycemia with serum glucose ranging between 148 mg/dL (8.2 mmol/L) and 381 mg/dL (21.2 mmol/L) three days after undergoing a complete repair of an atrioventricular defect. Hemoglobin A1c was 6.6%. The patient was born at term with a birth weight of 2.16 kg following a pregnancy complicated by gestational diabetes that was controlled with diet. The patient was initially started on a continuous intravenous insulin drip and subsequently placed on subcutaneous insulin (glargine, human isophane and regular insulin). Insulin was gradually decreased and eventually discontinued at seven months of age. Chromosomal microarray at 11 weeks of age showed XXY and a panel-based, molecular test for neonatal diabetes revealed a pathogenic heterozygous variant c.685G>A (p.Glu229Lys) in KCNJ11. The patient is now 34 months old and continues to have normal fasting and post-prandial glucose and HbA1C levels. The patient will need prospective follow up for assessment of his glycemic status. To our knowledge this is the second reported case of neonatal diabetes in an infant with Klinefelter syndrome. This case is the first due to a mutation in the KCNJ11 as the previously reported case of transient neonatal diabetes and Klinefelter syndrome had uniparental heterodisomy of chromosome 6.

Keywords: Neonatal diabetes, Klinefelter syndrome, KCNJ11

Conflict of interest: None declared

Received: 30.05.2017
Accepted: 31.07.2017

Address for Correspondence: Seema Kumar MD, Department of Pediatric and Adolescent Medicine, Division of Pediatric Endocrinology, Rochester, Minnesota, USA
Phone: +507-284-3300 E-mail: kumar.seema@mayo.edu ORCID ID: orcid.org/0000-0001-8004-4924

© Copyright 2018 by Turkish Pediatric Endocrinology and Diabetes Society
The Journal of Clinical Research in Pediatric Endocrinology published by Galenos Publishing House.
Neonatal diabetes is a monogenic form of diabetes, with an estimated incidence of 1 in 100,000 to 1 in 260,000 live births (8,9,10,11). It is defined as persistent hyperglycemia occurring in the first six months of life that lasts more than two weeks and requires insulin for management (9,12,13). Transient neonatal diabetes mellitus (TNDM) represents 50-60% of cases of neonatal diabetes, with the other 40-50% being permanent neonatal diabetes mellitus (14). The majority of cases of TNDM have paternal uniparental disomy of chromosome 6 or an unbalanced duplication of paternal chromosome 6 (15,16,17). Less frequent genetic abnormalities noted in patients with TNDM include activating mutations in the ATP-sensitive K+ channel encoding genes (KCNJ11 and ABCC8)(12). We report the first case of TNDM due to a mutation in the KCNJ11 gene in an infant with KS.

Case Report

A male infant was born at 39 + 3 weeks gestation via non-spontaneous vaginal delivery with a birth weight of 2.16 kg and length of 48 cm. Pregnancy was complicated by gestational diabetes that had been controlled with diet and labor was induced for intrauterine growth restriction. There were no complications in the neonatal period. External genitalia were normal. The infant was noted to have bilateral inguinal hernia at one month of age and underwent bilateral inguinal hernia repair at age 39 days. In the postoperative period, he developed tachypnea and respiratory distress. Physical examination was significant for a loud S1/S2 with no splitting of S2 and a harsh 2-3/6 systolic murmur, loudest at the left sternal border. Echocardiogram revealed a type A complete atrioventricular canal defect with moderate regurgitation of left and right sides of the common atrioventricular valve, multiple ventricular septal defects and a small secundum atrial septal defect. At age 42 days, a random blood glucose level was found to be 175 mg/dL and repeat blood glucose estimation on day 43 was 190 mg/dL. The mild hyperglycemia was attributed to stress. Furosemide and digoxin were started to improve acute heart failure. Subsequently, the infant exhibited poor weight gain and poor feeding. He underwent complete atrioventricular canal repair and secundum atrial septal defect closure at age 74 days. Postoperatively, on day 74, blood glucose values were noted to be between 148 mg/dL and 581 mg/dL. The patient was extubated by post-operative day three and weaned off vasopressors and inotropes by post-operative day four. Hyperglycemia persisted and hemoglobin A1C was noted to be 6.6%. C peptide was 2.7 ng/mL (reference range 1.1-4.4 ng/mL). Family history was significant for diabetes in father since his 20s who was successfully treated with metformin. Additionally, a maternal uncle had been diagnosed with diabetes at age 18 years and he was now on oral hypoglycemic agents previously.

Intravenous insulin infusion was started on day 78 at an initial dose of 0.03 units/kg/hour and was substituted with subcutaneous insulin injections (insulin glargine and regular insulin) at age 91 days. Though insulin glargine is not approved for use in infants, the release pattern of insulin glargine with no “peaks” makes it attractive during early infancy when infants are feeding frequently. There have been several reports regarding use of insulin glargine in infants with neonatal diabetes (18,19,20). Maximum hemoglobin A1c (HbA1C) was 7.2% at age five months. Glargine was subsequently replaced with intermediate acting isophane insulin due to insurance coverage reasons. Due to improvement in glucose values, insulin doses were gradually decreased and insulin was eventually discontinued at seven months of age. Oral sulfonylurea would be the preferred treatment in our patient given the KCNJ11 mutation. We planned to switch the patient from insulin to oral sulfonylurea after the results of the genetic analyses became available but elected not to do so when the insulin requirements began to decrease. We were able to discontinue insulin successfully with normal serum glucose values after discontinuation of insulin.

Fasting and post-prandial blood glucose values as well as HbA1C have subsequently been monitored and have been in the normal range. HbA1C was 6% at eight months of age, 6.2% at 11 months, 5.7% at 14 months, 5.4% at 19 months and 5.5% at 25 months of age. At the most recent follow up at 31 months of age, fasting glucose was 93 mg/dL and HbA1C was 5.4%. The patient has been gaining weight and growing normally and continues to make slow developmental progress. He started walking at 19 months of age and began receiving speech therapy for speech delay.

Genetic Analysis

Chromosomal microarray was performed at 11 weeks of age due to the complete atrioventricular canal defect (Cytogenetics Laboratories, Mayo Clinic, Rochester, MN). Chromosomal microarray revealed gain of the entire X chromosome. Limited chromosome study confirmed a diagnosis of KS/47, XXY. As external genitalia were unremarkable, gonadotropins and testosterone levels were not measured. The platform (Affymetrix CytoScan HD platform) used for microarray laboratory was single-nucleotide polymorphism based and did not detect stretches of homozygosity that would lead to possibility of uniparental disomy of chromosome 6. The laboratory was specifically asked to look for homozygosity at 6q24. Simultaneously, sequence analysis of 27 genes associated with neonatal...
diabetes was performed at the Genetic Services Laboratories of University of the Chicago (neonatal diabetes/Maturity Onset Diabetes of the Young sequencing panel). This panel based testing revealed a pathogenic heterozygous variant c.685G>A (p.Glu229Lys) in KCNJ11. There was also a heterozygous variant of unknown significance, c.713G>A (p.Arg238Gln) in BLK. Other genes included in the panel were ABC28, AKT2, CEL, CISD2, CP, EIF2AK3, FOXP3, GATP6, GCK, GLIS3, GLUD1, HADH, KCNJ11, KLF11, INSR, INS, IER3IP1, NEUROD1, NEUROG3, PAX4, PDX1, PTF1A, RFX6, SLC2A2, WFS1, and ZFP57.

Genetic testing for parents was recommended. The rationale for testing was discussed with the parents, as they may also have the KCNJ11 mutation detected in the patient and how this would influence their own medical care, but they have declined testing.

**Discussion**

We report the second case of TNDM in an infant with KS. Genetic testing in this patient revealed a pathogenic heterozygous variant in KCNJ11. This is the first reported case of neonatal diabetes due to a mutation in the KCNJ11 in a patient with KS.

The KCNJ11 encodes Kir6.2 subunit of the ATP-sensitive potassium channel in several tissues including pancreatic \( \beta \) cells, brain, heart and skeletal muscles (9,14). Mutations in KCNJ11 lead to a permanent opening of the potassium channel in the pancreatic \( \beta \) cells, thus preventing any activation of voltage-dependent calcium channel and glucose-induced insulin secretion leading to diabetes (21,22,23). The results of the genetic testing in this patient are noteworthy since heterozygous activating mutations in KCNJ11 are seen in only a small number of patients with TNDM (12) and instead comprise the most common cause of permanent neonatal diabetes (24,25,26). However, the pathogenic heterozygous variant c.685G>A (p.Glu229Lys) in KCNJ11 found in this patient has been previously associated with TNDM (27).

Mutations in KCNJ11 are de novo in 80 % of cases and inherited in an autosomal dominant pattern in the remaining cases (28). Family history in our patient was significant as his father had been diagnosed with diabetes since his 20s and he was doing well on metformin. Additionally, there was a history of gestational diabetes in the patient’s mother and diabetes requiring insulin therapy in a maternal uncle. Unfortunately, inheritance of the KCNJ11 variant in our child could not be determined as parents have declined testing. We continue to revisit with the parents with our recommendations on parental genetic analysis for better understanding of the significance of the KCNJ11 mutations in our patient. In contrast to our patient, the previously reported case of TNDM and KS had uniparental heterodisomy of chromosome 6 (7), the most common abnormality found in children with TNDM (7,15,16,17). Our patient did not have evidence for uniparental disomy on chromosome 6.

There are several similarities between our case and the previous report (7). Both infants had low birth weight as is expected in children with neonatal diabetes (8). The age at discontinuation of insulin was also similar, with insulin being discontinued at six months of age in the previously reported case and at seven months in our patient (7). Follow up data on both patients is limited and is available only until age two and a half years. These data suggest that there has been no recurrence of hyperglycemia. 40 % of patients with TNDM have recurrence of hyperglycemia later in life (8,17,29). Additionally, the hypogonadism associated with KS may lead to changes in body composition and a risk of developing metabolic syndrome and type 2 diabetes (2,6). Therefore, prospective follow up for assessment of glycemic status is warranted in our patient. If hyperglycemia were to recur, this patient would be a candidate for oral sulfonylurea therapy as oral sulfonylureas have been shown to result in improved glycemic control in patients with diabetes due to KCNJ11 mutations (9,23,30).

Another remarkable feature in our patient was the presence of an atrioventricular canal defect, ventricular septal defects and an atrial septal defect. Congenital cardiovascular anomalies are quite uncommon in children with KS. Those that have been reported include Tetralogy of Fallot and transposition of the great arteries (31,32). Parental genetic analyses will lead to better understanding of the significance of the KCNJ11 mutation.

**Ethics**

**Informed Consent:** Since this was a single case report, informed consent is not required by Mayo Clinic IRB guidelines.

**Peer-review:** External and internal peer-reviewed.

**Authorship Contributions**

Surgical and Medical Practices: Amanda Dahl, Seema Kumar, Concept: Amanda Dahl, Seema Kumar, Design: Amanda Dahl, Seema Kumar, Data Collection or Processing: Amanda Dahl, Radhika Dhamija, Seema Kumar, Analysis or Interpretation: Amanda Dahl, Radhika Dhamija, Seema Kumar, Alaa Al Nofal, Siobhan Pittock, Literature Search: Amanda Dahl, Seema Kumar, Writing: Amanda Dahl, Radhika Dhamija, Seema Kumar, Alaa Al Nofal, Siobhan Pittock, W Frederick Schwenk.
Financial Disclosure: The authors declared that this study received no financial support.

References

1. Groth KA, Skakkebaek A, Host C, Gravholt CH, Bojesen A. Clinical review: Klinefelter syndrome–a clinical update. J Clin Endocrinol Metab 2015;98:20-30.

2. Nielsen J, Johansen K, Yde H. Frequency of diabetes mellitus in patients with Klinefelter’s syndrome of different chromosome constitutions and the XXY syndrome. Plasma insulin and growth hormone level after a glucose load. J Clin Endocrinol Metab 1969;29:1062-1073.

3. Schattner A, Berrebi A. Klinefelter’s syndrome associated with autoimmune disease. J R Soc Med 1989;82:560.

4. Takeuchi Y, Murata Y, Sinsani J, Masukawa H, Nakamura R, Oi K, Kato Y, Niinomi M. Klinefelter’s syndrome accompanied by mixed connective tissue disease and diabetes mellitus. Intern Med 1999;38:875-881.

5. Bojesen A, Juul S, Birkebaek NH, Gravholt CH. Mortality in Klinefelter syndrome: a Danish register study based on hospital discharge diagnoses. J Clin Endocrinol Metab 2006;91:1254-1260.

6. Bojesen A, Kristensen K, Birkebaek NH, Fedder J, Mosekilde L, Bennett P, Laurberg P, Frystyk J, Flyvbjerg A, Christiansen JS, Gravholt CH. The metabolic syndrome is frequent in Klinefelter’s syndrome and is associated with abdominal obesity and hypogonadism. Diabetes Care 2006;29:1591-1598.

7. Felix TM, Tansey MJ, Patil SR, Murray JC, Dagle JM. Double paternal nondisjunction in an infant with transient neonatal diabetes mellitus and Klinefelter syndrome. Am J Med Genet A 2007;143A:895-898.

8. von Muhlendahl KE, Herkenhoff H. Long-term course of neonatal diabetes mellitus. N Engl J Med 1995;333:704-708.

9. Vaxillaire M, Bonnefond A, Froguel P. The lessons of early-onset monogenic diabetes for the understanding of diabetes pathogenesis. Best Pract Res Clin Endocrinol Metab 2012;26:171-187.

10. Vaxillaire M, D P, Bonnefond A, Froguel P. Breakthroughs in monogenic diabetes genetics: from pediatric forms to young adulthood diabetes. Pediatr Endocrinol Rev 2009;6:405-417.

11. Slingerland AS, Shields BM, Flanagan SE, Bruining GJ, Noordam K, Gach A, Mlynarski W, Malecki MT, Hattersley AT, Ellard S. Referral rates for diagnostic testing support an incidence of permanent neonatal diabetes in three European countries of at least 1 in 260,000 live births. Diabetologia 2009;52:1683-1685.

12. Aguilard-Bryan L, Bryan J. Neonatal diabetes mellitus. Endocr Rev 2008;29:265-291.

13. Polak M, Koralik M. Neonatal and very-early-onset diabetes mellitus. Semin Neonatol 2004;9:59-65.

14. Flanagan SE, Patch AM, Mackay DJ, Edghill EL, Gloy AL, Robinson D, Shield JP, Temple K, Ellard S, Hattersley AT. Mutations in ATP-sensitive K+ channel genes cause transient neonatal diabetes and permanent diabetes in childhood or adulthood. Diabetes 2007;56:1930-1937.

15. Herrmann R, Laine AP, Johansson C, Niederland T, Tokarska L, Dziatkowski H, Ionen J, Soltész G. Transient but not permanent neonatal diabetes mellitus is associated with paternal uniparental isodisomy of chromosome 6. Pediatrics 2000;105:49-52.

16. Temple IK, Shield JP. Transient neonatal diabetes, a disorder of imprinting. J Med Genet 2002;39:872-875.

17. Shield JP, Gardner R, Wadsworth E, Whiteford ML, James RS, Robinson DO, Baum JD, Temple IK. Aetiology and genetic basis of neonatal diabetes. Arch Dis Child Fetal Neonatal Ed 1997;76:F39-42.

18. Passanisi S, Timpanaro T, Lo Presti D, Mammi C, Caruso-Nicoletti M. Treatment of Transient Neonatal Diabetes Mellitus: Insulin Pump or Insulin Glargine? Our Experience. Diabetes Technol Ther 2014;16:880-884.

19. Barone JV, Tillman EM, Ferry RJ. Treatment of transient neonatal diabetes mellitus with subcutaneous insulin glargine in an extremely low birth weight neonate. J Pediatr Pharmacol Ther 2011;16:291-297.

20. Jeha GS, Venkatesh MP, Edelen RC, Kiensra KA, Karaviti L, Fernandes CJ. Neonatal diabetes mellitus: patient reports and review of current knowledge and clinical practice. J Pediatr Endocrinol Metab 2005;18:1095-1102.

21. McTaggart JS, Clark RH, Ashcroft FM. The role of the KATP channel in glucose homeostasis in health and disease: more than meets the islet. J Physiol 2010;588:3201-3209.

22. Klupa T, Skupien J, Malecki MT. Monogenic models: what have the single gene disorders taught us? Curr Diab Rep 2012;12:659-666.

23. Abbasi F, Saba S, Ebrahim-Habibi A, Sayahpour FA, Amiri P, Larjani B, Amoli MM. Detection of KCNJ11 gene mutations in a family with neonatal diabetes mellitus: implications for therapeutic management of family members with long-standing disease. Mol Diagn Ther 2012;16:109-114.

24. Gloy AL, Cummings EA, Edghill EL, Harries LW, Scott R, Costa T, Temple IK, Hattersley AT, Ellard S. Permanent neonatal diabetes due to paternal germline mosaic for an activating mutation of the KCNJ11 Gene encoding the Kir6.2 subunit of the beta-cell potassium adenosine triphosphate channel. J Clin Endocrinol Metab 2004;89:3932-3955.

25. Gloy AL, Pearson ER, Antcliff JF, Proks P, Bruining GJ, Slingerland AS, Howard N, Srivinasan S, Silva JM, Molnes J, Edghill EL, Frayling TM, Temple IK, Mackay D, Shield JP, Sumnik Z, van Rhijn A, Wales JK, Clark P, Gorman S, Eisenberg J, Ellard S, Bjornstad PR, Ashcroft FM, Hattersley AT. Activating mutations in the gene encoding the ATP-sensitive potassium-channel subunit Kir6.2 and permanent neonatal diabetes. N Engl J Med 2004;350:1838-1849.

26. Vaxillaire M, Populaire C, Busiah K, Cave H, Gloy AL, Hattersley AT, Czernichow P, Froguel P, Polak M. Kir6.2 mutations are a common cause of permanent neonatal diabetes in a large cohort of French patients. Diabetes 2004;53:2719-2722.

27. Edghill EL, Gloy AL, Goriely A, Harries LW, Flanagan SE, Rankin J, Hattersley AT, Ellard S. Origin of de novo KCNJ11 mutations and risk of neonatal diabetes for subsequent siblings. J Clin Endocrinol Metab 2007;92:1773-1777.

28. Naylor RN, Greeley SA, Bell GI, Philipson LH. Genetics and pathophysiology of neonatal diabetes mellitus. J Diabetes Investig 2011;2:158-169.

29. Shield JP, Baum JD. Transient neonatal diabetes and later onset diabetes: a case of inherited insulin resistance. Arch Dis Child 1995;72:56-57.

30. Pearson ER, Flechner I, Njolstad PR, Malecki MT, Flanagan SE, Larkin B, Ashcroft FM, Klimes I, Codner E, Iotova V, Slingerland AS, Shield J, Robert JJ, Holst JJ, Clark PM, Gorman S, Eisenberg J, Ellard S, Bjornstad PR, Ashcroft FM, Hattersley AT. Activating mutations in the gene encoding the ATP-sensitive potassium-channel subunit Kir6.2 and permanent neonatal diabetes. N Engl J Med 2006;355:467-477.

31. Swaminathan S, Goria SR, Barbour DS. Klinefelter syndrome in Association with Tertatoly of Fallet and Congenital Diaphragmatic Hernia. Pediatr Genet 2017;6:115-117.

32. Liu J, Moulic K, Mesia CI, Ge S, Obiri N, Anderson CE. Transposition of the great arteries in a neonate with Klinefelter syndrome- an incidental finding or a true association. Am J Med Genet A 2015;167A:268-270.