Title; Long-term follow-up of glycemic and neurological outcomes in an international series of patients with sulfonylurea-treated *ABCC8* permanent neonatal diabetes

Short running title; Outcomes of sulfonylurea treatment in ABCC8 PNDM

Authors; Pamela Bowman MBBS PhD^{1,2}, Frances Mathews MRES^{1,2}, Fabrizio Barbetti MD PhD^{3,4}, Maggie H. Shepherd PhD^{1,2}, Janine Sanchez MD⁵, Barbara Piccini MD⁶, Jacques Beltrand PhD^{7,8,9}, Lisa R. Letourneau-Freiberg, MPH, RD¹⁰, Michel Polak MD^{7,8,9}, Siri Atma W. Greeley MD PhD¹⁰, Eamon Rawlins², Tarig Babiker MRCP², Nicholas J. Thomas MRCP², Elisa De Franco PhD², Sian Ellard PhD², Sarah E. Flanagan PhD², Andrew T. Hattersley DM^{1,2}, for the Neonatal Diabetes International Collaborative Group.

- 1. Exeter NIHR Clinical Research Facility, Royal Devon and Exeter NHS Foundation Trust
- 2. Institute of Biomedical and Clinical Science, University of Exeter Medical School, Exeter
- 3. Dept. of Experimental Medicine, University of Rome Tor Vergata
- 4. Bambino Gesù Children's Hospital, IRCCS, Rome, Italy
- 5. Miller School of Medicine University of Miami, USA
- 6. Regional Center for Pediatric Diabetes, Meyer University Children's Hospital, Florence, Italy
- Service d'endocrinologie, gynécologie et Diabétologie pédaitrique, APHP centre, université de Paris, Paris France
- 8. INSERM U1016, Paris, France
- 9. Institut IMAGINE, Paris France
- 10. Kovler Diabetes Center, University of Chicago, Chicago, IL USA

Corresponding author;

Dr Pamela Bowman

Medical Research, Level 3 RILD, Barrack Road, Exeter, UK EX25DW

Telephone: 01392 408325 E-mail address: P.Bowman@exeter.ac.uk

Word Count: Abstract 251 Main text 2839

Figures and Tables (total 4); Figure 1 Parts A and B, Figure 2 Table 1, Table 2.

ABSTRACT

Objective

ABCC8 mutations cause neonatal diabetes that can be transient (TNDM) or less commonly permanent (PNDM); ~90% individuals can be treated with oral sulfonylureas instead of insulin. Previous studies suggested that people with *ABCC8*-PNDM require lower sulfonylurea doses and have milder neurological features than those with *KCNJ11*-PNDM. However, these studies were short-term and included combinations of permanent and transient forms of *ABCC8*-NDM. We aimed to assess the long-term glycemic and neurological outcomes in sulfonylurea-treated *ABCC8*-PNDM.

Research Design and Methods

We studied all 24 individuals with *ABCC8*-PNDM diagnosed in the UK, Italy, France or USA known to transfer from insulin to sulfonylureas before May 2010. Data on glycemic control, sulfonylurea dose, adverse effects including hypoglycemia, and neurological features were analysed using non-parametric statistical methods.

Results

Long-term data were obtained for 21/24 individuals (median follow-up 10.0 (4.1-13.2) years). 18/21 remained on sulfonylureas without insulin at most recent follow-up. Glycemic control improved on sulfonylureas (pre-sulfonylurea vs 1-year post-transfer HbA1c 7.2% vs 5.7%, p=0.0004) and remained excellent long-term (1-year vs. 10-year HbA1c 5.7% vs. 6.5%, p=0.04), n=16. Relatively high doses were used (1-year vs 10-year dose 0.37 vs 0.25mg/kg/day glyburide, p=0.50), without any severe hypoglycemia. Neurological features were reported in 13/21 individuals: these improved following sulfonylurea transfer in 7/13. The commonest features were learning difficulties (52%), developmental delay (48%), and ADHD (38%).

Conclusions

Sulfonylurea treatment of *ABCC8*-PNDM results in excellent long-term glycemic control. Overt neurological features frequently occur and may improve with sulfonylureas, supporting early, rapid genetic testing to guide appropriate treatment and neurodevelopmental assessment.

Introduction

The *ABCC8* gene encodes the SUR1 subunit of the pancreatic ATP-sensitive potassium (K_{ATP}) channel (1). SUR1 forms hetero-octameric complexes with Kir6.2, encoded by the *KCNJ11* gene (2). Mutations in K_{ATP} channel genes are the commonest cause of neonatal diabetes in non-consanguineous populations, with ~15-20% due to *ABCC8* mutations and ~25-30% due to *KCNJ11* mutations (3). Neonatal diabetes typically occurs in the first 6 months of life and can be permanent (PNDM), where diabetes persists lifelong, or transient (TNDM), where there is a period of remission of diabetes after 6-12 months followed by relapse in adolescence or early adulthood (4). *ABCC8* mutations cause TNDM in ~80% of cases and PNDM in ~20% cases; the opposite pattern is observed with *KCNJ11* mutations, which most frequently result in PNDM (5).

ABCC8 and *KCNJ11* mutations cause neonatal diabetes by preventing closure of pancreatic K_{ATP} channels in response to rising glucose (6, 7). This results in insulin deficiency, which historically required treatment with replacement doses of insulin. However, in ~90% of affected individuals sulfonylurea treatment can bypass the genetic defect by binding SUR1 and closing pancreatic K_{ATP} channels, promoting secretion of endogenous insulin and allowing patients to stop insulin injections and achieve excellent glycemic control and better quality of life (8-10). Sulfonylureas are the optimum treatment for *KCNJ11* PNDM long-term; in those patients who successfully transfer from insulin to sulfonylureas, metabolic control is maintained in >90% for at least 10 years with no serious adverse effects despite doses being ~2-10 times higher than those recommended in Type 2 Diabetes (T2D) (11). Short-term studies have suggested that lower doses of sulfonylurea are required to treat *ABCC8*-NDM in comparison to *KCNJ11*-NDM (9, 12). However, many previous studies contained both individuals with *ABCC8*-PNDM and *ABCC8*-TNDM, which have different clinical courses and treatment requirements (5, 6, 9). No study has assessed the long-term outcomes of sulfonylurea

ABCC8 and *KCNJ11* are both expressed in the brain as well as the pancreas (13, 14), therefore in addition to diabetes, central nervous system (CNS) features are observed in individuals with K_{ATP} channel mutations. These vary from the severe developmental delay, epilepsy and neonatal diabetes

(DEND) syndrome, to mild neuropsychological impairments detectable only on detailed neuropsychomotor testing (5, 15). In *KCNJ11*-PNDM, there is some correlation between the position of the variant in the protein and the clinical features. In contrast, in *ABCC8*-PNDM genotypephenotype relationships appear less distinct (16). In around half of individuals with *KCNJ11*-PNDM, sulfonylurea treatment results in partial improvement of the neurological features, which is thought to be due to the action of glyburide on K_{ATP} channels in the brain (11, 17).

Observational studies have suggested that the CNS features are not as common and /or severe in individuals with *ABCC8* mutations (5, 6, 18, 19), in comparison to those with *KCNJ11* mutations. However, as discussed above, previous research findings are based on cohorts containing both individuals with *ABCC8*-PNDM and *ABCC8*-TNDM. In addition, the majority of studies investigating the neurodevelopmental features associated with K_{ATP} channel mutations and the response of these features to sulfonylurea treatment have focused on patients with *KCNJ11*-PNDM.

Research relating to the CNS features in individuals with *ABCC8*-PNDM and the impact of long-term sulfonylurea treatment on both glycemic and neurological outcomes is crucial to establish and to inform clinical guidelines for this specific subtype.

Aim

To assess the long-term glycemic and neurological response to sulfonylureas in an international cohort of patients with PNDM due to *ABCC8* mutations.

Research Design and Methods

Patient cohort

Patients with a molecular genetic diagnosis of PNDM due to mutations in the *ABCC8* gene (NM_001287174.1) confirmed in laboratories in Exeter (UK), Rome (Italy), Paris (France) and Chicago (USA) known to transfer to sulfonylureas prior to 30th April 2010 with no period of remission of their diabetes were eligible for inclusion in the study (n=24). Three patients were lost to

follow-up in the first year after sulfonylurea transfer and were therefore excluded, leaving 21 patients with sufficient follow-up data (>4 years) for further analyses.

The study was conducted in accordance with the Declaration of Helsinki as revised in 2000. Patient data was collected during routine clinical care or through research surveys and was anonymised for use in the study. The study is registered with ClinicalTrials.gov, number NCT02624830.

Data Collection

Data were collected from the clinical records of participating patients or through research surveys completed by the participant or their carer(s). Data on glycemic control, sulfonylurea dose, and hypoglycemia, were collected before and after transfer from insulin to sulfonylureas, and annually until the most recent clinic follow-up. Clinicians and participants were asked to report side-effects or diabetes complications that occurred at any time point during the follow-up and if so to provide details about these.

Data on neurological features were collected before transfer from insulin to sulfonylureas, and after transfer at most recent follow-up. Clinicians or participants were specifically asked about the presence of developmental delay (DD), learning difficulties (LD), attention-deficit hyperactivity disorder (ADHD), epilepsy, sleep problems, muscle weakness, anxiety, autism, and spasticity as well as 'other' difficulties (11), and whether these features (if present) had improved on transfer to sulfonylureas.

Statistical Analysis

Data were analysed in Stata 16.0 using non-parametric statistical methods. Clinical characteristics of patients who remained on sulfonylurea alone were compared with those who required permanent reintroduction of insulin using the Mann-Whitney test for continuous data and 2-sample test of proportions for categorical data. For those patients who remained on sulfonylurea alone for the duration of the follow-up, paired data on metabolic control (HbA1c) and sulfonylurea dose were compared using the Wilcoxon signed-rank test. For those individuals with annual data available for

>50% of time points, longitudinal trends in HbA1c and sulfonylurea dose were plotted. Missing data were imputed as previously described (11).

Individuals who required insulin therapy only transiently or who were prescribed any other oral antidiabetic medication at any point in the follow-up were classified in the sulfonylurea only group. One individual (mutation, L1148R/R1380C) transferred from insulin to sulfonylureas twice aged 18 months (for 4 years) and again aged 26 years. His follow-up data relates to the second sulfonylurea transfer as no data were available after the first transfer 35 years ago.

For sulfonylureas other than glyburide, doses were converted to glyburide equivalent using percentage of maximum glibenclamide (glyburide) dose as per the British National Formulary (BNF) (20). Data are presented as median (range) unless stated otherwise. For all analyses, a p-value of less than 0.05 was used to denote statistical significance.

Results

Clinical characteristics

Clinical characteristics of the patients included in the study are shown in Tables 1 and 2.

Duration of follow-up

Median duration of follow-up was 10.0 (4.1-13.2) years, comprising a total of 205 patient years.

Sulfonylurea efficacy

At most recent follow-up, 18/21 (86%) patients remained on sulfonylurea therapy without insulin. For all three individuals who had restarted insulin, clinicians reported problems with adherence with prescribed medication and/or periods of loss to clinic/hospital follow-up. Clinical characteristics were similar between the individuals who remained on sulfonylureas vs. those who required reintroduction of insulin (Table 1).

Type of sulfonylurea prescribed

All patients who remained independent of insulin at most recent follow-up were prescribed glyburide

for the duration of the study. One patient was prescribed glyburide at initial transfer from insulin and was switched to tolbutamide at day 45 post-transfer (21); this individual subsequently required reintroduction of insulin therapy having stopped sulfonylurea treatment whilst lost to hospital follow-up.

Metabolic control and sulfonylurea dose

Paired data on pre-transfer HbA1c and HbA1c and sulfonylurea dose at year 1 (median 0.97, range 0.27-1.76 years) and year 10 (median 9.8, range 6.1-12.5 years) were available for 16 individuals. In these individuals, glycemic control improved on transfer to sulfonylurea [pre-transfer vs 1-year HbA1c 7.2 (5.3-9.5) vs 5.7 (5.0-7.3)% (55 (34-80) vs. 39 (31-56) mmol/mol), p=0.0004] and remained excellent at long-term follow-up [1-year vs. 10-year HbA1c 5.7 (5.0-7.3) vs. 6.5 (5.3-7.7)% (39 (31-56) vs. 48 (34-61) mmol/mol), p=0.04], figure 1a. High doses of sulfonylurea were used in most individuals [1-year vs 10-year dose 0.37 (0.01-1.25) vs 0.25 (0.03-1.30) mg/kg/glyburide, p=0.50], figure 1b. Only 3 individuals required doses under 0.1mg/kg/day glyburide at most recent follow-up. In 11 individuals who had sufficient annual data available, there was a gradual reduction in median sulfonylurea dose per kilogram of body weight over time despite relatively stable glycemic control, figure S1.

Side-effects

Diarrhea was reported in two individuals. One individual was diagnosed with irritable bowel syndrome (IBS). The second individual, previously reported by Codner et al, experienced transient diarrhea on glyburide; this stopped on switching to tolbutamide (21). No other adverse effects of sulfonylurea therapy were reported.

Hypoglycemia

There were no episodes of severe hypoglycemia, defined as losing consciousness or having seizures (22), reported over the course of the follow-up in patients treated with sulfonylurea alone. In one individual on glyburide treatment, an episode was reported whereby the blood glucose remained <4.0mmol/l even after treatment with fruit juice and third party assistance was required. Another

individual switched treatment from glyburide to tolbutamide (see above), due to episodes of asymptomatic hypoglycemia on glyburide treatment; these settled on tolbutamide (21).

Diabetes Complications

Microvascular complications occurred in 2 individuals; one had microalbuminuria, and one had microalbuminuria (normotensive) and proliferative retinopathy requiring intravitreal injections and photocoagulation, as well as mildly elevated LDL treated with a statin medication. These individuals transferred to sulfonylureas aged 10 years and 26 years (after a short period of sulfonylurea treatment as a child - see above). There were no macrovascular complications reported over the period of follow-up.

Body Mass Index (BMI)

In 10 individuals who remained independent of insulin and had paired height and weight data available, BMI remained normal [BMI standard deviation score (SDS) pre-sulfonylurea treatment - 0.13(-0.85-1.44) and at most recent follow-up on sulfonylureas -0.29 (-0.99-0.94), p=0.23].

Central Nervous System (CNS) Features

13/21 (62%) patients were reported to have CNS features before and after transfer to sulfonylurea, figure 3a and Table S1. The commonest features at most recent follow-up were developmental delay (DD) in 48%, learning difficulties (LD) in 52% and attention deficit hyperactivity disorder (ADHD) in 38%. Co-morbidity was common: 11 individuals had 3 or more specific CNS features together. .In five individuals, seizures were or may have been a result of factors other than the genetic mutation (Table S2).

In 7/13 (54%) there was some improvement (n=5) or complete resolution (n=2) noted in neurological features on starting sulfonylureas, figure 3a and Table S1. In the 2 patients whose neurological features completely resolved, both had DD pre-transfer (with LD in one case) but subsequently attained a developmental level expected for their age.

All 3 patients who required reintroduction of insulin treatment had neurological features (Table 1); there was improvement in the EEG background in one of these patients following initial transfer from insulin onto sulfonylureas. Age at transfer to sulfonylureas was similar in those patients with and without neurological features at most recent follow-up [6.6 (0.5-18.2 vs. 8.4 (0.9-26.0) years, p=0.53].

Discussion / Conclusions

In summary, in our 10-year study of 21 individuals with sulfonylurea-treated *ABCC8*-PNDM, 86% remained independent of insulin at their most recent follow-up. Furthermore, glycemic control was maintained on relatively high doses of sulfonylurea, without any reports of severe hypoglycemia or side effects. A large proportion of the cohort (62%) had overt neurological features both prior to sulfonylurea transfer and at most recent follow-up, and co-morbidity was common. There was partial improvement in some CNS features in just over half of individuals following transfer to sulfonylurea therapy.

The excellent long-term outcomes in sulfonylurea-treated *ABCC8*-PNDM are similar to those observed in *KCNJ11*-PNDM, Table S3 (11). Our data do not support the suggestion from previous studies that lower doses of sulfonylurea may be required in people with *ABBC8* mutations in comparison to those with *KCNJ11* mutations (9). This may be due at least in part to the inclusion of patients with transient and permanent forms of *ABCC8*-NDM in earlier cohort studies (9) and a lower sulfonylurea dose requirement in the patients with TNDM (23). In this study, behavioural and / or social factors are likely to explain the deterioration in glycemic control in the 3 individuals with *ABCC8*-PNDM who required reintroduction of insulin.

We have shown that overt CNS features in *ABCC8*-PNDM occur with relatively high frequency and that co-morbidity is common, in contrast with previous studies that included cohorts of patients with *ABCC8*-TNDM and PNDM (5, 6, 9). Indeed, the frequency and nature of CNS features in *ABCC8*-PNDM is similar to that observed in *KCNJ11*-PNDM, Table S3 (11), with the exception of autism which has been more frequently reported in *KCNJ11*-PNDM (11, 24, 25).

The partial improvement in neurological features in 7/13 individuals on transferring to sulforylureas is consistent with previous cases and suggests that the drugs may improve neurological function in some patients with ABCC8-PNDM, although a randomised controlled trial would be required to prove this definitively. It has been suggested the improvement in neurological features is greater the earlier sulfonylureas are started (17, 26), reflecting greater neuroplasticity at a younger age with a so-called 'sensitive period' occurring within the first 6 months of life (27). In our cohort, only 2 patients transferred under the age of 1 year and none transferred under the age of 6 months, therefore this crucial sensitive period for the action of sulfonylureas in the brain may have been missed. Studies in rats have suggested that glyburide is actively transported out of the brain (28); this may make it difficult to achieve therapeutic concentrations of the drug in the cerebrospinal fluid (CSF). As a result, recommended doses of glyburide in patients with KATP channel-related PNDM and severe neurological features are higher (at least ~1mg/kg/day) (29). Of those individuals who remained independent of insulin in our cohort, only 2 were on a dose this high at their most recent follow-up. Furthermore, there was a tendency for glyburide doses to fall over time, which may reflect lack of adjustment of total daily doses according to increases in body weight as children grow. This may also explain the slightly higher HbA1c at most recent follow-up when compared with year 1.

These factors emphasise the need for early genetic testing and identification of all patients with *ABCC8*-PNDM. Prompt genetic diagnosis facilitates early transfer to sulfonylureas, as well as systematic screening of all affected individuals for neurodevelopmental features at diagnosis and follow-up, and provision of appropriate support. Clinicians should consider higher doses of sulfonylureas if neurological features are present. They should also regularly adjust total daily dose to maintain the same dose per kilogram of body weight over time, thereby optimising treatment for both glycemia and neurological features.

Variable modes of inheritance are observed in our cohort; there is a mixture of dominant heterozygous *ABCC8* mutations (n=11) as well as compound heterozygous (n=7) and homozygous variants (n=3), in keeping with previous studies (30). This has implications for genetic counseling in relation to recurrence risk and carrier status in future offspring, which will be different for dominant vs recessive

inheritance. In this study, there were no differences in long-term outcomes between individuals with dominant heterozygous vs recessive (homozygous and compound heterozygous) mutations (Table 2). However, our study was not designed to address this question, and research in larger cohorts will be required to investigate the impact of mode of inheritance on clinical outcomes in *ABCC8*-PNDM. An additional limitation is the wide range of specific genetic variants included, which prevents the identification of strong genotype-phenotype relationships.

Finally, neurological features were not screened for systematically via repeated assessments in one center over the course of the follow-up, and comprehensive neuropsychological testing was not done as part of this study. Therefore, there is likely to be variable ascertainment and / or reporting of CNS features based on what was recorded in the clinical notes. This might result in an underestimation of the extent of neurological involvement in affected individuals. It is not possible to fully distinguish the relative contributions of mutant K_{ATP} channels in the brain and other factors to the neurological features will be due to a combination of different etiologies. Despite these limitations, this is the first study to assess the long-term treatment response and CNS features in an international cohort of patients with *ABCC8*-PNDM.

Further research in larger cohorts of individuals with *ABCC8* mutations will be required to investigate in more detail the CNS phenotype, genotype-phenotype relationships, and factors influencing the glycemic response to sulfonylureas e.g. specific physiological states such as puberty and pregnancy.

Conclusions

We have shown for the first time that sulfonylurea therapy is effective and safe long-term for people with PNDM due to *ABCC8* mutations, with excellent glycemic control maintained over 10 years without severe hypoglycemia or side effects, despite relatively high doses in most patients. Importantly, affected individuals frequently have multiple overt CNS features, which, in some, may partly improve with sulfonylureas. Rapid genetic diagnosis is crucial to facilitate early initiation of

precision therapy with sulfonylureas in *ABCC8*-PNDM, and enable prompt identification of neurodevelopmental features and provision of appropriate support for affected families.

Acknowledgements

Many thanks to the clinicians in the Neonatal Diabetes International Collaboration (Supplementary Appendix 1) for their contributions to data collection, and to the Exeter NIHR Clinical Research Facility for supporting the study. Many thanks to Professor Tamsin Ford for academic supervision of the first author during the study. Many thanks to Dr John Dennis for advice on statistical methods. Dr Pamela Bowman is the guarantor of this work and, as such, had full access to all the data in the study and takes responsibility for the integrity of the data and the accuracy of the data analysis.

Data access statement

The research data supporting this publication are provided within the manuscript and online supplemental material.

Funding

PB has a Sir George Alberti Clinical Research Training Fellowship funded by Diabetes UK (Grant Number 16/0005407). ATH is supported by a Wellcome Trust Senior Investigator award (Grant number 098395/Z/12/Z). SEF has a Sir Henry Dale Fellowship jointly funded by the Wellcome Trust and the Royal Society (Grant Number 105636/Z/14/Z). EDF has a RD Lawrence Fellowship funded by Diabetes UK (19/005971). MHS is a National Institute for Health Research (NIHR) Senior Nurse and Midwife Research Leader (NIHR4-SNMRL058) and is also supported by the Exeter NIHR Clinical Research Facility at the University of Exeter. The views expressed in this article are those of the author(s) and not necessarily those of the NIHR, or the Department of Health and Social Care. NJT is funded by a Wellcome Trust funded GW4 PhD. The University of Chicago Monogenic Diabetes Registry is supported by grants from the National Institute of Diabetes and Digestive and Kidney Diseases (R01DK104942 and P30DK020595).

Duality of Interest

None to declare.

Author Contributions

ATH, PB, FB, JB, MP, TB and NJT were involved in study design and protocol development. All authors were involved in data collection. EDF was involved in the genetic analysis. PB, FM, ER, ATH, and SEF were involved in data cleaning, analysis and interpretation. PB wrote the manuscript and all authors reviewed, critically revised and approved the manuscript for submission.

Figure Legends

Figure 1. Part A: HbA1c pre-sulfonylurea transfer, at year 1 and at most recent follow-up in 16 patients with data available at all 3 time points. Circles represent individuals and black horizontal lines represent group medians. Part B: Sulfonylurea dose at year 1 and at most recent follow-up in 16 patients included in Figure 1A. Circles represent individuals and black horizontal lines represent group medians.

Figure 2: Number of patients with *ABCC8*-PNDM with neurological features relative to sulfonylurea transfer. CNS = central nervous system.

Online Supplemental Material

Supplementary Appendix. Neonatal diabetes international collaboration.

Supplementary Table S1. Neurological features present before and after transfer to sulfonylureas, and features that improved on SU transfer.

Supplementary Table S2. Clinical details of patients in whom seizures / epilepsy may have been attributable to factors other than the genetic mutation.

Supplementary Table S3. Comparison of long-term outcomes in individuals with mutations in the *KCNJ11* and *ABCC8* genes.

Figure S1. HbA1c and sulfonylurea dose in 11 patients with at least 50% of annual longitudinal data

available for both variables. Data are presented as median (interquartile range). Missing data were

imputed as previously described (11).

References

1. Aguilar-Bryan L, Nichols CG, Wechsler SW, Clement JP, Boyd AE, Gonzalez G, et al. Cloning of the beta cell high-affinity sulfonylurea receptor: a regulator of insulin secretion. Science. 1995;268(5209):423.

2. Clement JP, Kunjilwar K, Gonzalez G, Schwanstecher M, Panten U, Aguilar-Bryan L, et al. Association and Stoichiometry of KATP Channel Subunits. Neuron. 1997;18(5):827-38.

3. 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(9997):957-63.

4. Flanagan SE, Patch AM, Mackay DJ, Edghill EL, Gloyn AL, Robinson D, et al. Mutations in ATPsensitive K+ channel genes cause transient neonatal diabetes and permanent diabetes in childhood or adulthood. Diabetes. 2007;56(7):1930-7.

5. Busiah K, Drunat S, Vaivre-Douret L, Bonnefond A, Simon A, Flechtner I, et al. Neuropsychological dysfunction and developmental defects associated with genetic changes in infants with neonatal diabetes mellitus: a prospective cohort study [corrected]. The lancet Diabetes & endocrinology. 2013;1(3):199-207.

6. Babenko AP, Polak M, Cave H, Busiah K, Czernichow P, Scharfmann R, et al. Activating mutations in the ABCC8 gene in neonatal diabetes mellitus. N Engl J Med. 2006;355(5):456-66.

7. Gloyn AL, Pearson ER, Antcliff JF, Proks P, Bruining GJ, Slingerland AS, et al. Activating mutations in the gene encoding the ATP-sensitive potassium-channel subunit Kir6.2 and permanent neonatal diabetes. N Engl J Med. 2004;350(18):1838-49.

8. Pearson ER, Flechtner I, Njolstad PR, Malecki MT, Flanagan SE, Larkin B, et al. Switching from insulin to oral sulfonylureas in patients with diabetes due to Kir6.2 mutations. N Engl J Med. 2006;355(5):467-77.

9. Rafiq M, Flanagan SE, Patch AM, Shields BM, Ellard S, Hattersley AT, et al. Effective treatment with oral sulfonylureas in patients with diabetes due to sulfonylurea receptor 1 (SUR1) mutations. Diabetes care. 2008;31(2):204-9.

10. Shepherd M. Transforming lives: transferring patients with neonatal diabetes from insulin to sulphonylureas. European Diabetes Nursing. 2006;3(3):137-42.

11. Bowman P, Sulen A, Barbetti F, Beltrand J, Svalastoga P, Codner E, et al. Effectiveness and safety of long-term treatment with sulfonylureas in patients with neonatal diabetes due to KCNJ11 mutations: an international cohort study. Lancet Diabetes Endocrinol. 2018.

12. Proks P. Neonatal diabetes caused by activating mutations in the sulphonylurea receptor. Diabetes Metab J. 2013;37(3):157-64.

13. Ashcroft FM. ATP-sensitive potassium channelopathies: focus on insulin secretion. The Journal of clinical investigation. 2005;115(8):2047-58.

14. Sakura H, Ammala C, Smith PA, Gribble FM, Ashcroft FM. Cloning and functional expression of the cDNA encoding a novel ATP-sensitive potassium channel subunit expressed in pancreatic beta-cells, brain, heart and skeletal muscle. FEBS Lett. 1995;377(3):338-44.

15. Gloyn AL, Diatloff-Zito C, Edghill EL, Bellanne-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(7):824-30.

16. Edghill EL, Flanagan SE, Ellard S. Permanent neonatal diabetes due to activating mutations in ABCC8 and KCNJ11. Rev Endocr Metab Disord. 2010;11(3):193-8.

17. Beltrand J, Elie C, Busiah K, Fournier E, Boddaert N, Bahi-Buisson N, et al. Sulfonylurea Therapy Benefits Neurological and Psychomotor Functions in Patients With Neonatal Diabetes Owing to Potassium Channel Mutations. Diabetes care. 2015;38(11):2033-41.

18. Hashimoto Y, Dateki S, Hirose M, Satomura K, Sawada H, Mizuno H, et al. Molecular and clinical features of KATP -channel neonatal diabetes mellitus in Japan. Pediatric diabetes. 2017;18(7):532-9.

19. Aittoniemi J, Fotinou C, Craig TJ, de Wet H, Proks P, Ashcroft FM. Review. SUR1: a unique ATP-binding cassette protein that functions as an ion channel regulator. Philosophical transactions of the Royal Society of London Series B, Biological sciences. 2009;364(1514):257-67.

20. <u>https://bnf.nice.org.uk/drug/glibenclamide.html</u>. [

21. Codner E, Flanagan SE, Ugarte F, Garcia H, Vidal T, Ellard S, et al. Sulfonylurea treatment in young children with neonatal diabetes: dealing with hyperglycemia, hypoglycemia, and sick days. Diabetes Care. 2007;30(5):e28-9.

22. Ly TT, Maahs DM, Rewers A, Dunger D, Oduwole A, Jones TW, et al. ISPAD Clinical Practice Consensus Guidelines 2014. Assessment and management of hypoglycemia in children and adolescents with diabetes. Pediatric diabetes. 2014;15 Suppl 20:180-92.

23. <u>https://www.diabetesgenes.org/about-neonatal-diabetes/transferring-patients-who-have-a-mutation-in-kcnj11-or-abcc8/</u>.

24. Bowman P, Broadbridge E, Knight BA, Pettit L, Flanagan SE, Reville M, et al. Psychiatric morbidity in children with KCNJ11 neonatal diabetes. Diabet Med. 2016;33(10):1387-91.

25. Svalastoga P, Sulen A, Fehn JR, Aukland SM, Irgens H, Sirnes E, et al. Intellectual Disability in KATP Channel Neonatal Diabetes. Diabetes Care. 2020;43(3):526-33.

26. Shah RP, Spruyt K, Kragie BC, Greeley SA, Msall ME. Visuomotor performance in KCNJ11related neonatal diabetes is impaired in children with DEND-associated mutations and may be improved by early treatment with sulfonylureas. Diabetes Care. 2012;35(10):2086-8.

27. Zeanah CH, Gunnar MR, McCall RB, Kreppner JM, Fox NA. Sensitive Periods. Monographs of the Society for Research in Child Development. 2011;76(4):147-62.

28. Lahmann C, Kramer HB, Ashcroft FM. Systemic Administration of Glibenclamide Fails to Achieve Therapeutic Levels in the Brain and Cerebrospinal Fluid of Rodents. PLoS One. 2015;10(7):e0134476-e.

29. <u>https://www.diabetesgenes.org/about-neonatal-diabetes/effects-of-sulphonylurea-on-the-brain/</u>.

30. Ellard S, Flanagan SE, Girard CA, Patch AM, Harries LW, Parrish A, et al. Permanent neonatal diabetes caused by dominant, recessive, or compound heterozygous SUR1 mutations with opposite functional effects. American journal of human genetics. 2007;81(2):375-82.

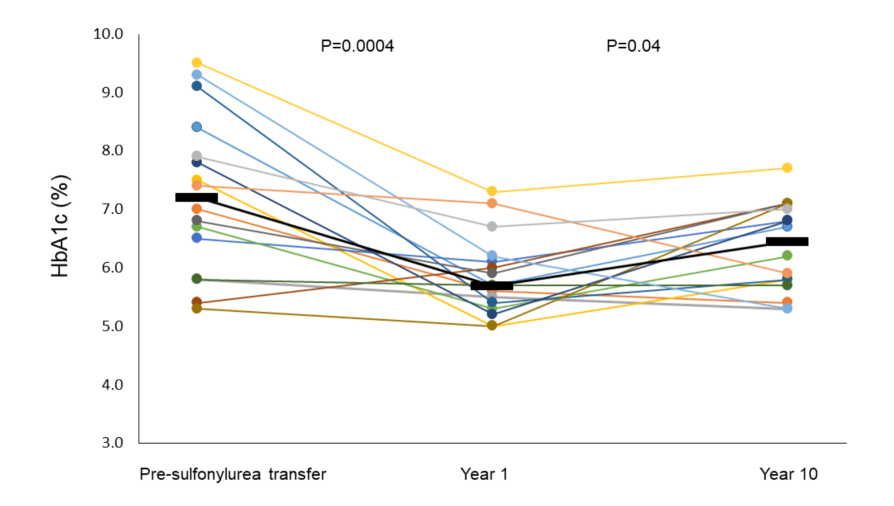
31. Day JO, Flanagan SE, Shepherd MH, Patrick AW, Abid N, Torrens L, et al. Hyperglycaemiarelated complications at the time of diagnosis can cause permanent neurological disability in children with neonatal diabetes. Diabet Med. 2017;34(7):1000-4.

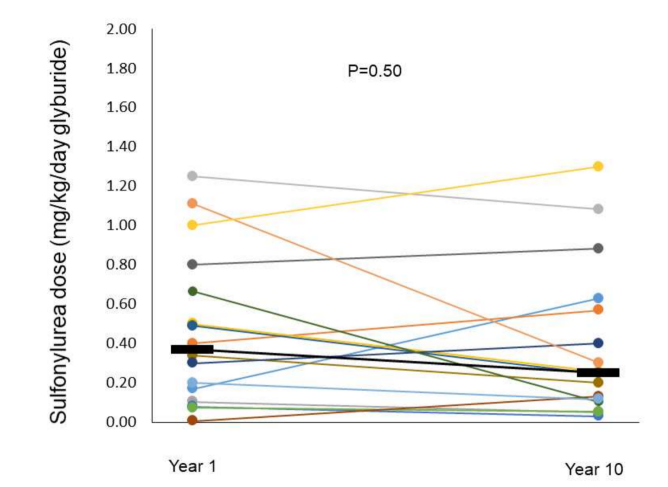
Clinical feature	On SU without insulin at most recent follow-up (n=18)	On insulin with or without SU at most recent follow-up (n=3)	P-value (on SU vs on insulin)	
Genotype	E208K/Y263D, V86G/N, D212I/N, P45L/G1401R, V86A/N, L1295F/N, T229I/V1532L, L225P/N, L135P/N, 2 E382K/E382K, L213R/N, R168C/G1256S, V215A/V215A, E208K/D1472N, V324M/W688R, L213P/N, L1148R/R1380C	D209E/N, R1380L/N, Q211K/N	N/A	
N male (%)	10 (56)	0 (0)	0.07	
Birth weight (g)	2750 (1510-3402) (n=16)	2700 (2400-2700) (n=3)	0.62	
Age at diagnosis (weeks)	7.5 (1.0-47.0) (n=18)	6.0 (5.0-17.0) (n=3)	1.00	
Age at transfer to SU (years)	7.6 (0.5-26.0) (n=18)	5.8 (2.8-9.4) (n=3)	0.46	
Current age (years)	21 (11-36) (n=18)	19 (16-22) (n=3)	0.58	
Pre-SU HbA1c (%)	7.5 (5.3-9.5) (n=18)	6.8 (6.7-7.2) (n=3)	0.52	
Pre-SU HbA1c (mmol/mol)	58 (34-80) (n=18)	51 (50-55) (n=3)	0.52	
Year 1 HbA1c (%)	5.7 (5.0-7.3) (n=16)	6.8 (6.7-6.9) (n=2)	0.13	
Year 1 HbA1c (mmol/mol)	39 (31-56) (n=16)	51 (50-52) (n=2)	0.13	
Most recent HbA1c (%)	6.7 (5.3-10.1) (n=18)	9.8 (5.9-10.3) (n=3)	0.17	
Most recent HbA1c (mmol/mol)	50 (34-87) (n=18)	84 (41-89) (n=3)		
Year 1 SU dose (mg/kg/day glyburide)	0.37 (0.01-1.25) (n=16)	0.03 (0.01-0.04) (n=2)	0.05	
Most recent SU dose (mg/kg/day glyburide)	0.35 (0.03 – 1.30) (n=18)	0.74 (0.02-1.45) (n=2)	1.00	
Neurological features (any)	10 (56)	3 (100)	0.15	
Pre-SU BMI SDS (kg/m2)	-0.13 (-0.85-1.44) (n=10)	-0.53 (-1.680.26) (n=3)	0.10	
Most recent BMI SDS (kg/m2)	-0.75 (-3.69 – 1.28) (n=13)	-1.34 (-1.940.24) (n=3)	0.19	

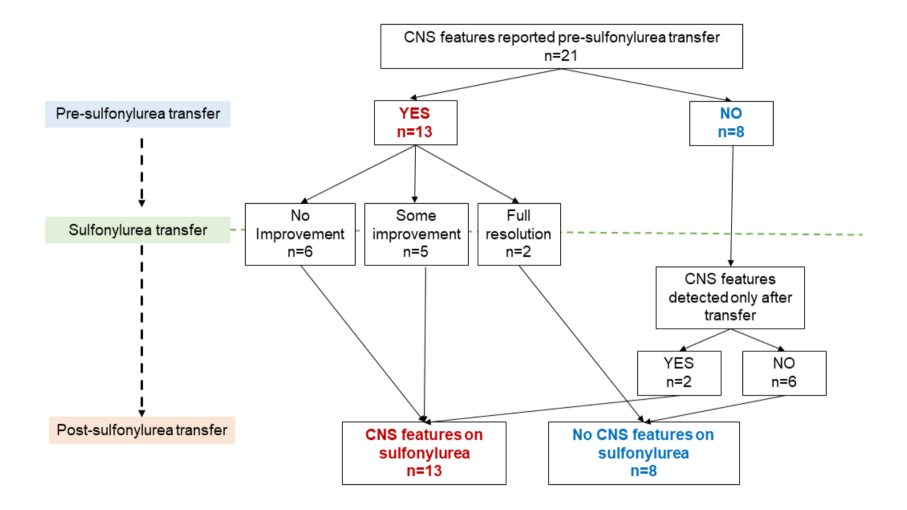
Table 1. Clinical features of whole cohort, including all data available at all time points. Data are presented as median (range). Year 1 is median duration 0.97 (0.27-1.76 years); data closest to one year used. Most recent duration for sulfonylurea dose and HbA1c median 10.0 (4.1-13.2) years. Most recent duration for BMI 9.8 (4.6-13.2) years (most recent time point at which both height and weight data available). N is different for each variable due to differences in amount of available data. Most recent sulfonylurea dose is different to that reported in the results section due to paired values for sulfonylurea dose and HbA1c at all time points being unavailable for 3 individuals included in the table. SU = sulfonylurea, BMI = body mass index, SDS = standard deviation score.

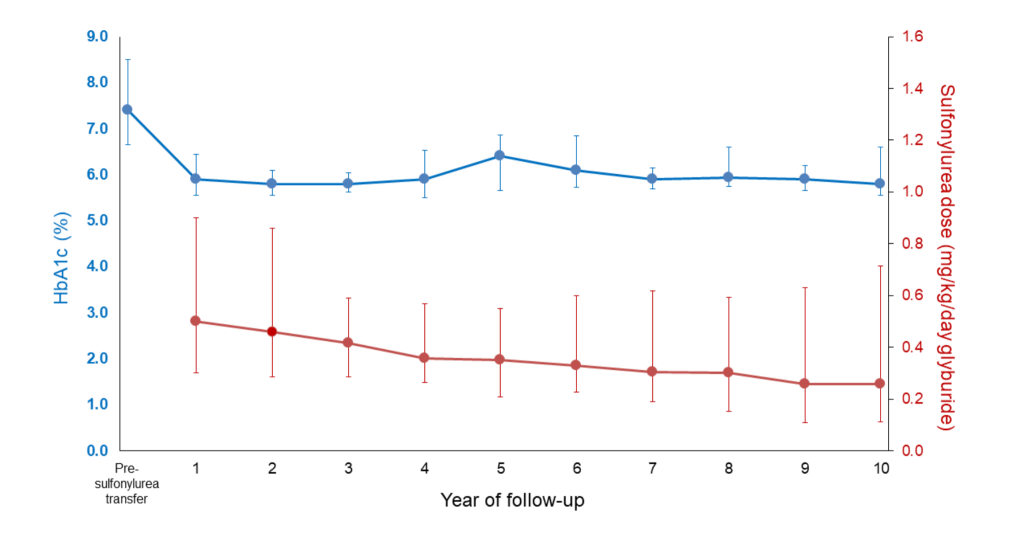
Clinical feature	Dominant heterozygous (n=11)	Recessive (compound heterozygous / homozygous) (n=10)	P-value
Genotype	D209E/N, D212I/N, L1295F/N, L135P/N, L213P/N, L213R/N, L225P/N, Q211K/N, R1380L/N, V86G/N, V86A/N	2 E382K/E382K, E208K/D1472N, E208K/Y263D, L1148R/R1380C, P45L/G1401R, R168C/G1256S, T229I/V1532L, V215A/V215A, V324M/W688R	N/A
Duration of follow-up (years)	11.2 (7.9-13.2)	9.1 (4.1-12.3)	0.02
N male (%)	4/11 (36)	6/10 (60)	0.27
Birth weight (g)	2700 (2100-3065) (n=9)	2750 (1510-3402) (n=10)	0.76
Age at diagnosis (weeks)	6 (2-17) (n=11)	11 (1-47) (n=10)	0.48
Age at transfer to SU (years)	5.8 (1.9-12.0) (n=11)	10.3 (0.5-26.0) (n=10)	0.12
Current age (years)	19 (15-23) (n=11)	22.5 (11-36) (n=10)	0.08
Pre-SU HbA1c (%)	6.7 (5.3-7.9) (n=11)	8.1 (5.8-9.5) (n=10)	0.001
Pre-SU HbA1c (mmol/mol)	50 (34-63) (n=11)	65 (40-80) (n=10)	0.001
Year 1 HbA1c (%)	6.0 (5.0-6.9) (n=10)	5.7 (5.0-7.3) (n=8)	0.92
Year 1 HbA1c (mmol/mol)	42 (31-52) (n=10)	39 (31-56) (n=8)	0.92
Most recent HbA1c (%)	7.0 (5.3-10.3) (n=11)	6.3 (5.3-10.1) (n=10)	0.32
Most recent HbA1c (mmol/mol)	53 (34-89) (n=11)	45 (34-87) (n=10)	0.32
Year 1 SU dose (mg/kg/day glyburide)	0.09 (0.01-1.25) (n=10)	0.50 (0.17-1.11) (n=8)	0.08
Most recent SU dose (mg/kg/day glyburide)	0.36 (0.02-1.45) (n=10)	0.35 (0.10-1.30) (n=10)	0.67
N on insulin at recent follow-up	3/11 (27)	0/10 (0)	0.08
N with neurological features any at most recent visit (%)	7/11 (64)	6/10 (60)	0.85

Table 2. Clinical features of individuals with dominantly vs recessively inherited variants in the *ABCC8* gene. Data are presented as median (range). N is different for each variable due to differences in amount of available data. SU = sulfonylurea, N = number.









CNS feature	Pre SU transfer (n)	Post SU transfer (n)	Post transfer only (n)	Improvement on SU (n)
Any	13	13	0	7
DD	10	10	1	3
LD	7	11	2	1
ADHD	6	8	2	1
Epilepsy	4	3	2	2**
Muscle weakness	2	2	0	0
Anxiety	2	2	0	0
Sleep problems	2	2	1	0
Spasticity	2	2	0	0
Autism	0	0	0	N/A
Other*	5	4	0	1***

Supplementary Table S1. Neurological features present before and after transfer to sulfonylureas, and features that improved on SU transfer

*'Other' CNS features prior to sulfonylurea transfer in addition to the specific features listed consisted of obsessive-compulsive disorder (OCD) with mild Tourette's, encopresis, hypertonia, hypotonia and an abnormal electroencephalogram (EEG) (in the absence of a diagnosis of epilepsy). These features were also present after sulfonylurea transfer with the exception of hypotonia (but not known if this was tested).

**both individuals had seizures at time of diagnosis only which may have been attributable to cerebral oedema (Table S2)

individuals treated with anti-epileptic medication not included as 'improved' (Table S2)

***improved background on EEG

CNS = central nervous system, SU = sulfonylurea, DD = developmental delay, LD = learning difficulties, ADHD = attention deficit hyperactivity disorder

Mutation in	Age at	Age at	Clinical history	Other neurological	Neurological features improved on
ABCC8 gene	diagnosis	transfer		features present in	transfer to SU
	of diabetes	to SU		addition to seizures /	
	(weeks)	(years)		epilepsy	
Individuals in wh	hom metabolic d	listurbance of	at diagnosis may have contributed to seizures		1
P45L/G1401R 6	6	8	Diabetic ketoacidosis at 6 weeks of age with	Muscle weakness,	Improvements in sleep, speech,
			severe dehydration, reduced consciousness,	hypertonia, spasticity,	concentration and schoolwork noted by
			opisthotonus and partial seizures - diagnosed with	DD, LD, and sleep	parents and teachers. No epilepsy at most
			cerebral edema (31)	problems	recent follow-up.
V215A/V215A 9	9	0.5	Focal seizures around time of diagnosis: in left	DD (mild), LD (mild)	No seizures since 2 months of age but on
			arm 2 days before admission and in left arm and		antiepileptic drugs. Other features (DD /
			leg 2 days after admission		LD) identified only after SU transfer.
V86G/N 5	5	3	Seizures only at time of diagnosis: none since.	DD, LD, ADHD,	Slight improvement. No anxiety post SU
				anxiety	transfer and no further seizures. Main
					problem currently is speech delay /
					difficulties at school.
Individuals in wl	hom seizures ar	e attributabl	e to another (non-metabolic) cause		
L1295F/N 1	12	6	One seizure due to starting treatment with	DD, LD, ADHD,	No change.
			dexmethylphenidate for ADHD (no further	muscle weakness	
			seizures on stopping drug)		
L135P/N	6	10	Viral meningoencephalitis at 6 weeks of age,	Spastic paraplegia, DD,	No change. Epilepsy not reported at recent
			treated with Depakine 200-300mg at clinic	LD, sleep problems	follow-up but has had treatment with
			follow-up prior to SU transfer		antiepileptic drugs

Supplementary Table S2. Clinical details of patients in whom seizures / epilepsy may have been attributable to factors other than the genetic

mutation. SU = sulfonylurea, DD = developmental delay, LD = learning difficulties, ADHD = attention deficit hyperactivity disorder

Outcome on SU treatment	ABCC8-PNDM (n=21)	<i>KCNJ11-</i> PNDM (n=81)
Patients independent of insulin at 10 years (%)	86	93
Median HbA1c at 10 years - paired (%)	6.5 (n=16)	6.4 (n=64)
Median SU dose required at 10 years - paired (mg/kg/day glyburide)	0.25 (n=16)	0.23 (n=64)
Median BMI at 10 years SDS (kg/m ²)	-0.75 (n=13)	-0.22 (n=72)
Frequency of neurological features at 10 years (%)	62 (n=21)	64 (n=81)
Improvement in neurological features after SU transfer (%)	54 (n=13)	47 (n=38)
Number of episodes of severe hypoglycemia on SU only over 10 years	0 (n=18)	0 (n=75)
Frequency of side effects (%)	11 (n=18)	14 (n=81)
Frequency of diabetes complications (%)	11 (n=18)	9 (n=81)

Supplementary Table S3. Comparison of long-term outcomes in individuals with mutations in the KCNJ11 and ABCC8 genes.

Comparative data on *KCNJ11*-PNDM taken from Bowman et al Lancet D&E 2018 (11) SU = sulfonylurea, PNDM = permanent neonatal diabetes mellitus, BMI = body mass index, SDS = standard deviation score