

Maturity-onset Diabetes of the Young: Update on Diagnosis and Treatment

Kyung Mi Jang

Department of Pediatrics, Yeungnam University College of Medicine, Daegu, Korea

Maturity-onset diabetes of the young (MODY) is characterized by a heterogeneous group of monogenic diabetes. MODY has autosomal dominant inheritance, a primary defect in pancreatic β -cell, and an early onset. Discriminating MODY from type 1 or type 2 diabetes is often challenging at first. To date, 14 different disease causing mutations have been identified in MODY patients worldwide. Targeted DNA sequencing is the gold standard to diagnose MODY and their asymptomatic relatives. Next-generation sequencing may help successfully to diagnose MODY patients and identify new MODY genes. In this review, the current perspectives on diagnosis and treatment of MODY and discrepancy in the disease-causing mutations between the Asian and Caucasian patients with MODY are summarized.

Key words: Diabetes mellitus, Genes, Sequencing, MODY

REVIEW ARTICLE

Received: September 18, 2020 Revised: October 12 2020 Accepted: October 21, 2020

Correspondence to: Kyung Mi Jang Department of Pediatrics, Yeungnam University College of Medicine, 170 Hyeonchung-ro, Nam-gu, Daegu 42415, Republic of Korea Tel: +82-53-620-3532 Fax: +82-53-629-2252 E-mail: fortune001j@gmail.com

ORCID Kyung Mi Jang: https://orcid.org/0000-0002-2226-9268

Copyright © 2021, Interdisciplinary Society of Genetic & Genomic Medicine

This is an Open Access article distributed under the terms of the Creative Commons Attribution Non-Commercial License (https://creativecommons.org/licenses/by-nc/4.0) which permits unrestricted non-commercial use, distribution, and reproduction in any medium, provided the original work is properly cited.

INTRODUCTION

Mutations in the genes that control insulin secretion cause maturity-onset diabetes of the young (MODY) [1,2]. MODY was first reported in 1974 as a mild familial diabetes with dominant inheritance [1]. MODY is characterized by β -cell dysfunction without autoantibodies, autosomal dominant inheritance and onset before 45 years of age [3]. MODY is not easily distinguished from type 1 diabetes and type 2 diabetes at diagnosis. Type 1 diabetes differs from MODY, in terms of pathogenesis of pancreatic β -cell auto-immunity. MODY patients have maintained β -cell function. Type 2 diabetes resembles MODY patients, but, type 2 diabetes patients are generally obese, and have insulin resistance.

To date, at least 14 MODY subtypes are reported (Table 1) [4-11]. MODY is rare disease, comprising between 1-5% of all diabetes cases in Europe [12,13]. Mutations in glucokinase (GCK), hepatocyte nuclear factor 1 α (HNF1A), HNF4A, and HNF1B are the most commonly identified causes in more than 95% cases of MODY, respectively, 32%,52%,10%, and 6% of diabetes patients in the United Kingdom [14]. However, there is a big discrepancy among Asian and Caucasian populations. Clinical MODY patients have been known with MODY-related gene defects (GCK, 22.8%; HNF1A, 13.9%; HNF4A, 3.8%, and HNF1B, 7.6%) in Japan, whereas the causative mutations were not identified in 51.9% [15]. This suggests that there are unidentified MODY related genes in Asia. Moreover, the exact prevalence of MODY in the Asian population has not been reported. This review summarizes the current perspective on understanding of MODY and discusses the Asian MODY study.

2 Journal of Interdisciplinary Genomics

Subtype	MODY gene	Chromosome location	Gene function	Pathophysiology	Other features	Treatment
MODY 1	HNF4a	20q13	Transcription factor	β -cell dysfunction	Hyperinsulinism during infancy, Low triglycerides level	Sulfonylureas
MODY 2	GCK	7q13	Enzyme in the first step of glucose metabolism	β -cell dysfunction	Mild fasting hyperglycemia	No medications, Diet
MODY 3	HNF1a	12q24	Transcription factor	β-cell dysfunction	Glycosuria	Sulfonylureas
MODY 4	PDX1	13q12	Transcription factor	β-cell dysfunction	Pancreatic agenesis in homozygote/compound heterozygote	Diet or OAD or insulin
MODY 5	HNF1β	17q12	Transcription factor	β-cell dysfunction	Renal anomalies, genital anomalies, pancreatic hypoplasia	Insulin
MODY 6	NEUROD1	2q31	Transcription factor	β -cell dysfunction	Neonatal diabetes, Neurological abnormalities in homozygote	OAD or insulin
MODY 7	KLF11	2q25	Transcription factor	β-cell dysfunction	Similar with type 2 diabetes	OAD or insulin
MODY 8	CEL	9q34	Controls exocrine and endocrine functions of pancreas	Pancreas endocrine and exocrine dysfunction	Exocrine dysfunction, lipomatosis	OAD or insulin
MODY 9	PAX4	7q32	Transcription factor	β-cell dysfunction	Possible ketoacidosis	Diet or OAD, or insulin
MODY 10	INS	11p15	Encode the proinsulin precursor	Insulin gene mutation	PNDM	Diet or OAD, or insulin
MODY 11	BLK	8p23	Tyrosine kinase functions in signal transduction	Insulin secretion defect	Overweight	Diet or OAD, or insulin
MODY 12	ABCC8	11p15	Regulating insulin release	ATP-sensitive potassium channel dysfunction	PNDM, TNDM	Sulfonylurea
MODY 13	KCNJ11	11p15	Regulating insulin release	ATP-sensitive potassium channel dysfunction	Neonatal diabetes in homozygote	OAD or insulin
MODY 14	APPL1	3p14	Insulin signal pathway	Insulin secretion defect	Dysmorphic phenotype, developmental delay	Diet or OAD, or insulin

Table 1. Causative genes and their clinical characteristics of MODY subtype

MODY, maturity-onset diabetes of young; HNF4α, hepatocyte nuclear factor 4α; GCK, glucokinase; PDX1, pancreatic and duodenal homeobox 1; HNF1β, hepatocyte nuclear factor1β; NEUROD1 neurogenic differentiation 1; KLF11, kruppel-like factor 11; CEL, carboxyl ester lipase; PAX4, pairedbox-containing gene 4; INS, insulin; BLK, B-lymphocyte kinase; ABCC8, ATP-binding cassette, subfamily C(CFTR/MRP), member 8; KCNJ11, potassium channel inwardly rectifying subfamily J, member 11, APPL1, adaptor protein, phosphotyrosine interacting with PH domain and leucine zipper 1; OAD, oral antidiabetic agent; PNDM, permanent neonatal diabetes; TNDM, transient neonatal diabetes.

MODY SUBTYPES AND THEIR CLINICAL FEATURES

HNF1A-MODY (MODY 3)

Hepatocyte nuclear factor 1 α is a homeodomain-containing transcription factor expressed in liver, kidney, pancreatic β -cell, and intestine. More than 400 HNF1A mutations have been reported in 1,247 families [16]. A mutation in exon 4 of the gene (P291fsinsC) is most frequently reported [16,17]. A missense mutation of HNF1A (R263L) has been reported in a Korean MODY, causing defective insulin secretion [18]. Mutations in HNF1A have high penetrance, and approximate 63% carriers develop diabetes by the age of 25, and almost all carriers develop diabetes by the age 55 [19]. Because HNF1A is expressed in various organs, patients with HNF1A MODY develop glycosuria even diabetes, due to a low renal threshold of glucose sensing [20]. Heterozygous mutation in HNF1A gene

cause β -cell dysfunction, and patients with this mutation have the similar risk of long-term complications over the time to those in type 1 and type 2 diabetes [21]. Therefore, tight glucose control and close monitoring for these patients are need. Patients with HNF-1MODY have high sensitivity to the sulfonylurea treatment, which is a five-greater response than metformin and is considered for first line treatment [22]. Many studies have been reported that many patients with HNF-1 MODY might switch from insulin to a sulfonylurea [23].

GCK-MODY (MODY 2)

GCK is an enzyme which catalyzes the conversion of glucose to glucose-6-phosphate in glucose metabolism, and controls glucose-related insulin secretion. Heterozygous mutations in GCK develop GCK-MODY, known as MODY-2. More than 600 mutations in GCK have been identified [24]. The heterozygous mutation in GCK cause elevated glucose threshold for insulin release, developing mild fasting hyperglycemia [25]. According to Japan study, MODY 2 is most common form in MODY subtype, comprising approximately 48% of patients with MODY [15]. This result is similar to that in European MODY patients [26]. Therefore, most of MODY 2 is diagnosed through routine examination, as urine glucose screening test. Hence, MODY 2 has high prevalence in countries where routine urinary screening test is performed [27,28]. However, only a small number (<5%) about MODY 2 was reported in Korea and China [29,30]. Therefore, this suggests that there is a big discrepancy between the Asian and Caucasian populations. The clinical manifestations comprise mild fasting hyperglycemia (5.5-8.0 mmol/L, glycosylated hemoglobin range of 5.6-7.3%) [25] which from birth, demonstrates slightly aggravations with getting old. Therefore, MODY 2 patients usually do not need treatment, outside pregnancy, because their diseases are non-progressive, and they rarely develop long-term complications [31].

HNF4A-MODY (MODY1)

HNF4A is also a transcription factor which is expressed in diverse organ such as the intestine, liver, pancreatic β -cell, and kidney. This regulates genes related to glucose transport and metabolism [32]. HNF4A-MODY is rare, comprising only approximately 5% cases of all MODY, more than 103 mutations have been identified [16,33]. The clinical manifestation of HNF4A is similar to that in HNF1A. Heterozygous HNF4A are usually not diagnosed before adolescence. Unlike HNF1A-MODY patients, MODY1 patients don't present glycosuria, low level of apolipoproteins might be a clue for diagnosis [34]. HNF4A-MODY has similar response to sulfonylurea, therefore, sulfonylurea should be considered as the first-line treatment [22].

PDX1-MODY (MODY 4)

PDX1 is known as insulin promoter factor 1 (IPF1) which is a homeodomain-containing transcription factor, regulating development of β -cell and insulin release in pancreas [35]. MODY 4 is very rare, and was first reported in 1997. Heterozygous IPF1 gene mutation causes β -cell dysfunction and homozygous mutation develop permanent neonatal diabetes due to pancreas agenesis [36].

HNF1β-MODY (MODY 5)

The transcription factor $HNF1\beta$ is associated with the organogenesis of the pancreas, liver, kidney, lung, gut and genitourinary tract [37]. Patients with mutation in HNF1 β have kidney disease such as renal cysts, renal tract malformations, and familial hypoplastic glomerulocystic kidney disease [38, 39]. It is called RCAD (renal cysts and diabetes) syndrome. Renal dysfunction is usually developed in affected patients by 45 years of age, and approximately half of these patients will progress to end-stage renal disease without diabetic renal disease [40]. Until now, more than 65 mutations have been identified. Approximate half of mutations comprise exon or complete gene deletions [41]. A heterozygous P159L mutation in HNF1B was reported in a Korean family and this pathogenesis was discovered [42,43]. Half of carriers develop diabetes in early adulthood. In contrast to patient with MODY 3, MODY 5 patients progress to an insulin-dependent condition due to pancreatic hypoplasia. Patients harboring same HNF1^β mutations show highly variable manifestations, which might exhibit different phenotype between family members.

NEUROD1-MODY (MODY 6)

NEUROD1 is a basic-loop-helix transcription factor, which is associated with pancreatic and neuronal development. Heterozygous mutations lead to diabetes in childhood or early adult. Whereas, homozygous mutations develop neonatal diabetes, neurological abnormalities [44,45].

WAY TO CORRECTLY DIAGNOSE PATIENTS WITH MODY

MODY shows similar clinical manifestations with both common types of diabetes. Therefore, discriminating MODY from type 1 diabetes or type 2 diabetes is often challenging at first visit [12,46,47]. The diagnostic criteria comprise 1) presence of overt diabetes at least three consecutive generations with autosomal dominant pattern, 2) diagnosis with diabetes before the age of 25, 3) absence of β -cell autoantibodies, 4) relatively preserved endogenous insulin secretion with a serum C-peptide level of > 200 pmol/L [12,46,47].

MODY can be detected by direct sanger DNA sequencing with up to 100% sensitivity [14,43]. Using targeted or wholeexome gene sequencing for MODY is gold standard to identify disease-causing mutations in MODY (Fig. 1). However, genetic testing for patients with MODY is not cheap, and may only be available in specialized laboratories. Molecular genetic testing is needed in targeted selection of individuals. Various algorithms have been developed to choose individual candidates who should get genetic testing for MODY [48,49]. Shield et al.

4 Journal of Interdisciplinary Genomics

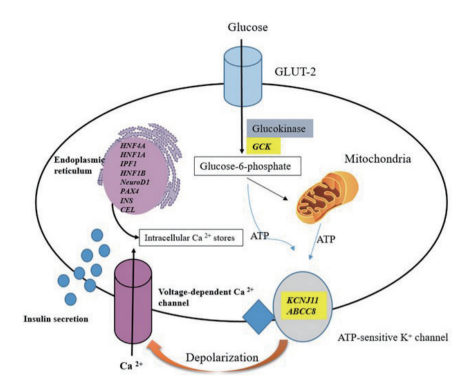


Fig. 1. Schematic representation of MODY-causing genes in pancreatic β cells and mechanism of glucose induced insulin secretion. HNF4A, hepatocyte nuclear factor 4a; HNF1A, hepatocyte nuclear factor 1a; IPF1, insulin promoter factor-1; HNF1B, hepatocyte nuclear factor1β; NeuroD1, neurogenic differentiation 1; PAX4, paired box-containing gene 4; INS, insulin gene; CEL, carboxyl ester lipase; GCK, glucokinase; KCNJ11, potassium channel inwardly rectifying subfamily J, member 11; ABCC8, ATP-binding cassette, subfamily C (CFTR/MRP), member 8.

[48] have developed a clinical model to discriminate MODY from type 1 and type 2 diabetes. According to this report, MODY has lower HbA1_C than type 1 diabetes and 23 times more family history of diabetes than type 1 diabetes. MODY has lower HbA1_C level, a lower body mass index (BMI), and younger age at diagnosis.

The genetic causes of MODY have been widely reported. Although MODY has been identified in Caucasian populations, the exact prevalence of MODY is not reported in Korea. According to recent Korea study, clinically suspected MODY patients underwent targeted MODY sequencing [50]. The diagnostic yield was similar to those in large study conducted in the United Kingdom (27%) [51]. In Korea study, 40 subjects as suspected MODY showed only 5% MODY3 and 2.5% MODY 2 [52]. This result is similar to those in China and Japan [30,53]. This reports suggest that East Asia might have high possibilities of a not yet identified 'MODY X' [30,50,53-56]. Therefore, there are attempts to identify new MODY gene by whole-exome sequencing [9]. Next generation sequencing is good tools to find unidentified genetic defects [56,57]. Furthermore, there are many attempts to discover new causative gene variants in MODY patients by whole exome sequencing in Korea [9].

CONCLUSION

MODY is a common cause of monogenic diabetes, comprising 1-2% of all diabetes causes. Despite of low incidence, identification of MODY gene has importance because correct diagnosis can help MODY patients receive individualized treatment.

Direct sequencing is best way to diagnose MODY patients, however, there are unidentified MODY gene. Therefore, efforts to identify new MODY gene and systemic approaches for MODY patients are needed for rapid diagnosis and proper treatment.

CONFLICTS OF INTEREST

No potential conflict of interest relevant to this article was reported.

REFERENCES

- 1. Tattersall RB. Mild Familial Diabetes with Dominant Inheritance. *QJM*: An International Journal of Medicine 1974;43:339-57.
- Yang Y, Chan L. Monogenic Diabetes: what it teaches us on the common forms of type 1 and type 2 diabetes. Endocr Rev 2016; 37:190-222.
- Ellard S, Bellanné-Chantelot C, Hattersley AT. European Molecular Genetics Quality Network Mg: Best practice guidelines for the molecular genetic diagnosis of maturity-onset diabetes of the young. Diabetologia 2008;51:546-53.
- 4. Froguel P, Vaxillaire M, Sun F, Velho G, Zouali H, Butel MO, et al. Close linkage of glucokinase locus on chromosome 7p to early-onset non-insulin-dependent diabetes mellitus. Nature 1992;356:162-4.
- 5. Yamagata K, Furuta H, Oda N, Kaisaki PJ, Menzel S, Cox NJ, et al. Mutations in the hepatocyte nuclear factor-4α gene in maturity-onset diabetes of the young (MODY1). Nature 1996;384: 458-60.
- 6. Horikawa Y, Iwasaki N, Hara M, Furuta H, Hinokio Y, Cockburn BN, et al. Mutation in hepatocyte nuclear factor-1 beta gene (TCF2) associated with MODY. Nat Genet 1997;17:384-5.
- Henzen C. Monogenic diabetes mellitus due to defects in insulin secretion. Swiss medical weekly 2012;142:w13690.
- Prudente S, Jungtrakoon P, Marucci A, Ludovico O, Buranasupkajorn P, Mazza T, et al. Loss-of-function mutations in APPL1 in familial diabetes mellitus. American Journal of Human Genetics 2015;97:177-85.
- 9. Shim YJ, Kim JE, Hwang SK, Choi BS, Choi BH, Cho EM, et al. Identification of candidate gene variants in Korean MODY families by whole-exome sequencing. Hormone Research in Paediatrics 2015;83:242-51.
- Shepherd M, Shields B, Hammersley S, Hudson M, McDonald TJ, Colclough K, et al. Systematic population screening, using biomarkers and genetic testing, identifies 2.5% of the U.K. Pediatric Diabetes Population With Monogenic Diabetes. Diabetes Care 2016;39:1879-88.
- 11. Urakami T. Maturity-onset diabetes of the young (MODY): current perspectives on diagnosis and treatment. Diabetes Metab Syndr Obes 2019;12:1047-56.
- 12. Thanabalasingham G, Pal A, Selwood MP, Dudley C, Fisher K, Bingley PJ, et al. Systematic assessment of etiology in adults with a clinical diagnosis of young-onset type 2 diabetes is a successful strategy for identifying maturity-onset diabetes of the young. Diabetes Care 2012;35:1206-12.
- Kim SH. Maturity-onset diabetes of the young: what do clinicians need to know? Diabetes & Metabolism Journal 2015;39: 468-77.
- 14. Kavvoura FK, Owen KR. Maturity onset diabetes of the young: clinical characteristics, diagnosis and management. Pediatr Endocrinol Rev 2012;10:234-42.
- 15. Yorifuji T, Fujimaru R, Hosokawa Y, Tamagawa N, Shiozaki M, Aizu K, et al. Comprehensive molecular analysis of Japanese patients with pediatric-onset MODY-type diabetes mellitus. Pediatr

Diabetes 2012;13:26-32.

- 16. Colclough K, Bellanne-Chantelot C, Saint-Martin C, Flanagan SE, Ellard S. Mutations in the genes encoding the transcription factors hepatocyte nuclear factor 1 alpha and 4 alpha in maturity-onset diabetes of the young and hyperinsulinemic hypoglycemia. Human Mutation 2013;34:669-85.
- 17. Ellard S, Colclough K. Mutations in the genes encoding the transcription factors hepatocyte nuclear factor 1 alpha (HNF1A) and 4 alpha (HNF4A) in maturity-onset diabetes of the young. Human Mutation 2006;27:854-69.
- Kim KA, Kang K, Chi YI, Chang I, Lee MK, Kim KW, et al. Identification and functional characterization of a novel mutation of hepatocyte nuclear factor-1alpha gene in a Korean family with MODY3. Diabetologia 2003;46:721-7.
- 19. Shepherd M, Ellis I, Ahmad AM, Todd PJ, Bowen-Jones D, Mannion G, et al. Predictive genetic testing in maturity-onset diabetes of the young (MODY). Diabet Med 2001;18:417-21.
- 20. Pontoglio M, Prie D, Cheret C, Doyen A, Leroy C, Froguel P, et al. HNF1alpha controls renal glucose reabsorption in mouse and man. EMBO reports 2000;1:359-65.
- 21. Steele AM, Shields BM, Shepherd M, Ellard S, Hattersley AT, Pearson ER. Increased all-cause and cardiovascular mortality in monogenic diabetes as a result of mutations in the HNF1A gene. Diabet Med 2010;27:157-61.
- 22. Pearson ER, Pruhova S, Tack CJ, Johansen A, Castleden HAJ, Lumb PJ, et al. Molecular genetics and phenotypic characteristics of MODY caused by hepatocyte nuclear factor 4α mutations in a large European collection. Diabetologia 200;48:878-85.
- 23. Pearson ER, Starkey BJ, Powell RJ, Gribble FM, Clark PM, Hattersley AT. Genetic cause of hyperglycaemia and response to treatment in diabetes. Lancet 2003;362:1275-81.
- 24. Osbak KK, Colclough K, Saint-Martin C, Beer NL, Bellanne-Chantelot C, Ellard S, et al. Update on mutations in glucokinase (GCK), which cause maturity-onset diabetes of the young, permanent neonatal diabetes, and hyperinsulinemic hypoglycemia. Human Mutation 2009;30:1512-26.
- 25. Steele AM, Wensley KJ, Ellard S, Murphy R, Shepherd M, Colclough K, et al. Use of HbA1c in the Identification of Patients with Hyperglycaemia Caused by a Glucokinase Mutation: Observational Case Control Studies. PLoS ONE 2013;8:e65326.
- 26. Feigerlová E, Pruhová S, Dittertová L, Lebl J, Pinterová D, Kolostová K, et al. Aetiological heterogeneity of asymptomatic hyperglycaemia in children and adolescents. Eur J Pediatr 2006; 165:446-52.
- 27. Estalella I, Rica I, Perez de Nanclares G, Bilbao JR, Vazquez JA, San Pedro JI, et al. Mutations in GCK and HNF-1alpha explain the majority of cases with clinical diagnosis of MODY in Spain. Clin Endocrinol (Oxf) 2007;67:538-46.
- 28. Codner E, Rocha A, Deng L, Martínez-Aguayo A, Godoy C, Mericq V, et al. Mild fasting hyperglycemia in children: high rate of glucokinase mutations and some risk of developing type 1 diabetes mellitus. Pediatr Diabetes 2009;10:382-8.
- 29. Hwang JS. MODY Syndrome. J Korean Soc Pediatr Endocrinol 2010;15:1-6.

6 Journal of Interdisciplinary Genomics

- 30. Xu JY, Dan QH, Chan V, Wat NM, Tam S, Tiu SC, et al. Genetic and clinical characteristics of maturity-onset diabetes of the young in Chinese patients. European journal of human genetics: EJHG 2005;13:422-7.
- 31. Hattersley A, Bruining J, Shield J, Njolstad P, Donaghue KC. The diagnosis and management of monogenic diabetes in children and adolescents. Pediatr Diabetes 2009;10(Suppl 12):33-42.
- 32. Stoffel M, Duncan SA. The maturity-onset diabetes of the young (MODY1) transcription factor HNF4alpha regulates expression of genes required for glucose transport and metabolism. Proceedings of the National Academy of Sciences of the United States of America 1997;94:13209-14.
- 33. Frayling TM, Evans JC, Bulman MP, Pearson E, Allen L, Owen K, et al. beta-cell genes and diabetes: molecular and clinical characterization of mutations in transcription factors. Diabetes 2001;50 (Suppl 1):S94-100.
- 34. Lehto M, Bitzén PO, Isomaa B, Wipemo C, Wessman Y, Forsblom C, et al. Mutation in the HNF-4alpha gene affects insulin secretion and triglyceride metabolism. Diabetes 1999;48:423.
- 35. Stoffers DA, Zinkin NT, Stanojevic V, Clarke WL, Habener JF. Pancreatic agenesis attributable to a single nucleotide deletion in the human IPF1 gene coding sequence. Nat Genet 1997;15: 106-10.
- 36. Schwitzgebel VM, Mamin A, Brun T, Ritz-Laser B, Zaiko M, Maret A, et al. Agenesis of human pancreas due to decreased half-life of insulin promoter factor 1. J Clin Endocrinol Metab 2003;88:4398-406.
- Barbacci E, Reber M, Ott MO, Breillat C, Huetz F, Cereghini S. Variant hepatocyte nuclear factor 1 is required for visceral endoderm specification. Development (Cambridge, England) 1999; 126:4795-805.
- 38. Bingham C, Ellard S, Allen L, Bulman M, Shepherd M, Frayling T, et al. Abnormal nephron development associated with a frameshift mutation in the transcription factor hepatocyte nuclear factor-1 beta. Kidney Int 2000;57:898-907.
- Edghill EL, Oram RA, Owens M, Stals KL, Harries LW, Hattersley AT, et al. Hepatocyte nuclear factor-1beta gene deletions--a common cause of renal disease. Nephrol Dial Transplant 2008;23: 627-35.
- 40. Bingham C, Bulman MP, Ellard S, Allen LI, Lipkin GW, Hoff WG, et al. Mutations in the hepatocyte nuclear factor-1beta gene are associated with familial hypoplastic glomerulocystic kidney disease. American Journal of Human Genetics 2001;68:219-24.
- 41. Chen YZ, Gao Q, Zhao XZ, Chen YZ, Bennett CL, Xiong XS, et al. Systematic review of TCF2 anomalies in renal cysts and diabetes syndrome/maturity onset diabetes of the young type 5. Chinese Medical Journal 2010;123:3326-33.
- 42. Kim EK, Lee JS, Cheong HI, Chung SS, Kwak SH, Park KS. Identification and functional characterization of P159L mutation in HNF1B in a family with maturity-onset diabetes of the young 5 (MODY5). Genomics & Informatics 2014;12:240-6.
- 43. Ellard S, Lango Allen H, De Franco E, Flanagan SE, Hysenaj G, Colclough K, et al. Improved genetic testing for monogenic diabetes using targeted next-generation sequencing. Diabetologia 2013;

56:1958-63.

- 44. Malecki MT, Jhala US, Antonellis A, Fields L, Doria A, Orban T, et al. Mutations in NEUROD1 are associated with the development of type 2 diabetes mellitus. Nat Genet 1999;23:323-8.
- 45. Gonsorcíková L, Průhová S, Cinek O, Ek J, Pelikánová T, Jørgensen T, et al. Autosomal inheritance of diabetes in two families characterized by obesity and a novel H241Q mutation in NEUROD1. Pediatr Diabetes 2008;9(4 Pt 2):367-72.
- Vaxillaire M, Froguel P. Monogenic diabetes in the young, pharmacogenetics and relevance to multifactorial forms of type 2 diabetes. Endocr Rev 2008;29:254-64.
- 47. Hattersley AT, Greeley SAW, Polak M, Rubio-Cabezas O, Njolstad PR, Mlynarski W, et al. ISPAD Clinical Practice Consensus Guidelines 2018: The diagnosis and management of monogenic diabetes in children and adolescents. Pediatr Diabetes 2018;19 Suppl 27:47-63.
- 48. Shields BM, McDonald TJ, Ellard S, Campbell MJ, Hyde C, Hattersley AT. The development and validation of a clinical prediction model to determine the probability of MODY in patients with young-onset diabetes. Diabetologia 2012;55:1265-72.
- 49. Carroll RW, Murphy R. Monogenic diabetes: a diagnostic algorithm for clinicians. Genes (Basel) 2013;4:522-35.
- Park SS, Jang SS, Ahn CH, Kim JH, Jung HS, Cho YM, et al. Identifying Pathogenic Variants of Monogenic Diabetes Using Targeted Panel Sequencing in an East Asian Population. J Clin Endocrinol Metab 2019.
- 51. Shields BM, Hicks S, Shepherd MH, Colclough K, Hattersley AT, Ellard S. Maturity-onset diabetes of the young (MODY): how many cases are we missing? Diabetologia 2010;53:2504-8.
- Hwang JS, Shin CH, Yang SW, Jung SY, Huh N. Genetic and clinical characteristics of Korean maturity-onset diabetes of the young (MODY) patients. Diabetes Res Clin Pract 2006;74:75-81.
- 53. Nishigori H, Yamada S, Kohama T, Utsugi T, Shimizu H, Takeuchi T, et al. Mutations in the hepatocyte nuclear factor-1 alpha gene (MODY3) are not a major cause of early-onset non-insulindependent (type 2) diabetes mellitus in Japanese. J Hum Genet 1998;43:107-10.
- 54. Iwasaki N, Oda N, Ogata M, Hara M, Hinokio Y, Oda Y, et al. Mutations in the hepatocyte nuclear factor-1alpha/MODY3 gene in Japanese subjects with early- and late-onset NIDDM. Diabetes 1997;46:1504-8.
- 55. Yamagata K, Nammo T, Moriwaki M, Ihara A, Iizuka K, Yang Q, et al. Overexpression of dominant-negative mutant hepatocyte nuclear fctor-1 alpha in pancreatic beta-cells causes abnormal islet architecture with decreased expression of E-cadherin, reduced beta-cell proliferation, and diabetes. Diabetes 2002;51:114-23.
- 56. Tanaka D, Nagashima K, Sasaki M, Funakoshi S, Kondo Y, Yasuda K, et al. Exome sequencing identifies a new candidate mutation for susceptibility to diabetes in a family with highly aggregated type 2 diabetes. Molecular Genetics and Metabolism 2013;109: 112-7.
- 57. Johansson S, Irgens H, Chudasama KK, Molnes J, Aerts J, Roque FS, et al. Exome sequencing and genetic testing for MODY. PLoS One 2012;7:e38050.