Project Details		
Project Code	MRCPHS24Ex Dempster	
Title	Investigating the epigenetic component of insulin secretion disorders	
Research Theme	Population Health Sciences	
Summary	During this PhD, you will use innovative bioinformatic techniques to	
	investigate how epigenetic processes contribute to rare genetic	
	disorders that affect insulin secretion. Critically, you will use these	
	disease-specific epigenetic patterns to develop classifiers termed	
	"episignatures" to assist with the interpretation of novel genetic variants	
	and as a diagnosis tool.	
Description	Mendelian disorders, while individually rare, are collectively common	
	affecting around 3.5% of the population. New sequencing technologies	
	have revolutionised medical genetics and enabled the identification of	
	rare mutations as the cause of many different disorders. However,	
	determining the exact genetic cause can still be challenging when	
	dealing with variants with unknown or uncertain clinical significance.	
	Recently there has been increasing evidence that epigenetic	
	modifications can assist in the interpretation of genetic variation.	
	Epigenetic modifications are biochemical modifications to DNA or DNA/protein complexes, which control gene expression independently	
	of DNA sequence variation. Epigenetic processes such as DNA	
	methylation are highly dynamic during development. Alterations in a	
	developmental trajectory due to a change in the underlying genetic	
	sequencing can therefore have profound effects on the epigenome.	
	Thus, an altered DNA methylation signature can be propagated and	
	maintained across multiple cell lineages resulting in a stable epigenetic	
	signature of a monogenic disease. The clinical laboratories in Exeter are	
	a national and international referral centre for genetic testing for insulin	
	secretion disorders; the Beta-cell Research Bank at Exeter provides a	
	unique resource of >25,000 patient samples with DNA sequencing data	
	available for thousands of these samples. Patients with congenital hyper-	
	and hypoglycaemia represent the extreme ends of the spectrum in	
	terms of insulin secretion. Studying the underlying genetic aetiology of	
	disease in these individuals provides a unique opportunity to improve	
	understanding of pancreas development and function. Great progress	
	has been made in understanding the genetic basis of disease in patients	
	with neonatal diabetes with 23 disease-causing genes reported.	
	Screening of these genes identifies a pathogenic mutation in >82% of	
	individuals. For patients with hyperinsulinaemic hypoglycaemia the	
	underlying genetic aetiology is only found in 50% of cases. Current	
	efforts have focused on using genome sequencing to search for the	
	'missing' mutations in insulin secretion disorders. The success of this	
	approach is however reliant on 'missing' causative mutations being	
	present in the DNA sequence, however epigenetic changes cannot be	
	detected by this approach. The importance of epigenetics in controlling	
	insulin secretion is well-established. DNA methylation defects at the	
	imprinted loci on chromosome 6q24 are reported in 70% of patients	
	with transient neonatal diabetes. Characterising the role of DNA	
	methylation in insulin secretion disorders will provide further insight into	
	the regulation of the genetic pathways controlling insulin secretion	
	which will be crucial for informing studies investigating models of beta-	

cell regeneration which are being developed to treat patients with more common forms of insulin dysregulation such as type 1 diabetes. We are currently profiling DNA methylation in the cohort using the Illumina DNA methylation array to create an epigenetic data resource for monogenic insulin secretion disorders. This PhD project aims to bring together expertise in monogenic disease and epigenetics, and the largest international cohort of patients with neonatal diabetes (n= >4000) and hyperinsulinism (n= >5000) to help the student achieve the following aims: 1, To perform a thorough review of the literature regarding epigenetics and insulin secretion disorders and the development and application of episignatures. 2, To use an integrated genomics/methylomic approach to investigate the role of DNA methylation in the aetiology of insulin secretion monogenic disorders. 3, To develop and test different methods to determine the optimal epigenetic classifier method. 4, To develop disease-specific episignatures that will aid variant interpretation and potentially provide patients with a genetic diagnosis that could have treatment and disease management implications.

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