# Centre for Cancer Research and Cell Biology

Ducio et Title				,
Project Title	Investigating an association between the DDRD molecular subgroup and expression of immune checkpoint pathway members in breast cancer			
Supervisor(s)	1. Prof Richard Kennedy			
	2. Dr Nuala McCabe			
School / Centre	Centre for Cancer Researc	ch and	Cell Biology (CCRCB)	
Principal	Email: r.kennedy@qub.ac.	uk	Tel: +44 (0)28 9097 2777	
Supervisor's Contact Details				
Degree Pathway	Medical Science	$\checkmark$		
for which project	Biochemistry			
is suitable (√)	Microbiology			
	General awards		Subject-specific awards	T
Is project of				
suitable standard			British Assoc Dermatologists	
/ subject for studentship	Wolfson Foundation		Digestive Disorders Foundation	
application? $(\checkmark)$	Jean Shanks Foundation		Pathological Society	
			Othor	
Background	Other         Other           We have identified a DNA damage response-deficient (DDRD) molecula			
	of the S-phase specific DNA damage response mechanism, the Fanconi Anaemia (FA)/BRCA pathway (Mulligan et al., 2014). A 44-gene assay identifying this subtype was validated as predicting benefit from DNA- damaging chemotherapy. Importantly, upregulation of interferon-related genes is observed within the DDRD molecular subtype (including the chemokines <i>CXCL10</i> and <i>CCL5</i> , and the immune inhibitory genes <i>PD-L1</i> and <i>IDO1</i> ).			DNA- related
	Recently we have shown that activation of the innate immune STING- mediated pathway is responsible for chemokine production in response to DNA damage <i>in vitro</i> , resulting in an inflammatory microenvironment in DDRD breast tumours (Parkes et al., 2017). Activation of this pathway and associated upregulation of the immune inhibitory gene, <i>PD-L1</i> , may explain the paradoxical lack of T-cell-mediated cytotoxicity observed in DDRD tumours. This provides a rationale for exploration of DDRD in the stratification of patients for immune checkpoint–based therapies.			
	Blockade of immune checkpoints is perhaps the most promising approach to activation of therapeutic anti-tumour immunity. Many of the immune checkpoints are initiated by ligand-receptor interactions at the interface between cancer cells and host immune cells, and as such, they can be readily blocked by antibodies or modulated by recombinant forms of ligands or receptors. Blockade of either PD-1 or its ligands has demonstrated consistent immune-potentiating effects and has been shown to have potent antitumor activity in both murine tumour models and clinical trials including non-small-cell lung cancer, melanoma, and renal- cell cancer. There are currently 6 anti-PD-L1 or PD-1 therapies in clinical trials for cancer (www.clinicaltrials.gov). In addition to the PD-1 pathway,			nmune terface can be rms of ds has been els and l renal- clinical

	there are dozens of other immune-modulating receptor-ligand				
	interactions that may be targeted clinically.				
Aims / objectives	We wish to investigate the expression of additional immune-inhibitory genes (e.g. <i>CTLA4</i> , <i>HVEM</i> , <i>LAG3</i> , <i>OX40</i> etc.) for an association with DDRD biology; this could provide a rationale for further exploration of immune checkpoint-modulating therapies within DDRD positive cancers.				
	To fulfil this aim we will perform real-time quantitative PCR (RQ-PCR) assays to analyse the mRNA expression of selected immune checkpoint gene targets within established isogenic cell line models. The BRCA1-mutant MDA-MB-436 cell line stably expressing either an empty vector control or wild type BRCA1 represents an <i>in vitro</i> model of DDRD-positivity and –negativity, respectively.				
	To investigate the requirement of the STING pathway and its component proteins for the regulation of immune-inhibitory gene expression we will perform siRNA knockdowns of pathway members (e.g. STING, cGAS, IRF3) in combination with DNA damaging chemotherapy and assess the expression of immune inhibitory gene targets.				
Techniques employed:	Routine cell culture, siRNA transfections, drug treatments				
employed.	<ul> <li>RNA extraction, cDNA synthesis, RQ-PCR and gene expression analysis</li> </ul>				

# **Centre for Experimental Medicine**

Project Title	Investigate novel therapies for Cardiovascular disease and Diabetes using patient specific induced pluripotent stem cells.			
Supervisor(s)	1. Dr Andriana Margariti			
	2.			
School / Centre	Medicine / CEM			
Principal Supervisor's Contact Details	Email: Email: a.margariti@qub.ac.uk		Tel: 44 (0)28 9097 6476	
Degree Pathway	Medical Science	V		
for which project	Biochemistry	V	7	
is suitable (√)	Microbiology		7	
	General awards	V	Subject-specific awards	
Is project of suitable standard / subject for studentship application? (√)	Wolfson Foundation Jean Shanks Foundation	V	British Assoc Dermatologists Digestive Disorders Foundation Pathological Society	
Background information:	worldwide and is charact Replacing damaged ECs of availability of appropriate of Recent ability to derive E cells) holds huge therape vascular cell therapy. Recently, a new paradigr been devised. We rease reprogramming, we cou which are induced by th types such as ECs und pluripotency[1]. However, the efficiency underlying mechanisms re- that the four reprogramm precisely activate Novel specific stimuli such as Fibroblasts Growth Fact knowledge has allowed u differentiated cells. Import reprogrammed ECs to en Together, these findings	Jean Shanks Foundation       Pathological Society         Other       Other         Cardiovascular disease and diabetes are the leading causes of death         worldwide and is characterised by endothelial cell (EC) dysfunction.         Replacing damaged ECs could be a potential therapeutic option but the         availability of appropriate cell types has been a major limitation.         Recent ability to derive ECs from induced-pluripotent stem cells (iPS cells) holds huge therapeutic potential for personalised medicine and vascular cell therapy.         Recently, a new paradigm of direct reprogramming strategy has also been devised. We reasoned that, at earlier time points during reprogramming, we could direct the epigenetically activated cells which are induced by the iPS cell factors into lineage specific cell types such as ECs under defined conditions without traversing		

Aims / objectives	This project will elucidate the signalling pathways which are regulated during the reprogramming of somatic cells to endothelial cell lineages using patients specific cells. <b>Experimental Design:</b> Patient specific Induced pluripotent stem cells and short reprogramming cells will be differentiated towards vascular cell lineages. The differentiation stage of the cells will be assessed in time point experiments by extracting RNA and Protein and further analysed, or stained and observed under a confocal microscope. Transfection experiments will be conducted to over-express potential genes which have an essential role in vascular cell differentiation. These studies will screen for and identify potential new therapies for vascular disease and regenerative medicine.
Techniques employed:	The student will have the opportunity to gain expertise in cutting- edge research field of induced pluripotent stem cells technology:
	<ol> <li>Pluripotent Stem cells and the powerful and novel approach of cell reprogramming; maintain them in undifferentiated conditions and induce differentiation towards endothelial cells.</li> <li>Fundamental techniques in molecular biology such as RNA isolation and Real time PCR, protein extraction and western blot, confocal microscope to assess the differentiation stage of the vascular cells.</li> <li>Transfection experiments and luciferase reporter assays to investigate the role of key signalling pathways implicated in vascular cell differentiation. Plasmid Purification.</li> <li>Experimental design, data summary, critical thinking, analysis and interpretation of the results, presentation skills, scientific writing.</li> <li>Overall, the proposed project will be an opportunity for an enthusiastic student who would like to gain experience in the excited filed of stem cells and cell reprogramming and their potential use in regenerative medicine. It will be provided training in pluripotent stem cells laboratory skills, molecular biology, and cell signalling. They will collaborate in a multi-disciplinary research centre. These skills will be an important foundation for whatever career path the student decides to follow in the future.</li> </ol>

#### Intercalated BSc in: **Medical Science Biochemistry** Microbiology **Project Title** Development of survey to explore pharmacological and nonpharmacological airway clearance treatments in bronchiectasis. Supervisor(s) 1.Prof Judy Bradley Judv.bradlev@gub.ac.uk 2. Dr Katherine O'Neill k.oneill@gub.ac.uk School / Centre Clinical Research Facility, Centre for Experimental Medicine, School of Medicine, Dentistry and Biomedical Science Principal Tel: Email: Judy.bradley@qub.ac.uk Supervisor's **Contact Details Degree Pathway** Medical Science Х for which project Biochemistry is suitable (√) Microbiology General awards Subject-specific awards Is project of suitable standard British Assoc Dermatologists / subject for Wolfson Foundation studentship **Digestive Disorders** Foundation application? $(\checkmark)$ Jean Shanks Foundation Pathological Society Other ..... Background Bronchiectasis is a chronic respiratory condition with symptoms including information: difficulty clearing secretions as well as frequent chest infections and declining lung function. A core part of treatment is pharmacological (nebulized therapies and/or oral medications to enhance airway clearance) and non-pharmacological airway clearance treatments (for example a range of physiotherapy chest clearance techniques). Although these treatments are recommended in the guidelines, it is not clear how adherent patients are to these treatments or how frequently they are able to access their healthcare team. We have currently developed a patient questionnaire to explore pharmacological and non-pharmacological airway clearance treatments. This questionnaire is due to be launched on a patient driven European Lung Foundation website. In parallel with this, it is important to understand the prescribing patterns of healthcare team, with regards to pharmacological and non-pharmacological airway clearance treatments. Together, data obtained from these questionnaires will enable us to explore attitudes and habits of patients with bronchiectasis and the healthcare team caring for them. Additionally this data will direct the areas of further research in bronchiectasis. Aims / objectives The aim of this project is to develop a questionnaire, to investigate physiotherapy use of Airway Clearance Treatments and their prescribing habits surrounding pharmacological and non-pharmacological Airway Clearance Treatments in Bronchiectasis. The developed questionnaire will eventually be used in a UK wide online platform to facilitate feedback. The primary objective is to compile a provisional questionnaire regarding Airway Clearance Treatments in Bronchiectasis. The secondary objective is to conduct a pilot test the questionnaire pack amongst experienced academic. research and clinical based physiotherapists to get their professional opinion regarding the content of

	the questionnaire and the language used in it, to ensure a user-friendly final version of the questionnaire. The methodology including recruitment will be facilitated by the academic supervisors. The feedback collected will direct changes to the questionnaire, allowing the final version of the questionnaire to be formed.	
Techniques employed:	<ul> <li>The student will have the opportunity to work on a MRC funded study and exposure to studies conducted within the clinical trial research facility.</li> <li>Questionnaire development</li> <li>Interviews</li> <li>Pilot testing</li> <li>Online survey launch activity</li> <li>Statistical analysis</li> <li>Literature review</li> </ul>	

Intercalated BSc in:	Medical Science Biochemistry Microbiology			
Project Title			d quantitative criteria to assess ndex data in bronchiectasis	sment
Supervisor(s)	<ol> <li>Dr Katherine O'Neill</li> <li><u>k.oneill@qub.ac.uk</u></li> <li>Prof Judy Bradley</li> <li><u>Judy.bradley@qub.ac.uk</u></li> </ol>			
School / Centre	Medicine, Dentistry and Bi		o for Experimental Medicine, Scho cal Science	ol of
Principal Supervisor's Contact Details	Email: <u>k.oneill@qub.ac.uk</u>		Tel:	
Degree Pathway	Medical Science	Х		
for which project	Biochemistry		]	
is suitable (√)	Microbiology			
Is project of suitable standard / subject for studentship	<i>General awards</i> Wolfson Foundation		Subject-specific awards British Assoc Dermatologists	
application? (✓)	Jean Shanks Foundation		Digestive Disorders Foundation	
			Pathological Society	
			Other	
Background information:	Lung clearance index is an emerging outcome measure that is increasingly being used in respiratory clinical trials and also in some clinical sites to explore early lung disease.			
	As it is a new measure, it is essential that there is appropriate quality control measures in place to ensure validity of the assessment.			lity
	The supervisors on this project have received funding from MRC to explore various outcomes including the use of a novel lung function measurement (lung clearance index) in bronchiectasis. This study is running over 8 sites in the UK. Belfast is acting as the core facility for quality checking (over reading LCI measurements).			n is
	A significant amount of dat available for quality assess		been collected in this study and is	5
	A recent paper Jensen et al 2016 PLOS ONE 11(6), 1-9, have developed a checklist and stepwise approach to facilitate the review of each lung clearance index trial. These approaches include assessment of quantitative criteria as well as qualitative criteria.			
	This project has the potent	ial to h	ave an abstract output.	
Aims / objectives	<ul> <li>The aim of this project will be to:</li> <li>a. Become familiarised with the steps included in the qualitative and quantitative criteria as detailed by Jensen et al 2016.</li> <li>b. Analyse the available data according to Jensen criteria.</li> </ul>			tive

Techniques employed:1. The student will have the opportunity to work on a MRC fu study and exposure to studies conducted within the clinical research facility.2. The student will become familiarised with lung clearance i which is a novel lung function measurement and increasin being used as an important outcome in clinical trials.3. The student will gain experience of how data is collected i multicentre clinical trials and transferred electronically to c sites.4. The student will use a range of statistical techniques to summarise relevant data	ıl trial ndex gly n

Intercalated BSc in:	Medical Science Biochemistry Microbiology			
Project Title	Exploring the relationship between Quality of life and lung function (FEV <sub>1</sub> and Lung Clearance Index) in Bronchiectasis (MRC funded clinimetrics project)			
Supervisor(s)		Prof Judy Bradley Judy.bradley@qub.ac.uk Dr Katherine O'Neill k.oneill@qub.ac.uk		
School / Centre	Clinical Research Facility, Medicine, Dentistry and Bio		for Experimental Medicine, Scho al Science	ol of
Principal Supervisor's Contact Details	Email: Judy.bradley@qub.ac.uk		Tel:	
Degree Pathway for which project is suitable (√)	Medical Science Biochemistry Microbiology	X		
Is project of suitable standard / subject for studentship application? (✓)	General awards Wolfson Foundation Jean Shanks Foundation		Subject-specific awards British Assoc Dermatologists Digestive Disorders Foundation Pathological Society Other	
Background information:	Bronchiectasis is a chronic respiratory condition with symptoms including difficulty clearing secretions as well as frequent chest infections and declining lung function. There are no licensed therapies for patients with bronchiectasis and it is not clear what outcome measures should be used in upcoming clinical trials. A large dataset of outcome measures including quality of life and lung function (FEV <sub>1</sub> and lung clearance index) has been collected from patients with bronchiectasis in the UK. Analysis of this database considering the relationship between quality of life and lung function (FEV <sub>1</sub> and lung clearance index), would yield an insight into the usefulness of these measures in monitoring patients with bronchiectasis. Analysis of results would be suitable for abstract submission and presentation.			
Aims / objectives	<ul> <li>The aim of this project will be to:</li> <li>a. Review the quality of life questionnaires used in terms of their validity, reliability and responsiveness in bronchiectasis.</li> <li>b. Become familiarised with lung clearance index data.</li> <li>c. Analyse the relationship between lung function and quality of life</li> </ul>			

Techniques employed:	<ol> <li>The student will have the opportunity to work on a MRC funded study and exposure to studies conducted within the Northern Ireland Clinical Research Facility.</li> </ol>
	<ol> <li>The student will become familiarised with outcome measures in bronchiectasis used in clinical trials.</li> </ol>
	3. The student will gain experience of how data is collected as part of a clinical trial.
	<ol> <li>The student will use a range of statistical techniques to summarise relevant data.</li> </ol>

# **Centre for Infection and Immunity**

Project Title	Pharmacological inhibition of DREAM to normalise the inflammatory immune response in cystic fibrosis			
Supervisor(s)	1. Dr Bettina Schock			
	2. Amal ElBanna			
School / Centre	Centre for Infection and In	nmuni	ty	
Principal Supervisor's Contact Details	Email: b.schock@qub.ac.uk		Tel: 02890 972258	
Degree Pathway	Medical Science	Х		
for which project	Biochemistry	Х		
is suitable (√)	Microbiology			
Is project of suitable standard / subject for studentship application? (√)	General awards Wolfson Foundation Jean Shanks Foundation	x	Subject-specific awards British Assoc Dermatologists Digestive Disorders Foundation Pathological Society Other	
Background information:	The ubiquitination protein A20 is an important negative regulator of the NF-kB pathway in inflammation. We have recently shown that A20 is reduced in Cystic Fibrosis (CF) chronic airway disease (epithelial cells). Further work indicated that the induction of A20 in CF might be repressed through increased expression of a repressor called DREAM and our recent ChIP data confirm a 'persistent binding of DREAM to its A20 promoter related binding site (DRE3 box). Through bioinformatical analyses of CF epithelial cells (connectivity mapping) we identified drugs predicted to reduce DREAM expression in CF epithelial cells and thereby to reduce CF airways inflammation.			
Aims / objectives	<ul> <li>This project will further investigate the expression and the role of DREAM in CF epithelial cells after stimulation of the cells with bacterial products (e.g. LPS).</li> <li>It is hypothesized that pharmacological reduction of DREAM will normalise the pro-inflammatory phenotype of CF airway epithelial cells, basally and after stimulation with bacterial products.</li> <li>This will be tested in airway epithelial cells from patients with CF and non-CF controls after inhibiting/reducing DREAM expression using drugs predicted by connectivity mapping. A20 expression (mRNA and protein) as well as the inflammatory response of these cells will be analysed.</li> <li>Using epithelial cells (cell lines) with and without a CFTR mutation, we</li> </ul>			
employed:	will use the following techniques: Tissue culture and sterile working techniques Quantitative real time PCR to detect A20, DREAM (and the A20 transcription factor USF1) mRNA Protein analyses for A20 and DREAM in treated cells using Western Blotting			

Project Title	Identification of the cytosolic DREAM binding protein in LPS stimulated cystic fibrosis epithelial cells.			
Supervisor(s)	1. Dr Bettina Schock			
	2. Amal ElBanna			
School / Centre	Centre for Experimental Medicine			
Principal Supervisor's Contact Details	Email:         Tel:           b.schock@qub.ac.uk         02890 972258		-	
Degree Pathway	Medical Science	Х		
for which project	Biochemistry	X	-	
is suitable (✓)	Microbiology	~	-	
	General awards	x	Subject-specific awards	
Is project of suitable standard / subject for	Wolfson Foundation	x	British Assoc Dermatologists	
studentship application? (✓)	Jean Shanks Foundation		Pathological Society	
	Other			
Background information:	The ubiquitination protein A20 is an important negative regulator of the NF-kB pathway. A20 is reduced in epithelial cells of patients with Cystic Fibrosis (CF) through increased expression of the repressor DREAM, which binds to the A20 promoter. Additionally, DREAM protein has cytosolic functions. Nuclear DREAM normally occurs as a dimer or a tetramer and regulates gene expression but our preliminary data also suggest that cytosolic DREAM binds to an approximately 20 kD protein. Mass spectroscopy analyses indicated some binding partner candidates that could be involved in the transport of DREAM into the nucleus.			
Aims / objectives	This project will further investigate the composition of cytosolic DREAM in LPS stimulated epithelial cells. The student will use sterile tissue culture techniques to grow epithelial cells and will investigate the expression of our candidate DREAM binding partner using quantitative real time PCR and immunoprecipitation/Western Blotting.			
Techniques employed:	Using epithelial cells (cell lines) with and without a CFTR mutation, we will use the following techniques: Tissue culture and sterile working techniques Quantitative RT-PCR Extraction of cytosolic proteins and protein analyses under non-reducing conditions for DREAM in using Western Blotting. Transferable skills (presentations/communication skills, organisation of work, working alone and in a team).			

# **Centre for Medical Education**

Due to at Titl			
Project Title	How can we prepare medical students and foundation trainees for safe and effective practice? PROJECT 1		
Supervisor(s)	1. Tim Dornan		
	2. Richard Conn/Richard McCrory		
School / Centre	Centre for Medical Education, School of Medicine		
Principal Supervisor's Contact Details	Email: t.dornan@qub.ac.uk	Tel: 02890 975773	
Degree Pathway	Medical Science		
for which project	Biochemistry		
is suitable (√)	Microbiology		
Is project of	General awards	Subject-specific awards	
suitable standard / subject for studentship	Wolfson Foundation	British Assoc Dermatologists Digestive Disorders Foundation	
application? ( $\checkmark$ )	Jean Shanks Foundation	Pathological Society Other	
Background information:	It is widely assumed that well designed undergraduate and foundation curricula can adequately prepare medical students and recently qualified doctors to practise safely, effectively, and happily. Evidence suggests otherwise. Efforts to strengthen curricula by intensifying teaching, assessment and introducing simulation have not yet proven successful. If you look at the nature of practice and the causes of clinical errors, that is unsurprising. Practice is far more complex than curricula suggest. Our group is investigating how people learn to prescribe (intravenous fluids, for example). We invite up to 2 intercalating students to join our team of medically qualified PhD students, social scientist, pharmacist, and senior education researcher. We will allocate two very motivated supervisors to each of them. Rather than specifying their exact projects now, we will tailor the projects to their individual wishes and needs. And according to where our research programme has reached, so that the research is very up-to-date. Interested students are invited to contact us for further information.		
Aims / objectives	<ul> <li>Methodically review relevant published articles relevant to a mutually agreed topic, probably related to prescribing but perhaps to some other topic in the domain of patient safety</li> <li>Conduct a piece of primary research related to the same topic, examining how learning could be made more effective</li> </ul>		
Techniques employed:	<ul> <li>Methodical literature survey; probably scoping review but, depending on the topic, an alternative methodology might be more appropriate</li> <li>Depending on the topic and the interests of the student(s), this may include interviewing students, qualified professionals, or patients; it may involve observing real clinical or educational practice; or it may involve some form of experimentation</li> <li>We hope the student(s) will present their work at conference(s); depending on the success of the project(s) these might also be published</li> </ul>		

Project Title	How can we prepare medic	al students and foundation trainees for safe		
Floject fille	and effective practice? PR			
Supervisor(s)	1. Tim Dornan			
	2. Richard Conn/Richard M	cCrory		
School / Centre	Centre for Medical Education	on, School of Medicine		
Principal	Email: t.dornan@qub.ac.uk	Tel: 02890 975773		
Supervisor's				
Contact Details				
Degree Pathway	Medical Science			
for which project	Biochemistry			
is suitable (√)	Microbiology			
Is project of	General awards	Subject-specific awards		
suitable standard		Dritich Asses Dress to be inte		
/ subject for	Wolfson Foundation	British Assoc Dermatologists Digestive Disorders Foundation		
studentship		Pathological Society		
application? ( $\checkmark$ )	Jean Shanks Foundation	Other		
Background	It is widely assumed that w	ell designed undergraduate and foundation		
information:	curricula can adequately prepare medical students and recently qualified			
	doctors to practise safely, effectively, and happily. Evidence suggests			
	otherwise. Efforts to streng	then curricula by intensifying teaching,		
		g simulation have not yet proven successful.		
		practice and the causes of clinical errors, that		
		far more complex than curricula suggest. Our		
		people learn to prescribe (intravenous fluids,		
		to 2 intercalating students to join our team of dents, social scientist, pharmacist, and		
		r. We will allocate two very motivated		
		1. Rather than specifying their exact projects		
		cts to their individual wishes and needs. And		
		earch programme has reached, so that the		
		Interested students are invited to contact us		
	for further information.			
Aims / objectives		evant published articles relevant to a		
		probably related to prescribing but perhaps to		
		domain of patient safety		
		hary research related to the same topic,		
	-	g could be made more effective		
Techniques		rvey; probably scoping review but,		
employed:		, an alternative methodology might be more		
	appropriate	and the interacts of the student(s) this man		
		and the interests of the student(s), this may dents, qualified professionals, or patients; it		
	3	real clinical or educational practice; or it may		
	involve some form of e			
		) will present their work at conference(s);		
		ess of the project(s) these might also be		
	published			

Project Title	Healthcare professionals Ireland: a scoping review		riences as immigrants to the U	K and
Supervisor(s)	1.Mairead Corrigan			
	2. Jenny Johnston			
School / Centre	CME			
Principal Supervisor's Contact Details	Email: <u>m.corrigan@qub.ac.uk</u> j.l.johnston@qub.ac.uk		Tel:	
Degree Pathway	Medical Science	х	-	
for which project is suitable (✓)	Biochemistry Microbiology		-	
	General awards		Subject-specific awards	
Is project of suitable standard			British Assoc Dermatologists	
/ subject for studentship	Wolfson Foundation		Digestive Disorders	
application? (√)	Jean Shanks Foundation		Foundation	
			Pathological Society	
Background information:	conduct a review of the em service students and perso topical and will offer a first tackle multiple cultural issu lives. This is an important p cultural diversity and media	pirical onnel ir step to les bot part of cal trai	D'Malley's scoping review framew literature around immigration of I in the UK and Ireland. This is high owards supporting students as the h in their professional and persor a current stream of research look ning. The student's work will cont int topic and help to develop appro	nealth ly ey nal king at ribute
Aims / objectives	2. Identify gaps for po	otentia		
Techniques employed:	underpinning epistemology Learn how to do database synthesise results	/ search bmit a	ostantive literature review, includi ning, manage large sets of results poster to a national conference ir blication	and

# **Centre for Public Health**

Project Title	Deep phenotyping and g	notic :	analysis for Behçet's disease – a	
	complex, multifactorial ra			
		ile uise	case	
Supervisor(s)	1. AJ McKnight			
	2 <mark>. Postdoc starting in Jan</mark>	<mark>uary 2(</mark>	017-2019	
School / Centre	SMDBS – Centre for Pub	lic Hea	lth	
Principal	Email: a.j.mcknight@qub.	ac.uk	Tel: 02890 638460 (shared line	e)
Supervisor's			, , , , , , , , , , , , , , , , , , ,	,
Contact Details		_		
Degree Pathway	Medical Science	Yes		
for which project	Biochemistry	Yes		
is suitable (√)	Microbiology	No		1
	General awards		Subject-specific awards	
Is project of		N.	British Assoc Dermatologists	Mar
suitable standard	Waltaan Faundation	Yes	British Assoc Dermatologists	Yes
/ subject for studentship	Wolfson Foundation		Digestive Disorders Foundation	Don't
application? (✓)	Jean Shanks	Yes	Pathological Society	know
	Foundation		Other	
Background	The European Union (EU)	) dofini	tion of a rare disease is one that	at affacts
	≤5 per 10,000 people, with ~30 million persons directly affected in the EU and ~106,000 affected in Northern Ireland. These diseases are individually rare, but collectively common and represent a significant public health problem. This project investigates inherited risk factors and the impact of living with a selected rare disease: Behçet's disease (BD). Genetic and environmental factors contribute to BD, but the process of diagnosis is challenging with inconsistent clinical manifestations of this disease. A recent survey of individuals living with rare disease(s) in Northern Ireland revealed ~50% of individuals receive ≥1 misdiagnosis with <sup>1</sup> / <sub>20</sub> seeing >10 doctors. Individuals with BD report a wide range of symptoms, which are variable in onset, severity, and frequency of flareups for this systemic vasculitis. This disease involves abnormal inflammation / immune responses and common features include recurrent ulcers, skin lesions, and serious eye inflammation.			
	practitioners revealed a 12.6/100,000 in the No expected 'UK' prevalence	much rthern e, and t	UK. Recent mapping through higher than expected preva Ireland population. This high he identification of several fam makes NI ideal to explore ger	lence of her than ilies with

	This project involves deep phenotyping and strategies to improve recognition of Behçet's disease, identify genetic risk factors, and improve data sharing.
Aims / objectives	The primary aim of this project is to survey patients affected by Behçet's disease, identify genetic risk factors associated with BD in Northern Ireland, and evaluate information sources for patients.
Techniques employed:	This project will involve generating data from online surveys and focus groups, as well as state-of-the-art genotyping (next generation sequencing and / or high density microarrays) to analyse more than one million unique genetic markers for association with Behcet's Disease in a Northern Ireland population.
	For the dedicated student, this project may also include working with multiple stakeholders and using mixed methodological approaches to evaluate access to appropriate information sources, the social impact of living with this rare disease, and evaluate mental health and wellbeing.

Project Title	Gene-environment intera Degeneration	actions	s in Age-related macular	
Supervisor(s)	Amy Jayne McKni	ght & F	Ruth Hogg	
School / Centre	Centre for Public Health			
Principal Supervisor's Contact Details	Email: a.j.mcknight@qub.ac.uk		Tel: (0)28 9097 6359	
Degree Pathway	Medical Science	Х		
for which project	Biochemistry			
is suitable (√)	Microbiology			
	General awards		Subject-specific awards	
Is project of				
suitable standard			British Assoc Dermatologists	
/ subject for studentship	Wolfson Foundation		Digestive Disorders Foundation	
application? ( $\checkmark$ )	Jean Shanks Foundation		Pathological Society	
			Other	
Background information:	significant proportion of the recent report highlighting 5 variants across 34 loci <sup>1</sup> , he also well established that is contribute to the developm the characteristic phenotyp have been to date no large cohorts well-phenotyped for demographic and environm the disease. The Northerm Aging provides such an op have underwent an extension health assessment which is retinal images (colour, OC field Optomap images) have phenotypes such as reticut deposits. This project wout data with environmental rist related phenotypes.	e herita 52 inde 52 inde 50 wever 50 orthoge 50 orthoge	s have proved successful in revea ability related to AMD, with the mo- pendently associated common ar missing heritability still remains. enetic and environmental factors AMD, but how these interact to re AMD is not well understood. The population based studies involvir as well as characterised for the sk factors known to be associated d Cohort for the Longitudinal Stud- ity as approximately 4,500 partici- me interview, dietary assessment multi-modal retinal imaging. The a-red, autofluorescence and ultra- n graded for AMD including novel budodrusen/subretinal drusenoid to relate genome-wide association for and presence of various AMD	ost nd rare It is esult in re ng many d with dy of pants and e wide
Aims / objectives	risk factors in AMD throug	h the u	een genetic risk loci and environm se of a genome-wide scan in a w udy in Northern Ireland (NICOLA	

Techniques employed:	Bioinformatics, multivariate statistical analysis, retinal grading.	

Project Title	Retinal evaluation in dial	betic k	idney disease – an early bioma	rker?
Supervisor(s)	1. Dr Gareth McKay			
	2. Prof Peter Maxwe			
School / Centre	Centre for Public Health			
Principal Supervisor's	Email:		Tel:	
Supervisor's Contact Details	g.j.mckay@qub.ac.uk		90978958	
Degree Pathway	Medical Science	$\checkmark$		
for which project	Biochemistry			
is suitable (√)	Microbiology			
	General awards		Subject-specific awards	
Is project of suitable standard			British Assoc Dermatologists	
/ subject for	Wolfson Foundation		Digestive Disorders Foundation	
studentship application? (✓)	Jean Shanks Foundation		Pathological Society	
Background	Clinical data and rational	imaaa	Others from the Northern Ireland D	iohotia
information:	Retinopathy Screening P respect to renal function ratio). These individuals w glomerular filtration (eGFF (chronic kidney disease C in whom eGFR measurem the preceding 5 year scree this "stable" kidney function data in individuals with pr group will contain individu <60mL/min/1.73m <sup>2</sup> . Retinat with risk of developing pro- research will help determin part of the diabetic retinan additional clinical utility in i renal failure. The retinal vasculature is invasive assessment enab- prior to clinically significant photography and <i>in sill</i> characterisation of retinal w <b>Improved non-invasive r</b> prognostic clinical tool.	rogram (serun ill be si kD sta hents ro ening p on grou ogressi ials in al vesso ogressi he if me nopath dentify directly bling de t eventsi <i>ico</i> mo vascula <b>isk str</b> a	me (NIDRSP) will be assesse in creatinine; urinary albumin/cre tratified into two groups using est titions. The stable kidney function ge 1 and stage 2) are those indiv- emained >60mL/min/1.73m <sup>2</sup> thro- beriod. Retinal vessel measurem- up will be compared with retinal sive kidney failure. This "kidney f whom eGFR declined over 5 ye el measurements will then be com- ve diabetic kidney disease (DKD easuring the retinal microvasculat y screening programme, would ing patients at highest risk of dever accessible to direct and repeate etection of early microvascular ch s. Recent improvements in digital easurement have enabled im- ar and structural parameters. <b>atification for DKD</b> is a highly de	d with atinine imated group viduals ughout ents in vessel failure" ears to related 0). This ure, as d offer eloping ed non- nanges retinal proved sirable
Aims / objectives	study will collect and analy novel prognostic indicator diabetes at increased ris	se retro s for e <b>k of re</b>	s recruited almost 300 participants ospective data from NIDRSP to ex early identification of <b>individual</b> s anal decline. inal variation is associated with	/aluate s with

	Techniques employed:	This study will use VAMPIRE (Vascular Assessment and Measurement Platform for Images of the REtina) software to identify and measure retinal microvasculature changes from digital photographs acquired prospectively. The standardised measured area is defined within the region 0.5-2.0 disc diameters from the optic disc margin. Fractal analysis will quantify geometric branching complexity and density of retinal vessels providing a holistic overview of retinal microvascular health. Logistic regression analysis will evaluate potential correlations between retinal microvascular variation and renal decline.
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Project Title	Can Autofluorescence be	e used	as a non-invasive marker of o	uter
	retinal dysfunction in dia			
Supervisor(s)	1. Ruth Hogg			
	2. Tunde Peto			
	2. Tunde Peto			
School / Centre	Centre for Public Health			
Bringing		L.	Tab 028 00071054	
Principal Supervisor's	Email: r.e.hogg@Qub.ac.u	K	Tel: 028 90971654	
Contact Details				
Degree Pathway	Medical Science	Х		
for which project	Biochemistry		]	
is suitable (✓)	Microbiology			
la marte de la	General awards		Subject-specific awards	
Is project of			British Assos Dermetalegista	
suitable standard / subject for	Wolfson Foundation		British Assoc Dermatologists	
studentship	Wonson i oundation		Digestive Disorders	
application? $(\checkmark)$	Jean Shanks Foundation		Foundation	
			Pathological Society	
			Other	
Background	Retinal auto fluorescence	(AF) in	haging has attracted much interes	t over
information:	the past 20 years as it prov	vides a	non-invasive method of assessir	
			retinal pigment epithelium (RPE).	
			mulates in the RPE during aging	
			of photoreceptor outer segments	
			and senescence. Patterns of incre een observed in a variety of retina	
			pathy. However its clinical applic	
			lity in making between-person	
			f ocular media changes not relate	d to
	the retina on the absolute	values	. Recently a method has been	
			ially available Heidelberg Spectra	
			that enables quantitative AF to b	e
			comparisons more feasible. This	ina
			nship between qAF and both age lish if qAF imaging could be used	
			dysfunction in the development of	
	Diabetic Retinopathy.	. canal		-
Aims / objectives			ip between qAF and age in a	
	population based of		of a A E diffore between these will	-
			of qAF differs between those with pared to age-matched controls, and	
			qAF differ between with increasin	
	severity of DR			3
	3. Clarify the relation		etween the extent of qAF and othe	
			ted to more severe DR such dura	ation
	of diabetes, body	nass i	ndex, hbA1c	

Techniques employed:	Retinal grading, multivariate statistics, literature review.
empioyeu.	

Project Title			neral retinal Autofluorescence in ion and diabetic retinopathy?	
Supervisor(s)	1. Ruth Hogg			
	2. Tunde Peto			
School / Centre	Centre for Public Health			
Principal Supervisor's Contact Details	Email: r.e.hogg@Qub.ac.uk		Tel: 02890971654	
Degree Pathway	Medical Science	Х		
for which project	Biochemistry			
is suitable (✓)	Microbiology			
	General awards		Subject-specific awards	
Is project of suitable standard / subject for	Wolfson Foundation		British Assoc Dermatologists	
studentship			Digestive Disorders	
application? ( $\checkmark$ )	Jean Shanks Foundation		Foundation	
			Pathological Society	
			Other	
Background information:	years to assess the accum product of photoreceptor of increased lipofuscin accum stress and inflammation, b development of age-relate retinopathy (DR). The relev conditions is being increas autofluorescent patterns is learning to evaluate Ultra-v Autofluorescent changes a these and traditional featur conventional imaging. The Cohort Longitudinal study	nulation nulation oth import d mace vance ingly ro- not wo wide file and inv res of <i>i</i> e data of Agin	ular degeneration (AMD) and diabeti of the peripheral retina in both ecognized but the role of peripheral ell understood. This project involves eld retinal images for peripheral retin estigating the relationship between AMD and DR captured using used will be from the Northern Irelan og (NICOLA) study.	ative ic S nal
Aims / objectives		heral f	luorescent changes in a population	
	changes and pres	ence o		
	<ol> <li>Investigate the relation of the set of the</li></ol>		ip between peripheral autofluoresce	ent
	changes and pres	ence o	I DR IEdiules.	

Techniques employed:	Retinal grading, epidemiology, multivariate statistical analysis, literature review.

Project Title	Impact of maternal weight gain during and post-pregnancy on maternal insulin resistance and neonatal outcome				
Supervisor(s)	1. Dr Una Graham				
	2. Prof. David McCance				
School / Centre	Centre for Public Health				
Principal	Email:		Tel:		
Supervisor's Contact Details	u.graham@qub.ac.uk		02890633423		
Degree Pathway	Medical Science	Υ			
for which project	Biochemistry				
is suitable (√)	Microbiology				
	General awards		Subject-specific awards		
Is project of					
suitable standard			British Assoc Dermatologists		
/ subject for studentship	Wolfson Foundation		Digestive Disorders Foundation		
application? ( $\checkmark$ )	Jean Shanks Foundation		Pathological Society		
			Other		
Background information:	Over 60% of pregnant mothers in Northern Ireland are overweight (BMI 25-30kg/m <sup>2</sup> ) or obese (≥30kg/m <sup>2</sup> ). Maternal obesity is a worldwide epidemic associated with a significant increase in maternal, fetal and neonatal complications including gestational diabetes mellitus (GDM). Offspring of mothers affected by GDM subsequently have an increased likelihood of diabetes and obesity at a young age. This transgenerational effect has the potential to create a vicious cycle with obesity and diabetes developing in future generations. Major public health efforts are needed to break this cycle. Whilst the association between maternal obesity and GDM is clear, the relative impacts of intra-pregnancy, interpregnancy and immediate post-natal weight changes on insulin sensitivity during and post-pregnancy have not been reported. This data would enable public health strategies to be directed towards women at a time when they, and their future offspring, are most likely to benefit from intervention.				
Aims / objectives	<ol> <li>To determine the impact of maternal weight changes at 3 time points on insulin sensitivity (as measured by the insulin response during the oral glucose tolerance test). Timepoints: 1. From early pregnancy (&lt;14weeks gestation) to late second trimester (24-28 weeks gestation),</li> <li>From early pregnancy (&lt;14 weeks) to early post-pregnancy (within 8 weeks of delivery) and 3. From index pregnancy (&lt;14 weeks) to subsequent pregnancy (&lt;14 weeks)</li> <li>To assess the impact of maternal weight changes and insulin sensitivity on neonatal weight and outcome</li> </ol>				

Techniques	This study will enable the student to develop extensive skills in data		
employed:	extraction, processing and interpretation; key skills for clinical research.		
	The student will be given an honorary contract within the BHSCT and		
	have the opportunity to attend specialist diabetes in pregnancy clinics,		
	and education sessions within the Regional Centre for Diabetes and		
	Endocrinology. Data relating to maternal weight changes, biochemical		
	results and neonatal outcome will be extracted from clinical databases		
	and combined with data from paper records to generate an extensive		
	dataset of women evaluated for GDM over the past decade. To assist		
	with this the student will be given the opportunity to attend training in		
	both excel and SPSS. Data analysis will be performed by the student		
	supported closely by the project supervisors and biostatisticians in the		
	CPH. It is anticipated that the student will present their results at local		
	and National conferences in Diabetes (Irish Endocrine Society Annual		
	Meeting, Diabetes UK Annual Conference); and assist in the preparation		
	of a manuscript for International publication.		
	of a manuscript for international publication.		

# <u>RISUS (Rugby Injury Surveillance</u> <u>in Ulster Schools) group and</u> <u>QUB Centre for Biomedical</u> <u>Sciences Education</u>

Intercalated BSc ir	n: Medical Science Biochemistry Microbiology			
Project Title	Modifying injury risk in Schools Rugby with implementation of TACKLESMART IRFU guidelines (Projects available for 3 students)			
Supervisor(s)	<ol> <li>Pooler Archbold (RISUS (Rugby Injury Surveillance in Ulster Schools) Group)</li> <li>2.Dr. Sean Roe QUB</li> </ol>			
School / Centre	Centre for Biomedical Sciences Education			
Principal Supervisor's Contact Details	Email: poolerarchbold@aol.com s.roe@qub.ac.uk	Tel: 02890972640		
Degree Pathway for which project is suitable (✓)	Medical Science Biochemistry Microbiology			
Is project of suitable standard / subject for studentship	General awards Wolfson Foundation Jean Shanks Foundation	Subject-specific awards British Assoc Dermatologists Digestive Disorders Foundation		
application? (✓)		Pathological Society Other		
Background information:	Other         Other           Identifying modifiable risk factors is central to injury surveillance and prevention initiatives. Recent studies have shown that poor tackling technique is associated with an increased risk of injury and concussion in Rugby Union.           To reduce the risk of injury in School's Rugby a new TACKLESMART program will be implemented at the start of the 2017/8 season.           The aim of this project is to implement the TACKLESMART program within the U-15 level of School's Rugby. Data collected will include player demographics, biometrics, and strength, previous history of injury, level of play and the use of protective equipment. The information gathered will assess if the implementation of new tackling guidelines and techniques reduce the severity and number of injuries including concussions in School's rugby.			