# Centre for Cancer Research and Cell Biology

Project Title	Examining the influence treatment pattern in Oese		loscopic Therapy on diagnosis eal Adenocarcinoma	and
Supervisor(s)	1. Helen Coleman			
	2. Richard Turkington			
School / Centre				
Principal Supervisor's	Email: h.coleman@qub.ac.uk		Tel: 02890 972756	
Contact Details				
Degree Pathway for	Medical Science	$\checkmark$	4	
which project is suitable (✓)	Biochemistry Microbiology	v	4	
Suitable (* )	General awards		Subject-specific awards	
Is project of				
suitable standard /			British Assoc Dermatologists	
subject for studentship	Wolfson Foundation		Digestive Disorders Foundation	
application? (✓)	Jean Shanks Foundation		Pathological Society	
			Other	
Background information:	Five-year survival rates for oesophageal adenocarcinoma (OAC) r poor at 15% and the incidence has increased six-fold over the pas decades. For tumours which have invaded beyond the submucos			st four
	standard of care involves s	surgica	I resection which preceded by ne	0-
	adjuvant chemotherapy. C	) esoph	ageal cancer surgery carries a high	gh
	morbidity (50%) and mortality (2-3% 30-day in hospital mortality) and			
	patients report a significant and long-lasting deterioration in health-			
	related quality of life following surgery. Standard care for high grade dysplasia (HGD) and early OAC tumours confined to the submucosa ha			
			on. However the introduction of	
	, ,		care over the last decade has	
			ns available for Barrett's oesoph	ວດເມດ
		-	agus with evidence low grade dy	-
		•	• • • •	•
		•	ident pathologists, or high grade	
			. For tumours confined to the	
		-	e oesophagus (stage T1a) endosc	opic
	resection combined with a	blative	e therapy is curative and organ-	
	preserving treatment. By	contra	st, patients with disease extendi	ng
	beyond the mucosal lining	and in	to the underlying supportive tiss	sues
	(stage T1b) are at risk of Ic	oco-reg	ional lymph node metastasis and	b
	these patients are therefo	re offe	red radical oesophageal resectio	n
			ment. The option of endoscopic	
	•		equirement of double reading of	
				may
	Barrett s Desopriagus blop	3185 10	confirm low grade dysplasia and	iiidy

	have led to a relaxation of the criteria for the diagnosis of high grade
	dysplasia as previously this would have led to surgical resection. We seek to examine the impact the introduction of endoscopic therapy has
	had on the trends in diagnosis of both low and high grade dysplasia.
	We will also examine the outcomes of endoscopic ablation and resection in comparison with international standards.
	The successful student would become fully integrated with the Cancer Epidemiology and Health Services Research Group in the Centre for
	Public Health, which would include attendance of weekly meetings to
	learn of other ongoing research in the group, and to evaluate
	epidemiological study designs at journal clubs.
	Students will be exposed to clinical collaborators in the Barrett's
	oesophagus research team including epidemiologists,
	gastroenterologists and pathologists who regularly meet, and so the student should have a strong interest in these medical specialties, or
	paediatrics. Full guidance and support will be provided for interpreting
Aime / chiectives	the statistical analysis and results.
Aims / objectives	Aim 1: A Descriptive Epidemiological Study of Endoscopic Therapy in NI
	Since the introduction of endoscopic therapy in Northern Ireland ten years ago over 300 ablation and resection procedures have been carried
	out. This number of procedures and the duration of follow up provides
	a unique opportunity to study the survival outcomes for these procedures and to compare these with standards in the published
	literature. We seek to integrate the Endoscopic Therapy register with
	the Northern Ireland Barrett's register in order to identify cases of
	Barrett's oesophagus which have progressed to dysplasia and required resection or ablation.
	Aim 2: A Concordance study of Endoscopy Records and Oesophageal Pathology Reports
	In order to develop a comprehensive database of endoscopic therapy
	for future work a concordance study will be performed to match
	endoscopy records with the pathology reports from oesophageal
	biopsy specimens.
	Aim 3: Determination of trends in referral of Dysplasia following the
	introduction of endoscopic therapy Current guidelines state that all HGD cases and cases where LGD is
	found on two separate occasions should be referred for endoscopic
	therapy. We seek to examine the referral patterns in NI following the
	introduction of the current guidelines for endoscopic therapy.

Techniques	Literature review
employed:	Critical appraisal of papers
	Scientific writing
	Knowledge in public health, gastroenterology and cancer epidemiology Data collection Statistical analysis

Intercalated BSc in	n: Medical Science Biochemistry			
	Microbiology			
Project Title			PK pathway in mediating resistan	ce to
	radiation/hormone therapy			
Supervisor(s)	1.Richard Kennedy			
	2. Nuala McCabe			
School / Centre	CCRCB			
Principal	Email: r.kennedy@qub.ac.u	k	Tel:	
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	Joon Charles Favor dation		Digestive Disorders Foundation	
application? (✓)	Jean Shanks Foundation		Dathalagical Society	
			Pathological Society	
			Other	
Background	Prostate cancer (PCa) is the	most	commonly diagnosed cancer in me	n in
information:	is radiation and androgen de therapies represents a majo This study aims to identify m radiation resistance and res androgen receptor signaling prostate cancer in 2012. We have identified a molecu by MAPK signalling and rep 50% metastatic prostate car number of roles involved in proliferation, migration and o of the MAPK pathway in the resistance using a panel of resistant PCa cell lines. We signaling and markers of ep parental and resistant cell lin Furthermore we aim to inves parental and radiation/enzal (Trametinib) and inhibitors of	epravai r barrie nechan istance l, appro ular sub resents nerstat drug re develo in hous aim to ithelial nes. stigate utamid of EMT nibition	re for stage III/IV prostate cancer p tion therapy; however resistance to er for the treatment of PCa. isms involved in the development of to enzalutamide, a potent inhibitor oved for the treatment of castrate re ogroup in prostate cancer which is of 30% of primary prostate cancers a The MAPK pathway is implicated in e cancer progression including sistance. We aim to investigate the opment of radiation and enzalutamic e generated radiation and enzaluta investigate differences in MAPK mesenchymal transition (EMT) between e resistant cell lines to MAPK inhibit (R428, Cabozantinib). Finally we w resensitises radiation/enzalutamide	these of of esistant driven and of a role de umide ween ition rill
Aims / objectives	<ul> <li>radiation/enzalutam</li> <li>2. Investigate the sensitive to inhibitors of</li> <li>3. Investigate the efficiency</li> </ul>	nide res sitivity MAPK acy of nide tre	ng and markers of EMT in sistant cell lines v parental cells. of radiation/enzalutamide resistant of and EMT signalling. MAPK/EMT inhibitors in combination atment. Do these agents resensitist ment?	on with
Techniques	Western blot; Quantitative	PCR;	cell culture; drug sensitivity assay	′S,
employed:			eration assays; migration assays;	
	invasion assays.			

Project Title	Optimising immune chec	kpoin	t therapy in ovarian cancer	
Supervisor(s)	1. Richard Kennedy 2. Eileen Parkes			
School / Centre	CCRCB			
	COROD			
Principal Supervisor's	Email: r.kennedy@qub.ac.	uk	Tel: 028 9097 2443	
Contact Details				
Degree Pathway	Medical Science			
for which project	Biochemistry	х		
is suitable (√)	Microbiology			1
1	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	Immune checkpoint-tar	geted	therapy (IO) has resulted in lin	nited
information:		-	varian cancer. A key research	
	-		onse to these agents in the cli	nic
	1 · · ·	• ·	innate immune pathway has b	
	identified as synergistic with anti-PD-1 therapy, and also a means			
	of overcoming resistance to IO. We have reported activation of			
	the cGAS-STING immune pathway as a result of cytosolic DNA			
	released in response to intrinsic and extrinsic DNA damage,			
	and that upregulation of PD-L1 in response to DNA damage is			
	dependent on STING.			
	Therefore, activating the STING pathway as a combination			
	treatment with IO could	result	in improved clinical response	s. We
			compounds, which can quickly	
			tion of PD-L1 expression in ov	
			vill then be validated for STING	
		•		
			mune checkpoints using prima	•
			curate model of tumour beha	viour
			er-associated ascitic fluid in	
	,		nds will be studied in combina	
	with IO using the ID8 Tr	o53-∕-E	Brca2-/- syngeneic mouse mode	el. In
	addition, compound tox	icity w	ill assessed using the normal	
	fallopian tube cell line FT190. Moreover, using ascitic fluid			
			ne cells from patients with ova	rian
	•		ation therapy using identified	
			umour response, and activatio	n of
	•		reatment combination selecte	
	initialie response. The	uedi li		u

	using these methods will be that with the least toxicity and
	greatest improvement in tumour response to IO.
Aims / objectives	(1) Identify drugs which activate the cGAS-STING innate immune
	pathway and subsequent PD-L1 gene expression
	(2) Validate identified hits in established and novel primary
	HGSOC cell lines by confirmation of cGAS-STING-PDL1 immune
	pathway activation.
Techniques	Cell Culture
employed:	qPCR
	Western blot
	In cell western
	High throughput screen
	Flow cytometry

Project Title	Identification of Targetak Oesophageal Adenocarc		diators of Drug Resistance in (OAC)
Supervisor(s)	<ol> <li>Richard Turkington</li> <li>Richard Kennedy</li> </ol>		
School / Centre			
Principal Supervisor's Contact Details	Email: r.turkington@qub.ac.uk		Tel: 02890 972756
Degree Pathway for	Medical Science	✓	
which project is	Biochemistry	✓	
suitable (✓)	Microbiology		
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:	poor at 15% and treatment not changed over the past resistance limits the effecti used to treat OAC and the resistance and novel agen aim to identify pathways of and analysis of suitable in Unravelling the mechanism	t strate two de veness discov ts to ta cispla vitro m ns of p ed in e	bhageal adenocarcinoma (OAC) remain gies for Her2-negative tumours have ecades. The development of drug s of current chemotherapeutic agents very of underlying mechanisms of arget these pathways is a priority. We tin resistance through the development nodels and pre-chemotherapy biopsies. rimary resistance will allow ineffective arly stage OAC and will also inform the tions of therapeutics.
Aims / objectives	Aim 1: Validation of Genes associated with Resistance to Chemotherapy in OAC To identify pathways and genes associated with resistance to chemotherapy in early stage OAC we have employed a systems biology approach. We have performed transcriptional profiling of 273 formalin fixed paraffin embedded pre-treatment endoscopic OAC biopsies using the Almac Diagnostics Xcel <sup>TM</sup> array. All OAC patients were treated with cisplatin-based neo-adjuvant chemotherapy followed by surgical resection between 2003 and 2014 at four UK centres. Following normalisation and filtering of the microarray data, pathway and functional enrichment analysis was applied to the resultant gene-set to determine clusters of significantly enriched pathways and Gene Ontology processes. Functional enrichment analysis was performed using Gene Set Enrichment Analysis (GSEA) on the differentially expressed gene lists. We hypothesize that pathways differentially regulated in relation to		

pathological response may be strong determinants of drug resistance in early stage OAC and so will be particularly relevant.

Genes related to the pathways of resistance are currently being assessed by focused siRNA (siRNA, Sigma) screen. We have selected an *in vitro* model representative of chemo-resistance in OAC by aligning transcriptional data according to published methods and those developed by Dr Jaine Blayney (Department of Bioinformatics,Queen's University Belfast). Candidate genes have been selected based on their fold change, biological importance in OAC and potential to be targeted. A focused screen of 84 genes will be performed in triplicate in Q1 2018 to study the effects on cell viability/cytotoxicity of gene silencing, either alone or in combination with cisplatin/5-FU, to discover targets which are not toxic in their own right but interact synergistically with chemotherapy. We anticipate that this screen will generate a number of promising leads and insights into drug resistance in OAC. The prospective student will select one of the candidates from this primary screen for further validation and development.

#### Aim 2: Discovery and mechanistic analysis of a novel drug target in OAC

A potential novel drug target will be validated in a panel of oesophagogastric cell lines with differing mutational contexts. Mechanistic analysis will be performed to discover their mode of action. We will determine the synergism of siRNA mediated knockdown of the selected target with cisplatin/5-FU in a panel of cell lines using MTT assays, combination index values and annexin V/propidium iodide flow cytometry. Western blotting will be performed for markers of apoptosis, such as PARP and cleaved caspase 3, and caspase activity assays will be carried out. Further examination of the effects of the target inhibition will be examined by Western blotting of relevant proteins, 14 day clonogenic assays and DNA repair assays eg comet assays. Should small molecule inhibitors be available for the selected targets these will also be evaluated for their apoptotic and mechanistic effects.

# Aim 3: Development of pre-clinical models representative of Cisplatin resistance in OAC.

Research into OAC is currently being hampered by a lack of in vitro cell lines which accurately model patient tumours and recapitulate clinical drug responsiveness. We are currently establishing novel primary cell lines using fresh OAC tissue collected during oesophageal staging and surgery at the Belfast City Hospital. This work will be carried out in collaboration with the OCCAMS consortium and will also include the storage of fresh frozen tissue for future research. Specimens will be transferred directly from the operating theatre to the research laboratory in complete DMEM media on ice, washed three times with 10ml of phosphate buffered saline, dissected into approximately 3mm<sup>3</sup> pieces with a scalpel and digested with trypsin-EDTA. Undigested segments will be removed by sedimentation and the clear supernatant spun at 600g for 5 minutes. Cell pellets will be cultured in complete DMEM under standard cell culture conditions. Immortalisation will be performed by lentivirus transfection and oncogene activation. Our group has already successfully established primary cell lines in breast and ovarian cancer and has developed

	optimised standard operating procedures for cell line generation. A cell line representative of cisplatin-resistance will then be used as a model to test the targeting of genes identified in Aim2. In this way we will develop models more representative of oesophageal tumours.
Techniques employed:	Cell Culture qPCR Western blot siRNA knockdown Flow cytometry

# **Centre for Experimental Medicine**

Project Title	INVESTIGATING THE INF ENDOTHELIAL PROGEN		ICE OF OXIDATIVE STRESS ON CELL FUNCTION	J
Supervisor(s)	1. Dr David Grieve 2. Dr Karla O'Neill			
School / Centre	СЕМ			
Principal Supervisor's Contact Details	Email: <u>d.grieve@qub.ac.ul</u>	<u>&lt;</u>	Tel: 028 9097 6468	
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 studentship	Wolfson Foundation	✓	Digestive Disorders Foundation	
application? ( $\checkmark$ )	Jean Shanks Foundation	✓	Pathological Society	
			Other	
Background information:	ischaemic cardiovascular of therapeutic potential of en- mobilised by ischaemia an group has characterised a endothelial colony-forming progenitor properties which health and disease. Oxidar are known to play a key ro evidence suggests that the Interestingly, we have sho stress, display differential	disease dotheli d are i distinc cells ( n prom tive str le in ca ey may wn tha gene e nodula	to influence the progression of e. Recent attention has focused of al progenitor cells (EPCs), which mportant in vascular homeostasis et EPC subtype, termed outgrowth ECFCs), with well-defined endoth ote new blood vessel formation in ess, and specifically NADPH oxid ardiovascular disease and emerg also regulate EPC function. t ECFCs are influenced by oxidat xpression compared to mature ted by hypoxia which is a charact vironment.	are s. Our n helial n both dases, ing ive
Aims / objectives	This project therefore aims to investigate the specific influence of oxidative stress and NADPH oxidases on in vitro ECFC function. It is hoped that the results will identify key pathways which may become dysregulated in disease and could represent potential targets to enhance the reparative capacity of these cells and their clear potential for the treatment of ischaemic cardiovascular disease.			
Techniques employed:	oxidases on ECFC functio treated with pro-oxidant co specific inhibitors of candio Expression of key signallin	n, stud mpour date pa ig gene nd in v	ts of oxidative stress and NADPH ies will be undertaken in cultured nds in the presence or absence o athways or after genetic manipula es will be quantified by real-time F ritro ECFC migration and prolifera ss functional effects.	cells f tion. RT-

Project Title       Testing the anti-microbial effect of leukotriene antagonist zafirlukast Mycobacterium xenopii and Mycobacterium malmoense         Supervisor(s)       1. Cecilia O'Kane         2. Danny McAuley       School / Centre	t on		
Supervisor(s)     1. Cecilia O'Kane       2. Danny McAuley			
2. Danny McAuley			
2. Danny McAuley			
Sahaal / Contro SMDDC Contro for Experimental Madicine			
School / Centre SMDBS Centre for Experimental Medicine			
Principal Email: Tel:			
Supervisor's c.okane@qub.ac.uk 02890976384			
Contact Details			
Degree Pathway Medical Science x			
for which project Biochemistry			
is suitable ( $\checkmark$ ) Microbiology x			
General awards Subject-specific awards			
Is project of			
suitable standard British Assoc Dermatologists			
/ subject for Wolfson Foundation			
studentship Digestive Disorders Foundation			
application? (✓) Jean Shanks Foundation Pathological Society			
Other			
<b>Background</b> There has been an exponential rise in the prevalence of non-			
information: tuberculous mycobacteria (NTM) in respiratory sputum isolates ove	r the		
past 3 decades. These organisms cause frequently intractable infec			
in patients with pre-existing structural lung disease, particularly COI			
	I D		
	and bronchiectasis. Within this group of infections, Mycobacterium malmoense and Mycobacterium xenopii, are associated with high level		
	morbidity and mortality. Treatment to date focuses on 3-4 antibiotics,		
	which are difficult to tolerate, and associated with poor rates of cure. In		
our lab we have recently found that a drug currently used for treatm			
of asthma (drug X) has significant anti-microbial activity against othe			
NTM species. This is incredibly exciting as these infections are high			
resistant to multiple antibiotics. Drug X is a safe, well-tolerated drug			
currently in use in clinical practice. If it has efficacy against M xenop			
and M malmoense this could potentially lead to shorter, more effect	live		
and more easily tolerated antimicrobial therapy for patients.			
Aims / objectives This study will test	4 - 1 - 11		
<ol> <li>the ability of drug X at clinically achievable concentrations,</li> </ol>	ιο KIII		
M xenopii in the laboratory	4 - 1 - 11		
<ol> <li>the ability of drug X at clinically achievable concentrations,</li> </ol>	to kill		
M malmoense in the laboratory			
Techniques Bacterial culture			
employed: Bacterial viability assays (Bacter-Glo)			
Bacterial quantification by spectrophotometry and colony counting			
If possible, depending on student's progress, some basic cell cultur	е		
and infection assays			

Project Title	A20 and DREAM in pulm	onary	fibrosis	
Supervisor(s)	<ol> <li>Dr Bettina C Schock (QUB, expertise: Inflammation, A20, DREAM)</li> <li>Amal ElBanna (QUB, Technical support, day-to-day laboratory supervision, expertise: cell culture, mRNA and protein analyses)</li> <li>Prof John Varga (Feinberg School of Medicine, Director, Northwestern Scleroderma Programme, expertise: scleroderma)</li> </ol>			
School / Centre	Centre for Experimental M	edicine	)	
Principal Supervisor's Contact Details	Email: b.schock@qub.ac.u	lk	Tel: 07828065833	
Degree Pathway	Medical Science			
for which project	Biochemistry			
is suitable (√)	Microbiology			
	General awards		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:	OtherSystemic sclerosis (Ssc) is a chronic a multi-organ (systemic) disease characterized by autoimmunity, vascular lesions and progressive fibrosis that affects predominately the skin and the lungs. To date, the disease is associated with a high mortality and there is no approved therapy (1). In Ssc, fibroblasts are responsible for abnormal extracellular matrix accumulation and skin biopsies have been used for gene expression profiling. To mechanistically investigate pro-fibrotic phenotype conversion, cultured fibroblasts are used. An underlying factor of Ssc fibroblasts is their persistent pro-fibrotic activation which is, in part, driven by persistent activation of the TGF $\beta$ / WNT pathway after TLR4 stimulation (2). A20 is a potent regulator of fibrotic and inflammatory pathways and in scleroderma this regulation may be compromised leading to chronic pro-fibrotic and pro-inflammatory stimulation. Pharmacological induction of A20 has anti-inflammatory effects (3), but the degree of A20 induction depends on the expression of the A20 repressor DREAM (4). In normal fibroblasts, TGFß induced sustained downregulation of A20, and abrogated its TLR4-dependent induction, while siRNA-mediated knockdown of A20 enhanced the amplitude of fibrotic responses elicited by TGFB. Moreover, adiponectin induced A20 in fibroblasts and reduced fibrotic outcome measures (5). Moreover, findings from our pilot work using publicly available gene arrays (controls n=38, Ssc n=76) are highly promising: A20 mRNA expression in skin biopsies from patients with scleroderma was significantly reduced while the A20 repressor DREAM was significantly increased.			

Aims / objectives	<ul> <li>Here we wish to investigate the effect of TGFß stimulation of lung fibroblasts on the expression of A20, DREAM, p21 (cell proliferation) and collagen I and III expression.</li> <li>Furthermore, we wish to examine the effect of A20 inducing drugs (e.g. gibberellic acid, myricetin) on proliferation and collagen expression.</li> <li>References: <ul> <li>(1) Allanore Y <i>et al.</i> Nat Rev Dis Primers. 2015 Apr 23;1:15002; (2)</li> <li>Bhattacharyya S <i>et al.</i> Arthritis Research &amp; Therapy 2016;18:216;</li> <li>(3) Reihill JA <i>et al.</i> Br J Pharmacol. 2016 Feb;173(4):778-89;</li> <li>(4) Tiruppathi C <i>et al.</i> Nat Immunol. 2014 Mar;15(3):239-47;</li> <li>(5) Bhattacharyya S, Varga J. Curr Rheumatol Rep.2015Jan;17(1):474;</li> </ul> </li> <li>This project will characterise A20 and DREAM expression in cultured lung fibroblasts (a commonly used model for Ssc lung fibrosis) in response to TGFß. We hypothesis that augmentation of A20</li> <li>(decreasing DREAM) will reduce proliferation and collagen expression in cultured lung fibrosblasts.</li> </ul>
	Fibroblasts will be grown in submersion, stimulated (TGF $\beta$ ) and A20, the repressor DREAM (mRNA, protein), proliferation marker p21 and collagen I and III (mRNA) will be determined by qRT-PCR and Western Blotting.
Techniques employed:	Tissue culture and sterile working techniques (culture of human lung fibroblasts, stimulation with TGFß2 (10 ng/ml) in the presence and absence of the predicted drugs), collection of total mRNA, conversion into cDNA and quantitative real time PCR. Protein analyses by Western Blotting. Statistical analyses of results. Transferrable skills: Working in a team and alone, presentation of data and communication to other members the laboratory and the wider scientific community and the collaborators.

# **Centre for Medical Education**

Project Title	How do medical students	s learn	to be 'good' doctors?	
Supervisor(s)	1. Tim Dornan			
School / Centre	Centre for Medical Education			
Principal	Email: Tel: 07712 528565			
Supervisor's Contact Details	timothy.dornan@gmail.com			
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	This intercalation offers on	e or m	aximum two highly motivated stude	ents
information:	the opportunity to work as members of a small, collegial research group, specialising in education research. The distinguishing feature of our work is that it is as much a social science as a medical science. The main strand of our work is research into how medical students learn to practise medicine amidst the social complexity of workplaces. This offers benefit on several quarters. It helps candidates learn to be good doctors; it helps them learn to teach; and it teaches them ways of thinking that are not so strongly promoted by the mainstream medical curriculum. We 'tailor' projects to the wishes and needs of individual students.			to od
Aims / objectives	<ul> <li>Projects we can offer this coming year include: <ul> <li>How do children experience hospitals and how can they contribute to medical students' and doctors' learning?</li> <li>How do medical students learn to 'hold their own' in hospital settings, and prescribe safely there</li> <li>How can medical humanities contribute to medical education</li> </ul> </li> </ul>			
Techniques employed:	We have expertise in a range of methodologies, chiefly qualitative research. This means interviewing or conducting discussions with people in order to learn about social situations, like practising medicine or prescribing.			ine
	<b>.</b> .		b learn how to conduct a rigorous one well, lead to publications.	
	We can offer training in su many other research techr		search, implementation science, a	nd

Project Title	Interprofessional simul review	ation	based education: a scoping	J
Supervisor(s)	1. Dr Briegeen Girvi	n		
	2. Dr Gerry Gormley			
School / Centre	1. School of Pharmacy			
	2. Centre for Medical Education			
Principal	Email:		Tel:	
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Contact Details				
Degree Pathway	Medical Science			
for which	Biochemistry			
project is	Microbiology			
suitable (√)			Out is at an a if a surround.	
le project of	General awards		Subject-specific awards	
Is project of suitable			British Assoc	
standard /	Wolfson Foundation		Dermatologists	
subject for	Wonson Foundation		Dermatologists	
studentship	Jean Shanks		Digestive Disorders	
application? $(\checkmark)$	Foundation		Foundation	
			Pathological Society	
			Other	
Background	BACKGROUND			<u> </u>
information:	Interprofessional educa	ation		
	•		Ithcare provision are comple	x and
	increasingly revolve arou	und t	eams of professionals, rathe	r than
	relying on individual prac	tition	ers. Competent individuals m	ay not
			ams. Professional developme	
			ce is known to improve the qu	
			on interprofessional skills mu	ust be
	established at undergrad	uate I	evel.	
	Simulation based education	ation		
			SBE) has emerged as a sign	ificant
		•	can advance student learnin	
			ssionals for practice. The evi	
			enefits that SBE can bring to p	
	care. The simulated	expei	rience provides a realistic	and
			ty that can prepare healthcare	teams
	to perform successfully ir	n real	clinical settings.	
	The emerging roles of <b>p</b>	oharn	nacy	
			Council Standards on the	initial
			rmacists advise that the MPI	
			experience of working with pa	
	carers and other health	care	e professionals. This is cu	rrently

	achieved through a mix of off-site placement visits, using patients, carers and other health care professionals in-class, and simulations. Qualified pharmacists are increasingly being involved in more clinical roles which involve close interprofessional working both in secondary and primary care. Examples include managing and prescribing for patients with long term conditions, such as hypertension, heart failure, diabetes, asthma and COPD. The expanding roles for pharmacists require competence in various skills such as clinical skills (including physical assessment), critical thinking, communication and team work skills. Pre-2011, most of the published literature around use of simulation in the education of health care professionals has been in medical and nursing schools and less often in pharmacy. Preliminary evidence shows that interprofessional learning through simulation enables participants to practice teamwork and communication skills that are essential for preventing errors and patient harm (Crea, 2011). A review of the published literature on the effectiveness of simulation
	in pharmacy education and particularly where simulation has been used in interprofessional education, would be a huge benefit to universities when planning training to prepare their students for work in practice.
Aims / objectives	<ul> <li>AIM</li> <li>The overall of this project is to establish an understanding of the role of simulation based education in the role of pharmacist training and interprofessional education.</li> </ul>
	<ul> <li>OBJECTIVES</li> <li>Undertake a <i>scoping review</i> to map current evidence relevant to a SBE in pharmacy and interprofessional education</li> <li>identify all relevant publications, and draw whatever conclusions the evidence supports in the use of SBE in pharmacy</li> <li>consider how this information may have an impact on educational policy and practice</li> <li>Identify topics for future educational research and development.</li> </ul>
Techniques employed:	The successful applicant will use a <i>scoping review</i> methodology to review the literature. Such a form of literature review is exploratory in nature that aims to map current evidence relevant SBE in pharmacy The review will follow the methodological steps for scoping
	<ul> <li>reviews devised by Arksey and O'Malley (2005)</li> <li>Namely:</li> <li>Step 1: Identifying the research question</li> <li>Step 2: Finding relevant articles</li> <li>Step 3: Selection of relevant articles</li> <li>Step 4: Charting the data</li> <li>Step 5: Collating and summarizing the data</li> </ul>

Step 6: Consultation exercise (focus groups)
<ul> <li>The proposed benefits to the successful applicant <ol> <li>Generate and synthesize knowledge that could be used to influence practice and policy</li> <li>Develop skills in critical thinking, research methods, searching the evidence base, interview skills, presentation skills</li> <li>If would be the hope that this work will lead to a publication in a scientific journal and presentation at academic conferences</li> </ol></li></ul>
Arskey H and O'Malley L. Scoping Studies: Towards a Methodological Framework. Int J social Research Methodology 2015; 8 (1): 19-32
Crea KA. Patient simulation. Practice skill development through the use of human patient simulation. American Journal of Pharmaceutical Education 2011; 75 (9): Article 188.

Project Title	The role of conversation	during	g intravitreal eye injections.	
Supervisor(s)	1. Dr Michael William	าร		
	2. Dr Catrin Rhys			
School / Centre	1. MW: Centre for Medical Education, SMDBS, QUB			
	2. CR: School of Cor	2. CR: School of Communication and Media, UUJ		
Principal	Email: m.williams@qub.ac.uk Tel: 07540386492			
Supervisor's				
Contact Details Degree Pathway	Madical Science			
for which project	Medical Science Biochemistry	Х	-	
is suitable ( $\checkmark$ )	Microbiology		-	
	General awards		Subject-specific awards	
Is project of				
suitable standard	-		-	
/ subject for				
studentship				
application? ( $\checkmark$ )				
Background	Intravitraal ava injactiona (		re the most commonly performed	
information:		,		
information.			eloped world, usually performed for	
	•		n or diabetic macular oedema. Patients	
			s they are driven by fear of losing vision	
			ss experience variable amounts of	
	anxiety before and during	•		
	-		ection room varies: for some patients it	
	-	may reassure and relax them, while some <i>injectors</i> may believe that		
	conversation distracts or even may increase infection risk. This project			
	will use 'conversation analysis' (CA) to explore the role of utterances in the intravitreal injection room. CA is an established empirical method			
	-		I disciplines to analyse the sequence	
			nsights gained may be both therapeutic	
	and linguistic in nature. Dr	Williar	ns will be closely supervising this and	
	another related CA-based	interca	lated project, with Dr Rhys providing	
	specialist linguistics input.			
Aims / objectives	To explore the role of conv	/ersatio	on during intravitreal injections using	
_	conversation analysis			
Techniques	An ethics application is in	prepara	ation, and ethical committee approval	
employed:	will hopefully be in place b	y the s	tart of the project.	
	The project will then involv	e ident	ifying participants, seeking their	
	consent, setting up record	ing equ	ipment, transcribing the conversations	
			ts using 'Jeffersonian notation', and	
			to analyse and write up findings.	
			-	

Project Title	The role of conversation du	ring v	sual acuity measurement.	
Supervisor(s)	<ol> <li>Dr Michael Williams</li> <li>Dr Catrin Rhys</li> </ol>			
School / Centre	MW: Centre for Medical Education, SMDBS, QUB CR: School of Communication and Media, UUJ			
Principal Supervisor's Contact Details	Email: m.williams@qub.ac.uk		Tel: 07540386492	
Degree	Medical Science	х		
Pathway for	Biochemistry			
which project	Microbiology			
is suitable (✓)				
Is project of suitable standard / subject for	General awards		Subject-specific awards	
studentship application? (√)	-		-	
Background	Visual acuity (VA) measureme	ent is c	one in every ophthalmic consulta	tion,
information:			staff. The VA result is an import	
		-	naking, for example on whether t	
			al injections. Conditions in which	
			as possible, but there is still son	
			•	
		-	in a patient's VA from one visit to	
	'noise'.	change	in their condition, but merely be	
	Conversation between nurse	and pa	tient is an inherent part of VA	
			the task, and encourages the pa	atient
		•	No protocols exist for the nature	
	conversation in this setting.			-
	0	ation ar	nalysis' (CA) to explore the role o	f
			CA is an established empirical m	
	0		sciplines to analyse the sequence	
	-		gained may be both therapeutic a	
		-	e closely supervising this and an	
	0			
	linguistics input.	i projec	ct, with Dr Rhys providing special	ISI
Aims / objectives	To explore the role of convers conversation analysis	sation o	luring visual acuity measurement	using
Techniques			n, and ethical committee approva	al will
employed:	hopefully be in place by the st			
			ng participants, seeking their con	
			scribing the conversations ('the d	
			sonian notation', and then working	g with
	the supervisors to analyse an	d write	up findings.	
	l			

# **Centre for Public Health**

Project Title	Dementia data analytics in	Northe	ern Ireland		
Supervisor(s)	1. Dr Bernadette McGuinne				
	2. Prof Peter Passmore				
School / Centre	СРН				
Principal	Email:b.mcguinness@qub	.ac.uk	Tel:90978959		
Supervisor's	<b>.</b> .				
Contact Details					
Degree Pathway	Medical Science	х			
for which project	Biochemistry				
is suitable (√)	Microbiology				
	General awards		Subject-specific awards		
Is project of			Dritich Asses Derresstals sists		
suitable standard			British Assoc Dermatologists		
/ subject for	Wolfson Foundation		Digestive Disorders Foundation		
studentship	laan Ohanka Estati				
application? (✓)	Jean Shanks Foundation		Pathological Society		
			Other		
Background			hropies, OFMDFM and Departmer	nt of	
information:			itia analytics project in Northern		
			of data from the Data warehouse,	GP	
			er service. Outputs will include:		
			, political and administrative decision	on	
	makers; will inform policy a				
			of diagnosis, treatment and care		
	across the country and Europe				
	3 Use to further develop Dementia National Strategy 4 Generate new research hypotheses				
		5 Link with other European and worldwide registries			
		nai pee	er-reviewed high impact journals		
Aims / objectives	Several projects will be car	ried or	It by the team including the interca	lated	
			of anticholinergic drug use and	alou	
	mortality in patients with de				
			ia with a high anticholinergic drug		
			ity rate compared to patients with		
	dementia not on anticholin	ergic d	rugs.		
			of GP prescriptions of anticholiner		
			ompared to patients with dementia	not	
	<b>a a</b>	nd mort	ality rates in both over a five year		
	period.				
Techniques	Statistical analysis of large	datase	ets		
employed:					

Project Title		and r	enal outcomes in UK Biobank	
Supervisor(s)	1. Dr Gareth McKay			
	2. Dr Bernadette McGuinness			
	3. Mr Euan Paterson			
School / Centre	Centre for Public Health			
Principal	Email: g.j.mckay@qub.ac.	uk	Tel: 90978958	
Supervisor's				
Contact Details				
Degree Pathway	Medical Science	$\checkmark$	_	
for which project	Biochemistry		_	
is suitable (√)	Microbiology		Outring ( and a file and a star	
le project of	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		Dathalagiaal Saciaty	
			Pathological Society	
			Other	
Background	There is a naucity of ev	idenc	e relating to primary and seco	ndary
information:			irment. The retina is derived	
			to the brain via the optic i	
			•	
		sharing cellular similarities to the central nervous system. Retinal		
	blood vessels are amenable to direct non-invasive visualisation			
	and measurement allowing investigation of early microvascular			
	changes prior to the development of conditions of a vascular			
	nature. Renal dysfunction and vascular disease have been			
	reported in association	with c	ognitive outcomes in several	cross-
	sectional and longitudina	al stuc	lies. Retinal vascular paramete	ers will
	be evaluated against co	gnitiv	e and renal function measures	s from
			ired retinal measures will ir	
			calibre, tortuosity, and brar	
			confounding variables will also	•
			vill help determine whether	
			offer additional clinical utili	
	•		educed cognitive functioning	
	renal impairment is a po		•	
		cinid		
Aims / objectives	We are receiving data for	r 6998	3 participants from UKBB. This	study
			ata to evaluate associations be	
			and cognitive and renal fu	
	outcomes.	01013		

identify and measure retinal microvasculature changes from digital photographs acquired from the UK Biobank. The standardised measured area is defined within the region 0.5-2.0 disc diameter from the optic disc margin. Fractal analysis will quantify geometri branching complexity and density of retinal vessels providing a holistic overview of retinal microvascular health. Statistical analyses will evaluate associations between retinal microvascular variation and cognitive and renal function with consideration of potential confounding variables.
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Project Title	Dietary patterns and microvascular health – a study of renal dysfunction in the UK Biobank cohort			
Supervisor(s)	<ol> <li>Dr Gareth McKay</li> <li>Prof Jayne Woodside</li> <li>Mr Euan Paterson</li> <li>Dr Charlotte Neville</li> </ol>			
School / Centre				
Principal Supervisor's Contact Details	Email: g.j.mckay@qub.ac.	uk	Tel: 90978958	
Degree Pathway	Medical Science	$\checkmark$		
for which project	Biochemistry		_	
is suitable (√)	Microbiology			
Is project of suitable standard	General awards		Subject-specific awards British Assoc Dermatologists	
/ subject for studentship	Wolfson Foundation		Digestive Disorders Foundation	
application? ( $\checkmark$ )	Jean Shanks Foundation		Pathological Society	
			Other	
Background information:	The retinal vasculature is accessible to direct and repeated non- invasive assessment enabling detection of early microvascular changes prior to clinically significant events. A good diet is associated with reduced chronic disease risk, but the association between diet and retinal vascular health is underexplored. Clinical data derived from the UK Biobank will be used to identify individuals with renal impairment (urinary albumin/creatinine ratio (ACR) > 3mg/mmol) for comparison with control individuals with normal ACR. Comparisons will examine retinal vessel			
	measurements, including microvascular parameters such as calibre, tortuosity, and branching patterns. A wide range of confounding variables will be considered in the analysis. This studentship will analyse dietary data to examine food, whole			ge of whole
	with retinal vessel health	n and	d explore whether diet is asso renal impairment in this popula	ition.
Aims / objectives	study will analyse cros	s sec e asso	98 participants from the UKBB. tional data from a subset of ociations between dietary pat al function.	these

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 from the UK Biobank. 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. Statistical analyses will evaluate potential associations between dietary patterns, microvascular variation and renal function with consideration of potential confounding variables.
	consideration of potential confounding variables.

Project Title	Deep phenotyping and genetic analysis for Behçet's disease – a complex, multifactorial rare disease			
Supervisor(s)	AJ McKnight			
School / Centre	SMDBS – Centre for Public Health			
Principal Supervisor's Contact Details	Email: <u>a.j.mcknight@qub.</u>	<u>ac.uk</u>	Tel: 02890 638460 (shared line	э)
Degree Pathway	Medical Science	Yes		
for which project	Biochemistry	Yes	1	
is suitable (√)	Microbiology	No	1	
	General awards	110	Subject-specific awards	
Is project of suitable standard / subject for studentship application? (√)	Wolfson Foundation Jean Shanks Foundation	Yes Yes	British Assoc Dermatologists Digestive Disorders Foundation Pathological Society	Yes Don't know
			Other	
Background information:	≤5 per 10,000 people, wit and ~106,000 affected individually rare, but col public health problem. Th the impact of living with a Genetic and environment diagnosis is challenging w disease. A recent surve Northern Ireland reveale with <sup>1</sup> / <sub>20</sub> seeing >10 doct symptoms, which are var ups for this systemic inflammation / immune recurrent ulcers, skin lesion BD is most often reported prevalence is reported 1.5/100,000 individuals in practitioners revealed a 12.6/100,000 in the No expected 'UK' prevalence			

	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 interactions in Age-related macular Degeneration					
Supervisor(s)	Amy Jayne McKnight & 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 for which project is suitable ( $\checkmark$ )	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			
Background information:	Jean Shanks Foundation       Pathological Society         Other       Other         Genome-wide association studies have proved successful in revealing a significant proportion of the heritability related to AMD, with the most recent report highlighting 52 independently associated common and rare variants across 34 loci <sup>1</sup> , however missing heritability still remains. It is also well established that both genetic and environmental factors contribute to the development of AMD, but how these interact to result in the characteristic phenotypes in AMD is not well understood. There have been to date no large-scale population based studies involving cohorts well-phenotyped for AMD as well as characterised for the many demographic and environment risk factors known to be associated with the disease. The Northern Ireland Cohort for the Longitudinal Study of Aging provides such an opportunity as approximately 4,500 participants have underwent an extensive home interview, dietary assessment and health assessment which include multi-modal retinal imaging. The retinal images (colour, OCT, infra-red, autofluorescence and ultra-wide field Optomap images) have been graded for AMD including novel phenotypes such as reticular pseudodrusen/subretinal drusenoid deposits. This project would seek to relate genome-wide association data with environmental risk factors and presence of various AMD					
Aims / objectives	related phenotypes. To explore the relationship between genetic risk loci and environmental risk factors in AMD through the use of a genome-wide scan in a well- phenotyped population-based study in Northern Ireland (NICOLA Study).					

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

Project Title	Improving the food environment in primary schools					
Supervisor(s)	1. Professor Jayne Woodside 2. Dr Michelle McKinley					
School / Centre	Centre for Public Health					
Principal Supervisor's Contact Details	Email: j.woodside@qub.ac.uk		Tel: 02890978942			
Degree Pathway	Medical Science	х				
for which project	Biochemistry	х	1			
is suitable (√)	Microbiology		1			
	General awards		Subject-specific awards			
Is project of suitable standard / subject for studentship	Wolfson Foundation		British Assoc Dermatologists Digestive Disorders Foundation			
application? $(\checkmark)$	Jean Shanks Foundation		Pathological Society			
			Other			
Background information:	There is growing concern about diet quality in childhood and how poor nutrition may impair brain development and cognitive function, with low quality diets being associated with lower academic achievement. Children, particularly in urban settings, also often have little knowledge of where their food comes from. This project will be based in a wider body of work aiming to investigate the potential impact of engagement with the food sector, and alteration of the food environment in the primary school setting on diet, health and wellbeing outcomes.					
Aims / objectives			f changes in the school food Ith and wellbeing outcomes.			
Techniques employed:	<ol> <li>Systematic literatu</li> <li>Qualitative researd</li> </ol>					