



UNNC – SIMM, CAS Doctoral Training Partnership

Available PhD topics

PhD topic 1	Al Enabled Drug Discovery of Anti-cancer Drugs
SIMM Supervisor(s)	Prof Xiaojie Lu
UNNC Supervisor(s)	Dr Bencan Tang
Short introduction & description of PhD	Machine learning is an interdisciplinary study of statistical model that solves problems by making data-based predictions and deductions. The significance of machine learning and artificial intelligence used in drug discovery has been emphasized by recent papers published in high ranking journals. This research aims to develop advanced machine learning algorithms for the screening of anti-cancer agents. It involves bench chemistry as well as artificial intelligence. A candidate with chemistry or medical chemistry background with strong interest in machine learning would be ideal.
Contact points	Please contact Dr Bencan Tang (<u>Bencan.Tang@nottingham.edu.cn</u>) and Prof Xiaojie Lu (<u>xilu@simm.ac.cn</u>) before sending the application via <u>'How to apply'</u> section.
PhD topic 2	Application of Graphene in quality control and production of Liver Organoids
SIMM Supervisor	Prof Guoyu PAN
UNNC Supervisor(s)	Prof Cheng Heng PANG
Short introduction & description of PhD	Background: The in vitro model of hepatocytes is an essential tool for the evaluation of drug metabolism and drug toxicity. Hepatocytes are easy to lose their original traits, morphology, structure and function due to the limited conditions of in vitro hepatocyte culture. As a result, it cannot reflect the efficacy and toxicity of in vivo drugs, and does not fulfil the needs of clinical and scientific research.
	3D culture can better reflect the interaction between cells and matrix, especially the recent rise of organoid technology, which may mimic the in vivo environment and reflect physiological changes when a diverse population of cells is present. With the help of novel organoid technology, 3D culture can better reflect the interaction between cells and substrates, simulate the in vivo environment, and reflect physiological changes. However, how to maintain the integrity and physiological function of organoids in vitro has always been the focus of translational medicine and materials science. From the perspective of new drug development, the best choice for hepatotoxicity and liver drug screening is hepatic-like cell organoids, which is capable of long-term preservation of liver function. However, this in vitro model is far from mature to be employed by pharmaceutical industry.

To solve this problem, a 3D culture material matrix with better biocompatibility is required; on the other hand, a rapid, comprehensive, and non-invasive quality control method is a prerequisite for industrialization of liver organoid culture.

Possibilities analysis of Graphene application in liver organoid culture and quality control:

Graphene is a honeycomb carbon structural material formed by carbon atoms connected by Sp2 hybridization, and has a hexagonal honeycomb layered structure. The surface properties, low toxicity to cells, and good biocompatibility of Graphene make it useful as a platform for cell adhesion and even inducing cell proliferation. As a scaffold, Graphene can increase the quantity of adhesion points and cell proliferation while increasing intracellular tension. Compared to 2D Graphene film, the 3D Graphene scaffold not only provides a porous structure, but also possesses microscale topographic features and electrical conductivity. Graphene can be used as an excellent carrier material for co-culture of various cells. It has a unique function for the morphological retention of organoids. However, the relationship between liver organoids and Graphene has not been systematically studied.

Non-invasive continuous quality control of organoids is a prerequisite for large-scale production of organoids which cannot be achieved by current quality control methods. Due to the unique biochemical components of cells, tissues and organs, Raman spectroscopy is expected to be a powerful tool for organoid culture quality control. Raman spectroscopy is restricted in its application to larger organoids because of its sensitivity. Surface-enhanced Raman spectroscopy can be more effective and more sensitive than conventional Raman spectroscopy to help us distinguish between normal and diseased cells or tissues. Due to the two-dimensional ultrathin structure and high specific surface area of Graphene, Graphene has strong adsorption to molecules. These excellent properties make it possible to chemically enhance the response signal of Raman spectroscopy. It was found that the surface plasmon properties of silver nanoparticles (AgNPs) can also enhance the absorption range of Raman light. It was reported that the combination of silver nanoparticles with Graphene can simultaneously exert the electromagnetic enhancement of AgNPs and the chemical enhancement of Graphene, the Raman signal of the detected molecule can be enhanced by several orders of magnitude. This provides a unique possibility for Raman spectroscopy application in quality control of liver organoid cultures.

Brief Research Proposal:

This project aims to establish a novel liver organoid culture method with novel materials in scaffolds. With the help of 3D Graphene or related materials, it may improve the metabolic enzyme activities and transport protein abilities in hepatic-like cell organoids, which can meet the needs of clinical and scientific research purposes. Silver Nanoparticles (AgNPs) composited with Graphene and alternative compositions may generate a novel surface-enhanced Raman spectroscopy material with high sensitivity and good stability for quality control in the production of organoid chip. It can realize the detection of physiological and pathological states of liver organoids under different culture conditions. The work may accelerate the pharmaceutical research and environmental toxicological investigation.

Contact points

Informal inquiries may be addressed to Prof Guoyu Pan (gypan@simm.ac.cn) and Prof Cheng Heng Pang (ChengHeng.Pang@nottingham.edu.cn), but formal applications should follow the instructions in 'How to apply' section.

PhD topic 3

Design and application of graphene coupled antibody detection kit

SIMM Supervisor	Prof Chunhe Wang
UNNC Supervisor(s)	Prof Cheng Heng PANG
Short introduction & description of PhD	Graphene has very bright prospects in biomedicine and drug delivery. Graphene's unique two-dimensional structure makes it have bright application prospects in the field of sensing and detection. Its large surface area makes it very sensitive to the surrounding environment, which shows that graphene has high detection sensitivity.
	Antibody, also known as immunoglobulin, is a large Y-shaped protein mainly secreted by plasma cells and used by the immune system to identify and neutralize foreign substances such as bacteria, viruses and other pathogens. An antibody can uniquely recognize a specific foreign object through its variable region, which is called an antigen.
	When the antibody is coupled with graphene, the detection kit can be made according to the targeting of antibody and the ultra-high sensitivity of graphene, which has great prospect and research value.
Contact points	Informal inquiries may be addressed to Prof Cheng Heng PANG (ChengHeng.Pang@nottingham.edu.cn) and Prof Chunhe Wang (wangc@simm.ac.cn), but formal applications should follow the instructions in 'How to apply' section.
PhD topic 4	DNA Encoded Library(DEL) Enabled Drug Discovery of Anti-cancer Drugs
SIMM Supervisor(s)	Prof Xiaojie Lu
UNNC Supervisor(s)	<u>Dr Bencan Tang</u>
Short introduction & description of PhD	DNA encoded library (DEL) technology has become a useful tool for the hit identification for many biologically interesting targets for the drug discovery. Several hit compounds discovery by the DEL has been further optimized and move forward to the phase1-2 clinical stage. The success of the DEL was heavily relied on the chemical diversity of libraries, more DNA compatible reactions and biologically interested scaffolds are urgently needed to expand the DEL chemical space. This research aims to develop more DNA encoded libraries for the screening of anti-cancer agents. It involves bench synthetic chemistry as well as computational chemistry. A candidate with chemistry or medical chemistry background with strong interest in computational chemistry would be ideal.
Contact points	Please contact Dr Bencan Tang (<u>Bencan.Tang@nottingham.edu.cn</u>) and Prof Xiaojie Lu (<u>xjlu@simm.ac.cn</u>) before sending the application <i>via</i> ' <u>How to apply'</u> section.
PhD topic 5	Drug Discovery of Anti-cancer Drugs
SIMM Supervisor(s)	Prof Wenhu Duan
UNNC Supervisor(s)	<u>Dr Bencan Tang</u>
Short introduction & description of PhD	The project will focus on the discovery of anti-cancer candidates. It involves the use medicinal chemistry knowledge to guide the synthesis of new molecules with potential to be developed into anti-cancer drugs. It will also involve the design of molecules for specific targets and the synthesis of designed molecules. A good medicinal chemist with master's degree would be an ideal candidate for this project.
Contact points	Informal inquiries may be addressed to Dr Bencan Tang (Bencan.Tang@nottingham.edu.cn) and Prof Wenhu Duan (whduan@simm.ac.cn), but formal applications should follow the instructions in 'How to apply' section.
PhD topic 6	Drug Discovery of Anti-cancer Drugs

SIMM Supervisor(s)	PI in drug discovery in SIMM - TBC
UNNC Supervisor(s)	Dr Bencan Tang
Short introduction & description of PhD	This project will use medicinal chemistry to guide the synthesis of new molecules with potential to be developed into anti-cancer drugs. The project will focus on the discovery of anti-cancer candidates. The project may involve the application of machine learning, computational chemistry or PROTAC. It will also involve synthesis of predicted molecules for testing by collaborators in biology. A good medicinal chemist or a synthetic chemist would be an ideal candidate for this project.
Contact points	Please contact Dr Bencan Tang (<u>Bencan.Tang@nottingham.edu.cn</u>) before sending the application via ' <u>How to apply'</u> section.
PhD topic 7	Intelligent drug delivery system and cancer immunotherapy
SIMM Supervisor	<u>Prof Haijun Yu</u>
UNNC Supervisor(s)	<u>Dr Binjie HU</u>
Short introduction & description of PhD	Immunotherapy has emerged as a promising clinical modality for cancer therapy due to its ability to initiate an antitumor immune response. However, current immunotherapy is severely impaired by immunosuppression of host T-cell antitumor activity through the programmed cell death 1 ligand (PD-L1) and programmed cell death receptor 1 (PD-1) (PD-L1/PD-1) immune checkpoint. In this project, we will dedicate to developing novel drug delivery system for tumor-specific delivery of various immune modulators. The drug delivery systems will be designed for on-demand drug release or activation at the tumor site, thus achieve precise immunotherapy. For example, Boolean logic nanoparticles will be developed for tumor-targeted co-delivery of immune modulators (e.g., immune activator and immune inhibitor) and combination immunotherapy. A library of stimuliactivatable units will be fabricated yielding YES/AND logic outputs by adjusting the input combinations, including extracellular matrix metalloproteins 2/9, intracellular acidity, and reduction in the tumor microenvironment. The drug delivery systems will be administrated through systemic injection, oral This Ph.D. program is a multi-disciplinary, which covers broad fields of materials sciences, nanomedicine, pharmaceutics and oncology. The potential candidate should be self-motivated and have strong background in material sciences, pharmacological sciences or onco-immunology.
Contact points	Informal inquiries may be addressed to Dr Binjie HU (<u>Binjie.HU@nottingham.edu.cn</u>) and Prof Haijun Yu (<u>hjyu@simm.ac.cn</u>), but formal applications should follow the instructions in 'How to apply' section.
PhD topic 8	Mathematical/computational neuroscience-based medicinal chemistry study
SIAT Supervisor	Prof Bing Xiong
UNNC Supervisor(s)	Dr Mainul HAQUE
Short introduction & description of PhD	Mathematical modelling of medicinal chemistry is a relatively new area, which plays an important role in clinical/chemical investigations. Our goal in this project is to develop and analyse a mathematical models along with laboratory experiments for a project related to multiple targeting drug development of chronic pain. This project will combine both mathematical/computational modelling and clinical approaches to the problem discussed above. The mathematical modelling will be undertaken under the supervision of Professor Mainul Haque at the School of Mathematical Sciences and the clinical investigations will be performed under the supervision of Prof Bing Xiong at the SIMM. The latter will train the applicant to do the necessary clinical experiments and

Contact points	his laboratory has all the necessary facilities to perform the experimental part of the programme. One explicit aim of this project would be to train the applicant in some of the appropriate clinical laboratory techniques combining mathematical/computational/statistical modelling to address clinical or public health questions of interest. Informal inquiries may be addressed to Dr Mainul Haque (Mainul.haque@nottingham.edu.cn) and Prof Bing Xiong (bxiong@simm.ac.cn), but formal applications should follow the instructions in 'How to apply' section.
PhD topic 9	Mathematical/computational modelling neural network regulating sleep homeostasis
SIMM Supervisor	Prof Guangsen Shi
UNNC Supervisor(s)	<u>Dr Mainul Haque</u>
Short introduction & description of PhD	Sleep homeostasis is essential for human health. It refers to the concept how the neural network regulates sleep. It is generally accepted that sleep homeostasis is achieved by the orchestration among varieties of sleep- and wake-promoting neurons, which eventually form a network. Each knot of this network can be potentially affected or interrupted by lots of factors, such as drugs, mood, behaviour drive, genetic variants etc. Some pathological factors are too strong therefore totally disrupt the network. This is how the disease such as insomnia occur. Mathematical modelling of gene regulatory networks is a relatively new area, which plays an important role in systems-biology investigations. Our goal in this project is to develop and analyse a mathematical models along with laboratory experiments for a specific gene regulatory the neural network regulates sleep. This project will combine both mathematical/computational modelling and laboratory approaches to the problem discussed above. The mathematical modelling will be undertaken under the supervision of Professor Mainul Haque at the School of Mathematical Sciences and the experimental investigations will be performed in the laboratory of professor Guangsen at SIMM. The latter will train the applicant to do the necessary experiments and his laboratory has all the necessary facilities to perform the experimental part of the programme. One explicit aim of this project would be to train the applicant in some of the appropriate laboratory techniques for measuring the effects of chemical and genetic risk factors on the sleep- and wake-promoting neurons network.
Contact points	Informal inquiries may be addressed to Dr Mainul Haque (Mainul.Haque@nottingham.edu.cn) and Prof Guangsen Shi (shiguangsen@zidd.ac.cn), but formal applications should follow the instructions in 'How to apply' section.
PhD topic 10	New drug development targeting brain tissue against stroke
SIMM Supervisor	Prof Baohong Jiang
UNNC Supervisor(s)	<u>Dr Binjie Hu</u>
Short introduction & description of PhD	Stroke continues to be the major global health problem associated with considerable mortality and morbidity. Developing new strategies to alleviate the lesion of stroke is urgent need. Our research will concentrate on new drug development against stroke. This research involves set up of animal model, efficiency evaluation based on infarct volume, neurobehavioral scores and histopathological examination. The new administration methods and drug delivery systems will be explored to penetrate blood brain barrier in order to target brain tissue for therapy of stroke.

quiries may be addressed to Prof Baohong Jiang (jiangbh@simm.ac.cn) and a (Binjie.HU@nottingham.edu.cn), but formal applications should follow the s in 'How to apply' section. If nano-particles and space distribution in cells and tissues of pry bowel disease visualized by synchrotron radiation micro-tomography and fluorescence micro-optical sectioning tomography Zhang Heng PANG Ory bowel disease (IBD) is a chronic and recurring long-term gastrointestinal lers, which causes severe destruction of the intestinal epithelial barrier. IBD ecognized as a global disease, accompanied by rising high incidence.
The property bowel disease visualized by synchrotron radiation microtomography and fluorescence micro-optical sectioning tomography Zhang Heng PANG Dry bowel disease (IBD) is a chronic and recurring long-term gastrointestinal lers, which causes severe destruction of the intestinal epithelial barrier. IBD
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colitis (UC) is one of the most common clinical forms of IBD which involves diarrhea, hematochezia and weight loss affecting life quality of patients the conventional treatment of IBD relies on large doses of oppressive and anti-inflammatory drugs, which cause serious adverse effects in specificity. To overcome these glitches, many researchers have designed es of colon-targeted preparations to improve drug accumulation at the Furthermore, there is a growing demand on understanding the thorough tectures, as well as, the accurate distribution patterns in vivo and retention as of the colon-targeted materials at three-dimensional (3D), in situ, and colo clevel. Thus, a novel strategy that reveals the spatial distribution of eted particles precisely at multi-scale ranging from sub-micron to the whole needs to be established.
o date, the cross-scale study of precise colon structure and its inner inhaled stribution still faces considerable obstacles. Although the in vivo imaging S) has been utilized as a routine method to provide the distributing follon-targeted particles at organ-scale, it only can ascertain the particles at at 2D level. Therefore, there still remains a lack of detection on the vatial distribution of colon-targeted particles at the whole-colon scale with resolution. If of PhD, high-precision cross-scale visualization of entire colon anatomy ed by the advanced Micro-Optical Sectioning Tomography (MOST) system the whole colon nissl-staining. In addition, suitable fluorescent dye labelled eted material were synthesized and characterized as model particles, and and colon-wide datasets were acquired by fluorescence-micro-optical tomography (f-MOST) system. More importantly, the sophisticated es of the mouse colon and the region-specific distribution patterns and associated deposition of the colon-targeted particles in colon were revealed usly for the first time, providing new approaches for understanding the ehavior and distribution of colon-targeted materials in vivo. This research is precise anatomical data for in vitro simulations and contribute to fill the en in vitro and in vivo analyzing results, which will facilitate the clinical
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Contact points	Informal inquiries may be addressed to Prof Jiwen ZHANG (jwzhang@simm.ac.cn) and Prof Cheng Heng PANG (chengheng.pang@nottingham.edu.cn), but formal applications should follow the instructions in 4How to apply section.
PhD topic 12	The morphology effect of Nano- and microspheres encapsulations on inhaled drug delivery and the application for lung disease therapy
SIMM Supervisor	Prof Yongzhuo Huang
UNNC Supervisor(s)	<u>Dr Binjie Hu</u>
Short introduction & description of PhD	Inhaled formulations provide a self-administered and safe method for lung disease therapy, with an advantage of local drug delivery that can directly target the organ and thus reduce the unwanted drug exposure to healthy tissues. Nano- or microspheres encapsulations can be used as inhaled delivery carriers for achieving drug extended release and protection for degradation. Their morphology plays an important role in determining the in vivo fate of the drugs. For example, the different size and shape could result in various biodistribution in the target organ. Furthermore, Surface modification of the carriers can render the functional alteration.
	In this project, the morphology effect of nano- and microspheres on the biodistribution and therapeutic efficacy will be explored, and subsequently, the morphology will be modified by a physical or chemical method to achieved an optimal treatment efficacy in various diseases (e.g., acute respiratory distress syndrome, pulmonary fibrosis, and chronic obstructive pulmonary disease). And inhaled vaccine delivery using of nano- and microspheres encapsulations will also be explored.
Contact points	Informal inquiries may be addressed to Dr Binjie Hu (binjie.hu@nottingham.edu.cn) and Prof Yongzhuo Huang (yzhuang@simm.ac.cn), but formal applications should follow the instructions in 4How to apply ' section.