RCNMH25Br Mastitskaya ow star cells of the brain help save the heart
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euroscience & Mental Health
lucagon-like peptide-1 is a hormone and neurotransmitter peptide nown for its role in appetite control and glucose metabolism. Other ealth benefiting effects of GLP-1, in particular prevention of damage om heart attack, are less understood. This project will study how GLP-1 ctivates neural connections in the brain to send protective stimuli to ne heart in an experimental rodent model of heart attack. The emerging alle of brain cells called astrocytes in modulation of these connections ill be clarified using state-of-the-art techniques of gene activation and lencing in vivo.
the GLP-1 analogue drugs took the market by storm, promising magic are for obesity. The main physiological role of endogenous gut-derived LP-1 is the stimulation of insulin release from the pancreas in response of food intake. There is also brain-derived GLP-1 that targets neural rcuits regulating satiety. Centrally produced GLP-1 and pharmacological LP-1 receptor agonists have potent effects on the processing of utonomic reflexes in the brainstem.
The have previously demonstrated that GLP-1 acts as a molecular factor is vagally mediated cardioprotection (PMID: 27702763). The vagus nerve critically important in cardiovascular health. Vagus nerve stimulation duces myocardial infarction (PMID: 22739118, PMID: 26918777), revents the progression of heart failure and improves exercise capacity MID: 32875170). It is difficult to engage vagus nerve activity in patients advanced age or in diabetes due to impairment of neural transmission the heart. Therefore, targeting the downstream molecular events narmacologically is the way to overcome these limitations. GLP-1 ceptor agonists, such as Ozempic, have proven to be incredibly ficient in the treatment of diabetes, reducing body weight and inimizing the other consequences of cardiometabolic disease. owever, the exact location of the GLP-1 receptor and the molecular athway(s) responsible for GLP-1's cardioprotective effects are not fully nderstood. We have preliminary data showing that GLP-1 protects the eart via modulation of autonomic reflexes at the level of the dorsal agal complex (DVC) in the brainstem. There is also substantial coperimental evidence suggesting a crucial role of brainstem astrocytes the processing of cardiovascular reflexes (PMID: 32132265) and in the gulation of GLP-1R mediated effects on energy balance (PMID: 7013681).

	The research aims of the project will include the following: 1) To characterise the cardioprotective effects and dose-response relationship of GLP-1R agonists administered systemically (i.v.) vs centrally (intracerebroventricular, i.c.v.) in an in vivo model of myocardial ischaemia/reperfusion injury. Pharmacological experiments will explore if CNS administered GLP-1R antagonists block cardioprotection evoked by systemically administered GLP-1R agonists and vice versa. 2) To determine the role of centrally produced endogenous GLP-1 in cardioprotection. The PPG neurones of the mouse brainstem will be ablated using Cre-dependent expression of diphtheria toxin fragment A (PMID: 30279161). 3) To study the effects of GLP-1R agonists and antagonists on neuroglial communication in brainstem in the presence of various neuro- and gliotransmitters in electrophysiology and Ca imaging experiments. 4) The role of brainstem astrocytes in the cardiovascular effects of GLP- 1R activation in vivo will be studied in experiments on their pharmacological inhibition (PMID: 27013681) or chemogenetic activation (PMID: 37370201). The student will receive comprehensive training in cardiovascular physiology and autonomic neuroscience, will become skilled in in vivo research and will obtain a Home Office animal licence. To support the development of data analysis skills they will enroll in coding courses in the first year of their PhD (R, Python, MatLab). The student will be encouraged to regularly engage in group meetings and research seminars both in Bristol and Exeter and will have the opportunity to work in both laboratories, in accordance with research needs, to benefit from the training and collaboration potential of the supervisory team. Informed by the outcome of each of the main research aims, the student can choose which aspects of GLP-1R mediated cardioprotection to explore. For example, if peripherally administered GLP-1R activation. This project will disentangle the cardiovascular effects of ce
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