## 科研伦理委员会伦理审查决议 Approval Letter of Research Ethics Committee

项目名称	LncRNA TNFRSF10A-AS1 在结直肠癌中的作用及机制		
Study Title	研究		
项目来源	and the state of		
Issued BY	研究生课题 		
主要研究者	学生: 王丹丹	科室	消化内科
Principal investigator	导师: 张晓岚	Department	
审查类别	初始审查	审查方式	快速审查
Category of Review		Type of Review	
送审日期	2021.3.1	审查决议编号	2021-R241
Date Submitted		Approval Letter No.	
会议地点		会议时间	
Meeting Location		Meeting Date	
送审资料	研究方案(版本号: 1.0, 版本日期: 2021.3.1) 知情同意书(版本号: 1.0, 版本日期: 2021.3.1) 主要研究者简历		
Document(s)			
Reviewed			
ACTIONOU			
投票结果 Comments	伦理委员会按照国家相关法规要求对送审材料进行		
	审阅和讨论,出席会议 $0$ 人,参与投票(Vote) $0$ 人。		
	投票结果:同意(Approved)_0_票;作必要的修正		
	后同意(Conditional approved) _0 票;作必要修正后重		
	审(Reviewd after revising) 0 票;终止或暂停已批准的		
	研究(Termination or suspension of approved studies) <u>0</u>		
	票;不同意(Rejected)_0_票。		
	根据国家卫健委《涉及人的生命科学和医学研究伦		
审查意见和结论	理审查办法》、《赫尔辛基宣言》和《涉及人的健康相		
	关研究国际伦理指南》的伦理原则, 经本伦理委员会审		
	查,同意(Approved)按所批准的临床研究方案、知情		
Review opinions and	同意书开展本项研究。		
1	conclusions 主审委员(签字)   主任委员(签字) 其序   日期: 2021年3月22日		
Conclusions			

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## 河北医科大学第二医院科研伦理委员会 Research Ethics Committee of the second hospital of Hebei Medical University

请遵循 GCP 原则、遵循伦理委员会批准的方案开展临床研究,保护受试者的健康与权利。研究过程中若变更主要研究者,对临床研究方案、知情同意书、招募材料等的任何修改,请申请人提交修正案审查申请。发生严重不良事件,及时提交严重不良事件报告。发生违反试验方案情况须及时报告本伦理委员会。

In order to protect the health and rights of the subjects, the applicant should follow the GCP principles and programs approved by the Ethical Committee during the clinical studies. If any changes are made to the primary investigator(s), the clinical research protocol, informed consent, recruitment materials, etc., the applicant is requested to submit an amendment for review. In the event of a serious adverse event, a serious adverse event report should be submitted in time. The violation of the protocol shall be promptly reported to the Ethics Committee.

请按照伦理委员会规定的年度/定期跟踪审查频率,在截止日期前1个月提交研究进展报告;申请人暂停或提前终止临床研究,请及时提交暂停/终止研究报告。完成临床研究,请申请人提交结题报告。

The applicant should submit the study progress report one month before the deadline according to the frequency of the annual / periodic follow-up examination. If the applicant suspends or prematurely terminates the clinical research, the suspension / termination report should be submitted in time. The applicant should submit the final report when they complete the clinical study.

研究纳入了不符合纳入标准或符合排除标准的受试者,符合中止试验规定而未让受试者退出研究,给予错误治疗或剂量,给予方案禁止的合并用药等没有遵从方案开展研究的情况;或可能对受试者的权益/健康以及研究的科学性造成不良影响等违背 GCP 原则的情况,请申办者/监察员/研究者提交违背方案报告。

If the study is not carried out in accordance with the scheme, such as the study included subjects who did not meet the inclusion criteria or met the exclusion criteria, subjects were not withdrawn from the study in accordance with the termination rule, the wrong treatment or dosage was given, the combination drugs prohibited by the scheme were given, or the rights / health of the subjects and the scientific nature of the study may be adversely affected, which violated the GCP principle, the sponsor / inspector / researcher is requested to submit the violation report.

本试验年度/定期跟踪审查频率为一年,本批件有效期为一年。

The annual / periodic follow-up examination frequency is one year and the validity of this document is one year.