Ethical approvals of all participating hospitals

This trial is to be conducted in 12 hospitals. The ethical review was firstly submitted to IRB of the principle organization, Guang'an men Hospital, and then to IRB of other participating hospitals. This trial has gained approval from all of the IRBs.

Ethical approvals are attached in the following sequence.

- 1. Guang'an men Hospital, China Academy of Chinese Medical Sciences
- 2. Xiyuan Hospital of China Academy of Chinese Medical Sciences
- 3. Dongzhimen Hospital affiliated to Beijing University of Chinese Medicine
- 4. Yueyang Hospital of Integrated Traditional Chinese and Western Medicine affiliated to Shanghai University of Traditional Chinese Medicine
- 5. West China Hospital of Sichuan University
- 6. First Teaching Hospital of Tianjin University of TCM
- 7. First Hospital of Hunan University of Chinese Medicine
- 8. Hengyang Hospital affiliated to Hunan University of Chinese Medicine
- 9. Hubei Provincial Hospital of TCM
- 10. Jiangsu Province Hospital of TCM
- 11. Shanxi Province Hospital of TCM
- 12. Shanxi Hospital of Integrated Traditional and Western Medicine



中国中医科学院广安门医院伦理委员会文件(EC_AF_022)

伦理审查批件

项目名称	电针治疗女性单纯性压力	力性尿失禁有效性	和安全性多中心随机对照试验	
项目编号	2012EC007	项目来源	"十二五"国家科技支撑计划	
牵头单位	中国中医科学院广安门医院			
申办者(如有)	/			
主要研究者	刘志顺			
审查类别	初始审查	审查方式	会议审查	
审查日期	2012.12.14	审查地点	广安门医院行政楼四楼会议室	
审查委员	殷海波,朴炳奎,林兰, 贞,沈瑞英	谢利民,曹炜,	冯玲,赵军,胡镜清,吴萍,顾[·	
 批准文件	研究方案(VERSION 1.0_	20121105),知情	同意书(V1.0)	
审查意见	食品物的 《药物的 《药物的 《药物的 《药物的 《克斯·斯里斯斯里斯斯里斯斯里斯斯里斯斯里斯斯里斯斯里斯斯里斯斯里斯斯斯里斯斯里斯斯	临查以究竟 主请件事规与现伦纳究究背 床床工《阿宾伦 要人以件定申任理入,的 G 研	股受益比的非预期不良事件,请 限踪审查频率,申请人在截止日期 1长单位伦理委员会提交各中心研 响试验进行或增加受试者危险的 面报告。 除标准的受试者,符合中止试验 或剂量,给予方案禁止的合并用 对受试者的权益/健康以及研究的 记,请申办者/监查员/研究者提交	
有效期	2012年12月20日~20)13年12月19日		
联系人与联系电话	乔洁 010-88001552			
主任委员签字	Trains		· 英院广英//	
		中国中医科学	隐白安门医院伦理委员会(盖章	
		日期:	2012年12月21日5	
			一种人理学	

Institutional Review Board Documentation of Guang'anmen Hospital of China Academy of Chinese Medical Sciences (EC_AF_022)

Ethics Approval of Guang'anmen Hospital of China Academy of Chinese Medical Sciences

Project Title	The Effect and Safety of Electro-acupuncture for Women with Pure Stress Urinary			
	Incontinence: a Multicenter, Randomized Controlled Trial			
Approval No.	2012EC007	Project Sponsor	the 12 th Five-year Plan of the National Key	
			Technology Support Program by the	
			Ministry of Science and Technology of the	
			People's Republic of China	
Leading Organization	Guang'anmen Hospital of	f China Academy of	Chinese Medical Sciences	
Applicant (if any)	/			
Site PI	Zhishun Liu			
Review Attribute	Initial Review	Review Methods	Review Conference	
Review Date	Sep 14, 2012	Review Place	Conference Room, the 4 th Floor of the	
			Administrative Building of Guang'anmen	
			Hospital	
Review Committee	Haibo Yin, Bingkui Piao, Lan Lin, Limin Xie, Wei Cao, Ling Feng, Jun Zhao, Jingqing Hu,			
	Ping Wu, Lizhen Gu, Ruiying Shen			
Approved Files	Study Protocol (VERSION 1.0_20121105), Informed Consent (V1.0)			
Review Comments	According to "ethical review methods for biomedical study involving human subjects"			
	issued by the Ministry of Health, "Good Clinical Practice", "Provisions for Clinical Trials of			
	Medical Device" and "Guidelines for Ethical Review Work of Drug Clinical Trials" issued			
	by State Food and Drug Administration (SFDA) of the People's Republic of China,			
	"management specifications for ethical review of TCM clinical studies" issued by State			
	Administration of Traditional Chinese Medicine, "Declaration of Helsinki", and			
	"International ethical guidelines for biomedical research involving human subjects" issued			
	by Council for International Organizations of Medical Sciences, this clinical research was			
	reviewed by the institut	tional review board	(IRB) of Guang'anmen Hospital of China	
	Academy of Chinese Me	dical Sciences. And	the study protocol, informed consent, and the	

Institutional Review Board Documentation of Guang'anmen Hospital of China Academy of Chinese Medical Sciences (EC_AF_022)

	Data: Dec 21, 2012
	IRB of Guang'anmen Hospital of China Academy of Chinese Medical Sciences (Seal)
Director Signature	Haibo Yin
Contact	Jie Qiao, +86 010 88001552
Validity Period	From Dec 20, 2012 to Dec 19, 2013
	prematurely.
	A final report should be submitted when the study is finished completely or terminated
	badly affected; the science of study was badly affected.
	medicine was used; 2) conditions that violate GCP principle: subjects' rights and health are
	the rules of withdrawal; incorrect treatment or dose was given; prohibited combined
	wrongly included in the study; subjects do not withdraw from the study when he/she meet
	the inclusion criteria, or should be excluded according to the exclusion criteria, were
	the following occurs: 1) conditions that violate the study protocol: subjects who did not meet
	A protocol deviation report should be submitted by the site PI/monitor/researcher if any of
	report should be submitted by the site PI to the IRB.
	greatly affect the progress of the study or increase the potential risk of the subjects, a written
	should be submitted by the site PI to the IRB of the leading site. In any condition which will
	according to ethical review frequency. A summary report of the study progress of each site
	Researchers should submit report of the study progress one month before the deadline
	other un-anticipated AE, which will affect the risk-reward ratio of this study, occurs.
	A report of the severe adverse events (SAE) should be submitted in time if any SAE or any
	modification of the study protocol, informed consent, or the recruitment files are made.
	An application should be submitted if a change of the principle investigator (PI), or any
	the whole study.
	approved by the IRB. The health and rights of the subjects should be protected throughout
	Please conduct this clinical study following the GCP principles and the study protocol
	recruitment files of this research were approved.

中国中医科学院西苑医院医学伦理委员会审查批件

批件号:中国中医科学院西苑医院医学伦理委员会 2013XL001-2

审查日期	2013年1月25日
审查地点	北京海淀区西苑操场1号中国中医科学院西苑医院医学伦理委员会
课题编号	2012BAI24B01
项目名称	电针治疗女性单纯性压力性尿失禁有效性和安全性 ——多中心随机对照试验
	伦理审查申请书、研究者手册、研究者履历、CRF、患者日记等;
审查文件	研究方案: 版本号: VERSION 1.0_20121106, 版本日期: 2012年11月6日
	修正的知情同意书:版本号:VERSION 1.0 20120220,版本日期: 2013 年2 月20 日
课题组织单位	国家科技部"十二五"国家科技部
临床研究单位	中国中医科学院广安门医院、北京中医药大学东直门医院、 四川大学华西医院、中国中医科学院西苑医院、湖南中医药大学附属衡阳医院、湖南省中医院、上海中医药大学附属岳阳中西医结合医院、天津中医药大学第一附属医院、陕西省中医医院、江苏省中医院、山西中医学院中西医结合医院、湖北省中医院
主要研究者	陆永辉(副主任医师) -本中心
会议审查委员	曹云、衷敬柏、尚晓泓、尹秀云、张广生、杨志旭、于振宣、韩梅、李涛、 房定亚、闫小平
审查意见	根据中华人民共和国国家药品食品监督管理局 2003 年颁布实施的《药物临泉试验质量管理规范》、2010 年 11 月颁布的《药物临床试验伦理审查工作指导原则》以及《赫尔辛基宣言》的伦理原则,经本伦理委员会会议审查,审查结果为"作业要的修正后同意",具体意见如下: (1) 主要研究者符合国家相关规定; (2) 研究方案的设计基本符合科学性、伦理性原则。临床筛选患者做残余易检查时不应让患者大量饮水。 (3) 修正后的知情同意书语言通俗易懂,信息充分。 (4) 给予伦理审查批件。如临床试验方案有任何修改,主要研究者更换等,需重新审查,获得批准后执行。 暂停/提前终止/完成临床研究,请及时通知伦理委员会。 本批件有效期一年,请于 2014 年 2 月 21 日前 1 个月提交跟踪审查申请报告。 发现影响受试者参加研究意愿的违反方案情况应及时报告
联系电话	伦理委员会秘书 管明杰 (010) 62835646
主任委员	曹云图一周
盖章	中国中医科学院西苑医院医学伦理委员会
日 期	2013 年 2 月 22 日

Ethics Approval of Xiyuan Hospital of China Academy of Chinese Medical Sciences

Approval Number: Institutional Review Board of Xiyuan Hospital of China Academy of Chinese Medical Sciences 2013XL001-2

Review Date	Jan 25, 2013
Review Place	
Review Place	Institutional Review Board of Xuyuan Hospital of China Academy of Chinese
	Medical Sciences, 1 Xiyuan Playground, Haidian, Beijing
Project No.	2012BAI24B01
Project Title	The Effect and Safety of Electro-acupuncture for Women with Pure Stress Urinary
	Incontinence: a Multicenter, Randomized Controlled Trial
Approval Files	Application Form of Ethical Review, Researchers' Handbook, Researchers' CV,
	CRF, Patients' Diary, etc.
	Study Protocol: Version No. VERSION 1.0_20121106; Version Date Nov 6, 2012
	Revised Informed Consent: Version No. VERSION 1.0 20120220; Version Date Feb
	20, 2013
Project Sponsor	the 12 th Five-year Plan of the National Key Technology Support Program by the
	Ministry of Science and Technology of the People's Republic of China
Clinical Sites	Guang'anmen Hospital of China Academy of Chinese Medical Sciences,
	Dongzhimen Hospital of Beijing University of Chinese Medicine, West China
	Hospital of Sichuan University, Xiyuan Hospital of China Academy of Chinese
	Medical Sciences, Hengyang Hospital of Hunan University of Chinese Medicine, the
	Second Affiliated Hospital of Hunan University of Chinese Medicine, Yueyang
	Hospital of Integrated Traditional Chinese and Western Medicine of Shanghai
	University of Traditional Chinese Medicine, the First Affiliated Hospital of Tianjin
	University of Chinese Medicine, Shaanxi Province Hospital of TCM, Jiangsu
	Province Hospital of TCM, Shanxi Hospital of Integrated Chinese and Western
	Medicine, Hubei Province Hospital of TCM
Site PI	Yonghui Lu (our center)
Review Committee	Yun Cao, Jingbo Zhong, Xiaohong Shang, Xiuyun Yin, Guangsheng Zhang, Zhixu
Review Committee	Yang, Zhenxuan Yu, Mei Han, Tao Li, Dingya Fang, Xiaoping Yan
Review Comments	According to the "Good Clinical Practice" issued by the State Food and Drug
Keview Comments	Administration of the People's Republic of China in 2003, the "Guidelines for
	Ethical Review Work of Drug Clinical Trials" issued in Nov, 2010, and the
	"Declaration of Helsinki", though our Review Conference, the results of this review
	was "agreed after revision", detailed comments are as following:
	(1) the site PI selected meets the correlated national regulation;
	(2) the design of this study protocol was scientific and ethical; For doing the residual
	urine volume, patients should not be required to drink water generously;
	(3) the revised informed consent was straightaway and informative;

	(4) the documentations were approved by the IRB, if any revision of the study		
	protocol, or a change of the main researcher was made, a new ethics review will be		
	needed.		
	Please inform the IRB in time if the study was paused/stopped		
	prematurely/completed.		
	This approval stands good for 1 year, please submit an application of continuing		
	review 1 month before Feb 21, 2014.		
	Please report in time if the condition that violate the study protocol occurs.		
Contact Phone	Secretary of the IRB: Mingjie Zi +86 010 62835646		
Committee Director	Yun Cao		
Seal	Institutional Review Board of Xiyuan Hospital of China Academy of Chinese		
	Medical Sciences		
Date	Feb 22, 2013		

北京中医药大学尔真门医院医学伦理委员会 IRB of Dongzhimen Hospital affiliated to Beijing University of Chinese Medicine

化连甲耳弧件

Appre	oval Noti	ce Template			
受理序号: ECSL-BDY-2013-04		-	批件号: EC	PJ-BDY-201	3-04
项目名称: 电针治疗女性单纯性压力性质	永失禁 有	可效性和安全	性多中心的	通机对照试	&
申办单位: 东直门医院			主要研究者	· 赵吉平	
项目类别: 国家科技支撑项目		批文号/	课题编号:	2012BAI24	301
方案版本号: 1.0_20121105		方案批准日	期: 201	2. 11. 5	
知情同意书版本号: V1.0	知情	司意书批准日	期: 2012	. 11. 5	
伦理审查方式: 〇会议审查	口快速	书查			
应到会 J 5 人,出席本次会议人员 C	7 人,	回避 0	人,缺席	€ A	
根据中华人民共和国国家食品药品监督管理局(SI物临床试验伦理审查工作指导原则)(2010年)、(口同意临床研	开究方案		
试验质量管理规范》(2003)、(中药品种保护指 (2009),世界医学会(赫尔辛基宣言)(2008),卫		口不同意临時	未研究方案		<i>:</i>
及人的生物医学研究伦理审查办法》(2007). 国家 理局《中医药临床研究伦理审查管理规范》(2010		口终止临床	研究方案		
际医学科学组织委员会(人体生物医学研究国际) (2002) 的伦理原则 经本伦理委员会审查决定	(德指南)	□暂停临床码	研究方案	-	. •

同意临床研究

审査意见:

注:本批件自签发目期有效期一年,研究负责人必须严格使用经审查同意的知情同意书文本和研究方案。如伦理审查批件失效时不能完成所有的临床研究(包括统计分析),请在本批件失效前一个月,递交持续审查申请。如研究结束并在审查有效期内,请递交研究结题报告。研究中发生涉及受试者或其他人风险的任何预期或非预期的不良事件,应立刻报告本伦理委员会,任何研究方案、知情同意书的修改包括研究人员得变更,必须递交研究方案修改申请表,经伦理委员会审查获得批准后执行。

北京中医药大学东直门医院医学伦理委员会 IRB of Dongzhimen Hospital affiliated to Beijing University of Chinese Medicine

会议签到表

Meeting attendance sheet

项目名称	电针治疗女性单纯性压力性尿失禁有效性和安全性多中心随机 对照试验		
会议时间	2月1日	会议地点	第一会议室

成员	性别		专业	签名
李澎涛	男	主任	 脑病	
子伊彻	77	.T.].L	DE17P3	
高颖	女	副主任	脑病	2003-
叶永安	男	副主任	消化	
柳红芳	女	副主任	肾病内分泌	1 home
张永涛	男	委员	呼吸	2 237]
王新月	女	委员	消化	
杨博华	男	委员	周围血管	,
鲁卫星	男	委员	心血管	En Z
王蓬文	女	委员	药理学	200
曹俊岭,	男	委员	药剂学	Mason de
刘凯	男	委员	法律代表	
贺海东	男	委员	医疗器械	Ryz
张胜利	男	委员	群众代表	3 Pau \$ 系)
陈信义	男	委员	血液肿瘤	1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1
彭淑莲	女	委员	乳腺外科	洲人

Institutional Review Board of Dongzhimen Hospital Affiliated to Beijing University of Chinese Medicine

Approval Notice Template

Accepted No. ECSL-BDY-2013-04 Approval No. ECPJ-BDY-2013-04

Project Title: The Effect and Safety of Electro-acupuncture f	for Women with Pure Stress Urinary	
Incontinence: a Multicenter, Randomized Controlled Trial		
Application Center: Dongzhimen Hospital	Site PI: Jiping Zhao	
Project Attribute: the National Key Technology Support Pro	ogram Project No. 2012BAI24B01	
Protocol Version No. 1.0_20121105	Protocol Approval Date: Nov 5, 2012	
Informed Consent No. V1.0	nformed Consent Approval Date: Nov 5, 2012	
Review Method: √ Review Conference	Quick Review	
Member: Anticipated 15 persons, participated 9 persons,	avoided 0 persons, absented 6 persons.	
According to the "Guidelines for Ethical Review Work of Drug		
Clinical Trials" (2010), the "Good Clinical Practice" (2003), the	$\sqrt{approved}$	
"Guiding Principle of Herb Variety Protection" (2009) issued by		
State Food and Drug Administration (SFDA) of the People's		
Republic of China, "Declaration of Helsinki" (2008), the "ethical	not approved	
review methods for biomedical study involving human subjects"		
(2007) issued by the Ministry of Health, the "Management		
specifications for ethical review of TCM clinical studies" (2010)	terminated	
issued by State Administration of Traditional Chinese Medicine,		
and the "International ethical guidelines for biomedical research	paused	
involving human subjects" (2002) issued by Council for		
International Organizations of Medical Sciences, our IRB agreed		
that:		
Review Comments:		

APPROVED

Note: The validity of this approval is 1 year. Site PI must abide by the approved documents. If the trial could not accomplish before the validity (including the statistical analysis), please submit for continuing review 1 month before the deadline. If the trial accomplished in the validity, please submit the final report. If there is any adverse event related to the trial occurred, please reported to the committee. If there is any change about the protocol, informed consent, or investigators, modification application must be submitted to the committee and get approved.

Director √Assistant Director S	Signature:	Date: Feb 5, 2013
Institutional Review Board of Don	gzhimen Hospital Affiliated to	Place: the 1 st meeting room
Beijing University of Chinese Med	licine	
Continuing Review Frequency:	3 mon 6 mon $\sqrt{12}$ mon Conta	ct: Jianwei Shang +86 010 84013229

Institutional Review Board of Dongzhimen Hospital Affiliated to Beijing University of Chinese Medicine

Meeting Attendance Sheet

Project Title	The Effect and Safety of Electro-acupuncture for Women with Pure Stress Urinary		
	Incontinence: a Multicenter, Randomized Controlled Trial		
Meeting Date	Feb 1, 2013 Meeting Place the 1 st meeting room of Dongzhimen Hospital		

Member	Gender	Position	Major	Signature
Pengtao Li	Male	Director	Cerebropathy	
Ying Gao	Female	Assistant Director	Cerebropathy	
Yong'an Ye	Male	Assistant Director	Gastroenterology	
Hongfang Liu	Female	Assistant Director	Nephropathy Endocrine	
Yongtao Zhang	Male	Committee Member	Respirology	
Xinyue Wang	Female	Committee Member	Gastroenterology	
Bohua Yang	Male	Committee Member	Surrounding blood-vessel	
Weixing Lu	Male	Committee Member	Angiocardiopathy	
Pengwen Wang	Female	Committee Member	Pharmacology	
Junling Cao	Male	Committee Member	Pharmacy	
Kai Liu	Male	Committee Member	Legal Representative	
Haidong He	Male	Committee Member	Medical Equipment	
Shengli Zhang	Male	Committee Member	People's Representative	
Xinyi Chen	Male	Committee Member	Hemooncology	
Shulian Peng	Female	Committee Member	Breast Surgery	

Attachment 4: Ethical approval of Yueyang Hospital of Integrated Traditional Chinese and Western Medicine affiliated to Shanghai University of Traditional Chinese Medicine

上海中医药大学附属岳阳中西医结合医院伦理委员会

IRB of Yueyang Hospital of Integrated Traditional Chinese and Western Medicine, Shanghai University of TCM

伦理审查批件

Approval Notice Template

伦理审议批件号:上海中医药大学附属岳阳中西医结合医院伦理委员会 2013 伦理审查 033 号 (20 β - 03 号)

研究名称	电针治疗女性单纯性压力性尿失禁有效性和安全性多中心随机对照试验		
研究类型	临床试验	研究周期	两年
研究单位和研究	者 上海中医药大学附属品	岳阳中西医结合医院	陈跃来
伦理委员会审议	成员 金利国、王雪文、徐政 正、任力	令 玲、史晓、常时新	f、陈云飞、樊民胜、 居
伦理委员会地址	上海市虹口区甘河路 1	10 号	
审议时间	2013年4月27日		e/ WITE
审议结论	物临床试验质量管理热学组织委员会颁布的原则。本伦理委员会颁布的原则。本伦理委员会的1、临床课题伦理审查2、研究方案(1.0 20123、知情同意书(2.0 204、主要研究者简历5、招募广告6、CRF表(1.0 2012.17、研究人员名单	5、招募广告 6、CRF 表(1.0 2012.11.09)	
	性压力性尿失禁有效性 并要求:上述资料未经 程中如发生严重不良事	注和安全性多中心随 圣本委员会批准,不 耳件,应立即(24 力 青同意书及研究者有	F展"电针治疗女性单约机对照试验"; 。得作任何修改;试验过、时内)报告本委员会; 「任何更改,应及时通知
主任委员签字	SALA		
该批件有效期一年,自批件生效日 备注 伦理委员会提交跟踪审查申请。 联系人: 肖夏懿 电话: 65161782**		产 申请。	月内未完成研究的,请向

上海中医药大学附属岳阳中西医结合医院 医学伦理委员会(盖章)

上海中医药大学附属岳阳中西医结合医院伦理委员会

IRB of Yueyang Hospital of Integrated Traditional Chinese and Western Medicine, Shanghai University of TCM

会议签到表

Sign-in Sheet of Full Board Meeting

会议日期: 2013年4月27日

审查项目: 电针治疗女性单纯性压力性尿失禁有效性和安全性多中心随机对照试验 伦理委员会到会委员签名:

10年安贝公刘云安贝3	DZ 71 +		
姓名	性别	专业情况	签名
金利国	男	医学、管理	3-11
陈云飞	男	医学	Dene
常时新	男	医学	The same
徐玲玲	女	药学	1 3 7/12
史 晓	女	中医学	
王雪文	女,	护理学	3 - 35
樊民胜	男	社会科学、伦理学	TOPPER
周正	男	法律	10) 701
任 力	男	法律	127

Institutional Review Board of Yueyang Hospital of Integrated Traditional Chinese and Western Medicine, Shanghai University of TCM

Approval Notice Template

Approval No. IRB of Yueyang Hospital of Integrated Traditional Chinese and Western Medicine, Shanghai University of TCM 2013 Ethics Approval No. 003 (2013-033)

Duainat Titla	The Effect and Sefety of Floater and	aunatura for Warran	with Dura Strazz
Project Title	The Effect and Safety of Electro-acupuncture for Women with Pure Stress		
	Urinary Incontinence: A Multicenter, Ra		
Research Attribute	Clinical Trial	Research Period	2 years
Site and Site PI	Yueyang Hospital of Integrated Tradi	tional Chinese and W	estern Medicine,
	Shanghai University of TCM Yu	elai Chen	
Review Committee	Liguo Jin, Xuewen Wang, Lingling Xu	, Xiao Shi, Shixin Cha	ng, Yunfei Chen,
	Minsheng Fan, Zheng Zhou, Li Ren		
IRB Address	110 Ganhe Road, Hongkou, Shanghai		
Review Date	Apr 27, 2013		
Review Comments	According to the "Good Clinical Prace Administration (SFDA) of the Pect "Declaration of Helsinki", and the biomedical research involving human International Organizations of Medical reviewed and discussed the files as follows: 1. Ethics Approval Application Form of 2. Study Protocol (1.0 2012.11.06) 3. Informed Consent (2.0 2013.05.10) 4. Main Researchers' CV 5. Recruiting Advertisement 6. CRF (1.0 2.12.11.09) 7. All Researchers' List 8. Other Files: Case Screening Form, R. Urination Diary The IRB approved your study "The Enforthment of Trial" through voting; The modify unless a permission was obtain event occurred during the study should If there is any change about the protocomodification application must be submined.	repole's Republic of a "International ethica an subjects" issued Sciences, all the menowing: Clinical Trial recording Sheet of the Unified and Safety of Electron Safety of Electron Safety of Electron Safety and Safety and Safety and Safety of Electron Safety and Safet	China in 2003, I guidelines for by Council for observation of our IRB Urinal Pad Using, extro-acupuncture of the notal lowed to any severe adverse within 24 hours; or investigators,
Director Signature			
Note	The validity of this approval is one	year, if the trial could	l not accomplish
	before the validity, please submit for co	ntinuing review before	the deadline.
	Contact: Xiayi Xiao Phon	ne: +86 021 65161782-	-2419

Institutional Review Board of Yueyang Hospital of Integrated Traditional Chinese and Western Medicine, Shanghai University of TCM Sign-in Sheet of Full Board Meeting

Meeting Date: Apr 27, 2013

Project Title: The Effect and Safety of Electro-acupuncture for Women with Pure Stress Urinary

Incontinence: A Multicenter, Randomized Controlled Trial

Committee Signature:

Name	Gender	Major	Signature
Liguo Jin	Male	Medicine, Management	
Yunfei Chen	Male	Medicine	
Shixin Chang	Male	Madicine	
Lingling Xu	Female	Pharmacy	
Xiao Shi	Female	Traditional Chinese Medicine	
Xuewen Wang	Female	Nursing	
Minsheng Fan	Male	Social Sciences, Ethics	
Zheng Zhou	Male	Law	
Li Ren	Male	Law	

Attachment 5: Ethical approval of West China Hospital of Sichuan University

四川大学华西医院临床试验与生物医学伦理专委会审查批件

2013年 审(7)号

科室(专业)	: 中西医结合科	项目负责人姓名及职称: 李宁 副主任医师
项目名称	电针治疗女性单纯性压力	性尿失禁有效性和安全性一多中心随机对照试验
研究方案	版本号:无	版本日期: 2013.1.7
知情同意书	版本号:修订版	版本日期: 2013.1.29

审查意见:

- 1. 研究者资质符合伦理要求。
- 2. 研究方案及知情同意书基本符合伦理要求。

审查结果: ■同意 □作必要修正后同意 □修正后再审 □不同意 □终止或暂停

请遵循我国相关法律、法规和规章(SFDA《药物临床试验质量管理规范》(2003)、《医疗器械临床试验规定》(2004)、WMA《赫尔辛基宣言》和CIOMS《人体生物医学研究国际道德指南》、卫生部《涉及人的生物医学研究伦理审查办法(试行)(2007)》),遵循伦理委员会批准的方案和知情同意书开展临床试验(研究),保护受试者的健康与权利。

在试验(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,请申请人及时提交严重不良事件报告;紧急报告之后,尽快提交详细的严重不良事件随访报告。

请递交年度和定期跟踪审查报告;当出现任何可能显著影响试验(研究)进行或增加受试者危险的情况时,请申请人及时向伦理专委会提交书面报告。

试验(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(研究)规定而未让受试者退出试验(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伦理原则与规范的情况,请申办者/监查员/研究者提交违背方案报告。

申请人暂停或提前终止临床试验(研究),请及时提交暂停/终止试验(研究)报告。 完成临床试验(研究),请申请人提交结题报告。



Institutional Review Board of West China Hospital of Sichuan University Ethics Review Approval

Approval No. 2013-7

Department:	Integrated Traditional	Chinese	Site PI:	Ning Li
	and Western Medicine			Associate Chief Physician
Project Title	The Effect and Safety o	of Electro	-acupuncture for	Women with Pure Stress
	Urinary Incontinence: a M	lulticenter	, Randomized Co	ntrolled Trial
Study Protocol	Version No. /		Version Date: Ja	an 07, 2013
Informed Consent	Version No. the Edited Ve	ersion	Version Date: Ja	an 29, 2013
Review Comments:				
1. The Site PI selec	cted met the requirements of	f ethics.		
2. The study protoc	col and informed consent me	et the requ	uirements of ethics	S.
Review Result:	approved \square approved aft	ter revisio	n □ reviewed ag	gain after revision
	not approved □terminated	or suspen	ided	
Researchers must obe	ey the related laws and regu	lations suc	ch as SFDA "Goo	d Clinical Practice (2003)",
"Provisions for Clini	"Provisions for Clinical Trials of Medical Device (2004)", WMA "Declaration of Helsinki", CIOMS			ation of Helsinki", CIOMS
"International ethical	guidelines for biomedical	research i	nvolving human s	subjects (2007)". The study
should perform accor	rding to the protocol and in	formed co	onsent approved b	by this IRB. The health and
right of the subjects s	should be protected.			
If a change of the Site	e PI, or any modification of	the proto	col/informed cons	sent was made, a new ethics
approval application of the modified files must be submitted.				
Researchers should report the severe adverse event (SAE) in time if any SAE occurred during the				
study. After the report, a detailed follow-up report of the SAE should also be submitted in time.				
Please submit the ann	Please submit the annual or regular follow-up review report in time. In any condition which will greatly			condition which will greatly
affect the progress of	affect the progress of the study or increase the risk of the subjects, a written report should be submitted			report should be submitted
to the IRB.				
The applicant/monitor/researcher should submit a protocol deviation report if any of the following				

condition occurs: 1) subjects who did not meet the inclusion criteria, or should be excluded according

to the exclusion criteria were wrongly included in the study; 2) subjects do not withdraw from the study

when he/she meet the rules of withdrawal; 3) incorrect treatment or dose was given; 4) prohibited

combined medicine was used; 5) subjects' rights and health are badly affected; 6) the science of study

was badly affected.

A concluding report should be submitted when the study is completely done or stopped prematurely.

IRB of West China Hospital of Sichuan University

Director Signature:

Date: Feb 22, 2013

天津中医药大学第一附属医院医学伦理委员会 IEC of The First Affiliated Hospital of Tianjin University of Traditional Chinese Medicine

审 查 批 件

Approval Notice

伦理批件号: TYLL2013[E]字 001

根据卫生部《涉及人的生物医学研究伦理审查办法》(2007)、国家中医药管理局《中医药临床研究伦理审查管理规范》(2010)、国家食品药品监督管理局《药物临床试验伦理审查工作指导原则》(2010)、《药物临床试验质量管理规范》(2003),以及世界医学会《赫尔辛基宣言》(2008)、国际医学科学组织理事会《人体生物医学研究国际伦理指南》(2002)的伦理原则,经天津中医药大学第一附属医院医学伦理委员会 2013 年 3 月 8 日快速审查,同意由申办者天津中医药大学第一附属医院和主要研究者傅立新共同申请的电针治疗女性单纯性压力性尿失禁有效性和安全性-多中心随机对照试验项目开展临床研究工作。

请申办者、研究人员严格遵循 GCP 规定和本伦理委员会批准的方案(版本号: VERSION1.0_20121106 版本日期: 20121106)、知情同意书(版本号: VERSION1.0_20121106 版本日期: 20121106)开展临床研究。在研究开始前,须完成临床试验注册。该项目进行中如发生下列情况,须及时书面报告本伦理委员会:①对临床方案、知情同意书等的任何修改;②更换主要研究者;③发生严重不良事件;④出现任何可能影响试验进行或增加受试者危险的情况;⑤出现违反方案情况;⑥暂停或提前终止临床研究。

本伦理委员会将对该项目跟踪审查。

请于2014年3月8日前1个月提交研究进展报告。

该项目完成后,请向本伦理委员会提交结题报告。

本批件有效期为 2013 年 3 月 8 日至 2016 年 3 月 8 日

天津中医药大学第一个阴唇医院医学伦理委员会

主任委员签字:

日

期: 2013.3.8

IEC of The First Affiliated Hospital of Tianjin University of Traditional Chinese

Medicine

Approval Notice

Approval No. TYLL2013[E] 001

According to the "ethical review methods for biomedical study involving human subjects" (2007)

issued by the Ministry of Health, "Good Clinical Practice" (2003) and "Guidelines for Ethical Review

Work of Drug Clinical Trials" (2010) issued by the State Food and Drug Administration (SFDA) of the

People's Republic of China, "management specifications for ethical review of TCM clinical studies"

issued by State Administration of Traditional Chinese Medicine, WMA "Declaration of Helsinki"

(2008), "International ethical guidelines for biomedical research involving human subjects" (2002)

issued by Council for International Organizations of Medical Sciences, through the fast review from

the IEC of the First Affiliated Hospital of Tianjin University of Traditional Chinese Medicine, the

research of "The Effect and Safety of Electro-acupuncture for Women with Pure Stress Urinary

Incontinence: a Multicenter, Randomized Controlled Trial" applied by the First Affiliated Hospital of

<u>Tianjin University of Traditional Chinese Medicine</u> and <u>Lixin Fu</u> was approved to perform.

The applicant and researchers should strictly abide by the GCP, the approved protocol (VERSION

1.0_20121106, Version Date: Nov 6, 2012), and the approved informed consent (VERSION

1.0_20121106, Version Date: Nov 6, 2012). The applicant/researcher should register this clinical trial

before the performance of the study. Written report should submit to our IEC if any of the following

occurs: 1) any modification of the study protocol, or the informed consent; 2) change of the site PI; 3)

sever adverse event occurs; 4) subjects' rights and health are badly affected; 5) protocol deviation; 6)

research paused or terminated prematurely.

Our IEC will continue reviewing the research.

The report of the study progress should be submitted 1 month before Mar 8, 2014.

Final report should be submitted after the research completion.

The validity of this approval ranges from Mar 8, 2013 to Mar 8, 2016.

IEC of The First Affiliated Hospital of Tianjin University of Traditional Chinese Medicine

Director Signature: Jinzhong Zhang

Date: Mar 08, 2013

Contact: Jingyun Jia **Phone:** +86 022 27432276

编号: AF/SC-08/01.0

伦理审查批件

批件号	湖南中医药大学第一附属医院伦理委员会 HN-LL-KY-2013-001-01			
项目名称	电针治疗女性单纯性压力性尿失禁有效性和安全性-多中心随机对照试 验			
项目来源	"十二五"国家科技支	撑计划 2012BAI24B	01	
研究单位	中国中医科学院广安门医院、湖南中医药大学第一附属医院等			
主要研究者	章薇			
审查类别	初始审查	审查方式	会议审查	
审查日期	2013.1.23	审查地点	医院会议室	
审查委员	郭志华, 贺菊乔, 赵艳玉谢海波, 钟 晓, 管小		张志国,张月娟,谭 劲	
批准文件	临床研究方案(版本号: 知情同意书(版本号:V			

审查意见

根据卫生部《涉及人的生物医学研究伦理审查办法(试行)》(2007)、SFDA《药物临床试验质量管理规范(2003)》、《医疗器械临床试验规定(2004)》、WMA《赫尔辛基宣言》和 CIOMS《人体生物医学研究国际道德指南》的伦理原则,经本伦理委员会审查,同意按所批准的临床研究方案、知情同意书、招募材料开展本项研究。

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。

研究开始前,请申请人完成临床试验注册。

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, 请申请人及时提交严重不良事件报告。

请按照伦理委员会规定的年度/定期跟踪审查频率,申请人在截止日期前 1 个月提交研究进展报告;申办者应当向组长单位伦理委员会提交各中心研究进展的汇总报告;当出现任何可能显著影响试验进行、或增加受试者危险的情况时,请申请人及时向伦理委员会提交书面报告。

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/监查员/研究者提交违背方案报告。

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。



年度/定期跟踪审查频率	12 个月
有效期	自批件下发之日起一年内有效
联系人与联系电话	赵鸿 王华 0731-85369233
主任委员签字	郭孝子.
伦理委员会	湖南中医药大学第一附属医院伦理委员会(盖章
日期	2013年1月24日



湖南中医药大学第一附属医院伦理委员会

审查会议签到表

日期: 2013年1月23日 伦理委员会到会委员签名:

		20 20 20 A7 . 102 //	
姓名	性别	专业情况	签名
郭志华	男	湖南中医药大学第一附属医院 心血管内科 主任医师 教授	新产品
贺菊乔	男	湖南中医药大学第一附属医院 中医外科 主任医师 教授	5/04
赵艳玲	女	湖南中医药大学第一附属医院 主任医师 教授	# Fol
陈其华	男	湖南中医药大学第一附属医院 中医外科 主任医师 教授	433
黄孟君	男	湖南中医药大学第一附属医院 中医消化 教授	为3.3
张月娟	女	湖南中医药大学第一附属医院 护理 主任护师 教授	2/2mx
张志国	男	湖南中医药大学第一附属医院 药学 主任药师 教授	别是到
谭劲	男	湖南中医药大学第一附属医院 中西医结合口腔 主任医师 教授	福动
谢海波	男	湖南中医药大学第一附属医院 中医内科 副主任医师 副教授	B7474
管小平	男	律师, 融源律师事务所	M. T
钟晓	女	保险 太平人寿湖南分公司,业务经理	基本 晚
赵鸿(秘书)	女	湖南中医药大学第一附属医院 护理 副主任护师 副教授	AIN

No. AF/SC-08/01.0

Ethics Review Approval

Accepted No.	Institutional Review Board	of the First Affiliate	d Hospital of Hunan University of
	Chinese Medicine HN-LL-KY-2013-001-01		
Project Title	The Effect and Safety of El	ectro-acupuncture fo	r Women with Pure Stress Urinary
	Incontinence: a Multicenter	r, Randomized Contr	olled Trial
Project Sponsor	the 12th Five-year Plan of t	he National Key Tec	hnology Support Program by the
	Ministry of Science and Te	chnology of the Peop	ole's Republic of China
	2012BAI24B01		
Research Site	Guang'anmen Hospital of China Academy of Chinese Medical Sciences, the First		
	Affiliated Hospital of Hunan University of Chinese Medicine		
Site PI	Wei Zhang		
Review Attribute	Initial Review	Review Methods	Review Conference
Review Date	Jan 23, 2013	Review Place	Meeting Room of the Hospital
Review Committee	Zhihua Guo, Juqiao He, Yanling Zhao, Qihua Chen, Mengjun Huang, Zhiguo Zhang,		
	Yuejuan Zhang, Jin Tan, Haibo Xie, Xiao Zhong, Xiaoping Guan		
Approved Files	Study Protocol (Version No	o. VERSION 1.0-201	21106)
	Informed Consent (Version	No. VERSION 1.0-	201201109)

Review Comments

According to the "ethical review methods for biomedical study involving human subjects" (2007) issued by the Ministry of Health, "Good Clinical Practice" (2003) and "Provisions for Clinical Trials of Medical Device" (2004) issued by SFDA, WMA "Declaration of Helsinki", and the COIMS "International ethical guidelines for biomedical research involving human subjects", through the review of our IRB, the study protocol, informed consent, and related recruitment files were approved.

Please conform to the principle of GCP, and conform to the protocol approved by our IRB, and protect the health and rights of the subjects.

The applicant or PI should register this clinical trial online before the start of the study.

An application should be submitted if a change of the site PI, or any modification of the study protocol, informed consent, or the recruitment files are made.

The report of the severe adverse event (SEA) should be submitted in time if any SAE occurs.

Please do the follow-up review annually or termly according to the stipulation of our IRB. The report of the study progress should be submitted one month before the deadline. A summary report of the study progress should be submitted to the IRB of the leading site. In any condition which will greatly affect the study progress or increase the potential risk of the subjects, a written report should be submitted by the applicant/researcher to IRB.

A protocol violation report should be submitted by the applicant/monitor/researcher in the following conditions: 1) conditions that violate the study protocol: subjects who did not meet the inclusion criteria, or should be excluded according to the exclusion criteria were wrongly included in the study; subjects do not

Institutional Review Board of the First Affiliated Hospital of Hunan University of Chinese Medicine

withdraw from the study when he/she meet the rules of withdrawal; incorrect treatment or dose was given; prohibited combined medicine was used; 2) conditions that violate GCP principle: subjects' rights and health are badly affected; the science of study was badly affected.

A final report should be submitted when the study is completely done or stopped prematurely.

Review Frequency	12 months
Validity	1 year
Contact	Hong Zhao / Hua Wang +86 0731 85369233
Director	
IRB	Institutional Review Board of the First Affiliated Hospital of Hunan University of Chinese Medicine
Date	Jan 24, 2013

Institutional Review Board of the First Affiliated Hospital of Hunan University of Chinese Medicine

Sign-in Sheet of the Review Conference

Date: Jan 23, 2013

Review Committee Signature:

Name	Gender	Major	Signature
Zhihua Guo	Male	Chief Physician, Professor, Cardiovascular medicine, the first	
		affiliated hospital of Hunan University of Chinese Medicine	
Juqiao He	Male	Chief Surgeon, Professor, Traditional Chinese Surgery, the first	
		affiliated hospital of Hunan University of Chinese Medicine	
Yanling Zhao	Female	Chief Physician, Professor, the first affiliated hospital of Hunan	
		University of Chinese Medicine	
Qihua Chen	Male	Chief Surgeon, Professor, Traditional Chinese Surgery, the first	
		affiliated hospital of Hunan University of Chinese Medicine	
Mengjun	Male	Chief Physician, Professor, Gastrointestinal Medicine, the first	
Huang		affiliated hospital of Hunan University of Chinese Medicine	
Yuejuan Zhang	Female	Chief Nurse, Professor, Nursing, the first affiliated hospital of	
		Hunan University of Chinese Medicine	
Zhiguo Zhang	Male	Chief Pharmacist, Professor, Pharmacy, the first affiliated hospital	
		of Hunan University of Chinese Medicine	
Jin Tan	Male	Chief Dentist, Professor, Integrated Chinese and Western Oral	
		Medicine, the first affiliated hospital of Hunan University of	
		Chinese Medicine	
Haibo Xie	Male	Assistant Chief Physician, Assistant Professor, Traditional	
		Chinese Medicine, the first affiliated hospital of Hunan University	
		of Chinese Medicine	
Xiaoping Guan	Male	Lawyer, Rongyuan Law Office	
Xiao Zhong	Female	Business Manager, Insurance, Tai Ping Life Hunan Branch Office	
Hong Zhao	Female	Assistant Chief Nurse, Assistant Professor, Nursing, the first	
(secretary)		affiliated hospital of Hunan University of Chinese Medicine	

伦理审查批件

项目編号 2013EC001 项目来源 "十二五"国家科技支撑计划 牵头单位 湖南中医药大学附属衡阳医院 中办者(如有) 主要研究者 岳增辉 申查方式 医院门诊楼 11 楼会议室 申查日期 2013.2.17 审查地点 医院门诊楼 11 楼会议室 申查委员 王诚喜、龙双才、邹岳萍、徐基平、钟新林、匡肇、董秋萍、贺新民、谢军、万贤明、谢春完 代联区区区、02121105),知情同意书(V1.0) 根据卫生部《涉及人的生物医学研究伦理审查办法(试行)》、国家药品自品监督管理局《药物临床试验质量管理规范》、《医疗器械临床试验规定》、《多物临床试验伦理审查工作指导原则》、国家中医药管理局《中医药临床研究伦理审查管理规范》以及《赫尔辛基宣言》和国际医学科学组织委员会颁布的《人体生物医学研究国际道德指南》的伦理原则《多年伦理委员会申查,同意按例批准的临床研究方案、知情同意书、招募材料开展本项研究。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。 如发生严重不良事件以及影响研究风险受益比的非预期不良事件,请申请人及时提交严重不良事件报告。 请按照伦理委员会规定的年度/定期跟踪审查频率,申请人在截止日期前一个月提交研究报告;申报者应当向组长单位伦理委员会提交各中心的研究过展的汇总报告。当出现任何可能显著影响试验进行或增加受试者危险的情况时请申请人及时向伦理委员会提交书面报告。 研究纳入了不符合纳入标准或符合排除标准的受试者,符合中止试验规是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污有递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设成不良影响等违背 GCP 原则的情况,请申办者/监察员/研究者提交违背方案指告、提前终止或完成临床研究,请及时提交结题报告。	项目名称	电针治疗女性单纯性	生压力性尿失禁有效性	性和安全性多中心随即对照试验	
學具单位 申办者(如有) 主要研究者 中查类別 申查方式 申查共別 申查方式 申查共別 申查方式 申查共別 申查方式 申查共別 申查方式 申查共別 申查方式 申查地点 医院门诊楼11 楼会议室 申查 表	项目编号				
主要研究者 岳增辉 审查类别 初始审查 审查方式 会议审查 医院门诊楼 11 楼会议室 审查日期 2013.2.17 审查地点 医院门诊楼 11 楼会议室 审查日期 2013.2.17 审查地点 医院门诊楼 11 楼会议室 可查委员 王诚喜、龙双才、邹岳萍、徐基平、钟新林、匡肇、董秋萍、贺新民、谢军、万贤明、谢春亮 批准文件 研充方案:(VERSIONI.020121105),知情同意书(VI.0) 服据卫生部《涉及人的生物医学研究伦理审查办法(试行)》。国家结品信题管理局《药物临床试验质量管理规范》、《医疗器械临床试验规定》、《委物临床试验伦理审查工作指导原则》、国家中医药管理局《中医药临床研究伦理审查管理规范》以及《赫尔辛基宣言》和国际医学科学组织委员会颁布的《人体生物医学研究国际道德指南》的伦理原则,经本伦理委员会审查,同意按例批准的临床研究方案、知情同意书、招募材料开展本项研究。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。 如发生严重不良事件以及影响研究风险受益比的非预期不良事件,请申请人及时境定严重不良事件以及影响研究风险受益比的非预期不良事件,请申请人及时是定严重不良事件报告。 "有接账伦理委员会规定的年度/定期跟踪审查频率,申请人在截止日期前一个月提交研究报告,申出现任何可能显著影响试验进行或增加受试者危险的情况时请申请人及时向伦理委员会提交书面报告。 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资有遵从方案开展研究的情况,或可能处试者的权益/健康以及研究的科学性遗成不良影响等选背。CCP 原则的情况,请申办者/监察员/研究者提交违背方案指传。 提前终止或完成临床研究,请及时提交结题报告。 2013 年 2 月 19°2014 年 2 月 18 日 跃系人及电话 谢军,0734-8137737	牵头单位	湖南中医药大学附属衡阳医院			
主要研究者 岳增辉 审查类别 初始审查 审查方式 会议审查 医院门诊楼 11 楼会议室 审查日期 2013.2.17 审查地点 医院门诊楼 11 楼会议室 审查日期 2013.2.17 审查地点 医院门诊楼 11 楼会议室 可查委员 王诚喜、龙双才、邹岳萍、徐基平、钟新林、匡肇、董秋萍、贺新民、谢军、万贤明、谢春亮 批准文件 研充方案:(VERSIONI.020121105),知情同意书(VI.0) 服据卫生部《涉及人的生物医学研究伦理审查办法(试行)》。国家结品信题管理局《药物临床试验质量管理规范》、《医疗器械临床试验规定》、《委物临床试验伦理审查工作指导原则》、国家中医药管理局《中医药临床研究伦理审查管理规范》以及《赫尔辛基宣言》和国际医学科学组织委员会颁布的《人体生物医学研究国际道德指南》的伦理原则,经本伦理委员会审查,同意按例批准的临床研究方案、知情同意书、招募材料开展本项研究。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。 如发生严重不良事件以及影响研究风险受益比的非预期不良事件,请申请人及时境定严重不良事件以及影响研究风险受益比的非预期不良事件,请申请人及时是定严重不良事件报告。 "有接账伦理委员会规定的年度/定期跟踪审查频率,申请人在截止日期前一个月提交研究报告,申出现任何可能显著影响试验进行或增加受试者危险的情况时请申请人及时向伦理委员会提交书面报告。 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资有遵从方案开展研究的情况,或可能处试者的权益/健康以及研究的科学性遗成不良影响等选背。CCP 原则的情况,请申办者/监察员/研究者提交违背方案指传。 提前终止或完成临床研究,请及时提交结题报告。 2013 年 2 月 19°2014 年 2 月 18 日 跃系人及电话 谢军,0734-8137737	申办者(如有)				
申查日期 2013. 2.17 审查地点 医院门诊楼 11 楼会议室 王诚喜、龙双才、邹岳萍、徐基平、钟新林、匡肇、董秋萍、贺新民、谢军、万贤明、谢春亮 研究方案: (VERSIONI. 020121105),知情同意书 (VI. 0) 根据卫生部《涉及人的生物医学研究伦理审查办法(试行)》、国家药品信品监督管理局《药物临床试验质量管理规范》、《医疗器械临床试验规定》、《经物临床试验处理审查工作指导原则》、国家中医药管理局《中医药临床研究伦理审查管理规范》以及《赫尔辛基宣言》和国际医学科学组织委员会颁布的《人体生物医学研究国际道德指南》的伦理原则,经本伦理委员会师有。《内体生物医学研究国际道德指南》的伦理原则,经本伦理委员会而有,同意按例批准的临床研究方案、知情同意书、招募材料开展本项研究。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。 如发生严重不良事件以及影响研究风险受益比的非预期不良事件,请申请人及时提交严重不良事件以及影响研究风险受益比的非预期不良事件,请申请人及时提处严重不良事件以及影响研究风险受益比的非预期不良事件,请申请人及时提交严重不良事件和告。 请按照伦理委员会规定的年度/定期跟踪审查频率,申请人在截止日期前一个月提交研究报告: 当出现任何可能显著影响试验进行或增加受试者危险的情况时请申请人及时向伦理委员会提交书面报告。 研究纳入了不符合纳入标准或符合排除标准的受试者,符合中止试验规是而未让受试者退出研究,给予节或剂量,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设成不良影响等违背 GCP 原则的情况,请申办者/监察员/研究者提交违背方案指告。 提前终止或完成临床研究,请及时提交结题报告。 2013 年 2 月 19°2014 年 2 月 18 日 谢军,0734-8137737	主要研究者	岳增辉			
中古委员 王诚喜、龙双才、邹岳萍、徐基平、钟新林、匡肇、董秋萍、贺新民、谢军、万贤明、谢春亮 他准文件 研究方案:(VERSIONI.020121105),知情同意书(VI.0) 审查意见 根据卫生部《涉及人的生物医学研究伦理审查办法(试行)》、国家药品自品监督管理局《药物临床试验质量管理规范》、《医疗器械临床试验规定》、《经物临床试验危理审查工作指导原则》、国家中医药管理局《中医药临床研究伦理审查管理规范》以及《赫尔辛基宣言》和国际医学科学组织委员会颁布的《人体生物医学研究国际道德指南》的伦理原则,经本伦理委员会而在《人体生物医学研究国际道德指南》的伦理原则,经本伦理委员会而不完,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。 如发生严重不良事件以及影响研究风险受益比的非预期不良事件,请申请人及时提交严重不良事件报告。 请按照伦理委员会规定的年度/定期跟踪审查频率,申请人在截止日期前一个月提交研究报告;申报者应当向组长单位伦理委员会提交各中心的研究过展的汇总报告;当出现任何可能显著影响试验进行或增加受试者危险的情况时请申请人及时向伦理委员会提交书面报告。 研究创入区符合纳入标准或符合排除标准的受试者,符合中止试验规是而未让受试者退出研究,给予背误治疗或剂量,给予方案禁止的合并用药等发育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指告。 提前终止或完成临床研究,请及时提交结题报告。 2013 年 2 月 19 2014 年 2 月 18 日	审查类别	初始审查	审查方式	会议审查	
开资明、谢春亮	审查日期	2013. 2. 17	审查地点	医院门诊楼 11 楼会议室	
根据卫生部《涉及人的生物医学研究伦理审查办法(试行)》、国家药品信品监督管理局《药物临床试验质量管理规范》、《医疗器械临床试验规定》、《教物临床试验伦理审查工作指导原则》、国家中医药管理局《中医药临床研究伦理审查管理规范》以及《赫尔辛基宣言》和国际医学科学组织委员会颁布的《人体生物医学研究国际道德指南》的伦理原则,经本伦理委员会审查,同意按射批准的临床研究方案、知情同意书、招募材料开展本项研究。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。 如发生严重不良事件以及影响研究风险受益比的非预期不良事件,请申请人及时提交严重不良事件报告。 请按照伦理委员会规定的年度/定期跟踪审查频率,申请人在截止日期前一个月提交研究报告;申报者应当向组长单位伦理委员会提交各中心的研究过展的汇总报告;当出现任何可能显著影响试验进行或增加受试者危险的情况时请申请人及时向伦理委员会提交书面报告。 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递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设成不良影响等违背 GCP 原则的情况,请申办者/监察员/研究者提交违背方案指告。 提前终止或完成临床研究,请及时提交结题报告。 1	审查委员		邓岳萍、徐基平、钟新	析、匡肇、董秋萍、贺新民、谢军、	
品监督管理局《药物临床试验质量管理规范》、《医疗器械临床试验规定》、《多物临床试验伦理审查工作指导原则》、国家中医药管理局《中医药临床研究伦理审查管理规范》以及《赫尔辛基宣言》和国际医学科学组织委员会颁布的《人体生物医学研究国际道德指南》的伦理原则,经本伦理委员会审查,同意按例批准的临床研究方案、知情同意书、招募材料开展本项研究。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。 如发生严重不良事件以及影响研究风险受益比的非预期不良事件,请申请人及时提交严重不良事件报告。 请按照伦理委员会规定的年度/定期跟踪审查频率,申请人在截止日期前一个月提交研究报告;申报者应当向组长单位伦理委员会提交各中心的研究过展的汇总报告;当出现任何可能显著影响试验进行或增加受试者危险的情况时请申请人及时向伦理委员会提交书面报告。 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沒有递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设成不良影响等违背 GCP 原则的情况,请申办者/监察员/研究者提交违背方案报告。 提前终止或完成临床研究,请及时提交结题报告。 1	批准文件	研究方案: (VERSIO	N1. 020121105),知情	情同意书 (V1.0)	
联系人及电话	审查意见	品监督管理局域的 管理是是是一个的事件。 一个的事件。 一个的事件。 一个的事件。 一个的事件。 一个的事件。 一个的事件, 一个的事件。 一个的事件。 一个的事件, 一个的事件。 一个的事件, 一个的事件。 一个的事件, 一个的事件。 一个的事件, 一个的事件。 一个的事件, 一个的事件。 一个的事件, 一个的事件。 一个的事件, 一个的事件。 一个的。 一个的。 一个的。 一个的。 一个的。 一个的。 一个的。 一个的	你怎么就是我们的一个一个一个一个一个一个一个一个一个一个一个一个一个一个一个一个一个一个一个	观范》、《医疗器械临床试验规定》、《药家中医药管理局《中医药临床研究伦理和国际医学科学组织委员会颁布的《人原则,经本伦理委员会审查,同意按所愿材料开展本项研究。 此准的方案开展临床研究,保护受试者临床研究方案、知情同意书、招募材料。 证本的方案、知情同意书、招募材料。 证实验述的非预期不良事件,请申请跟踪审查频率,申请人在截止日期前1 跟踪审查频率,申请人在截止日期前1 跟踪审查频率,申请人在截止日期前1 战论进行或增加受试者危险的情况时, 证实验进行或增加受试者危险的情况时, 证实验进行或增加受试者危险的情况时, 证实验过行或增加受试者危险的情况时, 证实验过行或增加受试者危险的情况时, 证实验过行或增加受试者危险的情况时, 证实验过行或增加受试者危险的情况时, 证实验过行或增加受试者危险的情况时, 证实验过行或增加受试者危险的情况时, 证实验过行或增加受试者危险的情况时, 证实验过行或增加受试者危险的情况时,	
注任委员签字 湖南中医药大学附属衡阳医院伦理委员会(盖章)	有效期	2013年2月19~201	4年2月18日		
湖南中医药大学附属衡阳医院伦理委员会(盖章)	联系人及电话	谢军,0734-813773	7	人 联药大心	
	主任委员签字		2	S COMMANDER OF THE PARTY OF THE	
2013年2月19日			湖南中医药力	大学附属衡阳医院伦理委员会(盖章)	
			20	13年2月19日	

$Institutional\ Review\ Board\ Documentation\ of\ Hengyang\ Hospital\ affiliated\ to Hunan\ University\ of$ $Chinese\ Medicine\ (EC_AF_2013001)$

Ethics Review Approval

Project Title	The Effect and Safety of	Electro-acupuncture f	For Women with Pure Stress Urinary Incontinence: a		
	Multicenter, Randomized Controlled Trial				
Approval No.	2013EC001	Project Sponsor	the 12 th Five-year Plan of the National Key		
			Technology Support Program by the Ministry of		
			Science and Technology of the People's		
			Republic of China		
Clinical Site	Hengyang Hospital Affilia	ated to Hunan Univer	sity of Chinese Medicine		
Applicant (if any)	/				
Site PI	Zenghui Yue				
ReviewAttribute	Initial Review	ReviewMethods	Meeting Review		
Review Date	Feb 17, 2013	Review Place	Meeting Room, the 11 th Floor of the Clinic		
			Building of Yueyang Hospital		
Review Committee	Chengxi Wang, Shuangca	ai Long, Yueping Zo	u, Jiping Xu, Xinlin Zhong, Zhao Kuang, Qiuping		
	Dong, Xinmin He, Jun Xi	e, Xianming Wan, Ch	nunliang Xie		
Approved Files	Study Protocol (VERSIO	N 1.0_20121105), Inf	Formed Consent (V1.0)		
Review Comments	According to "ethical rev	riew methods for bior	medical study involving human subjects" issued by		
	the Ministry of Health,	"Good Clinical Prac	ctice", "Provisions for Clinical Trials of Medical		
	Device" and "Guidelines	for Ethical Review V	Work of Drug Clinical Trials" issued by State Food		
	and Drug Administration	(SFDA) of the People	e's Republic of China, "management specifications		
	for ethical review of TCM clinical studies" issued by State Administration of Traditional Chinese Medicine, Declaration of Helsinki, and "International ethical guidelines for biomedical research				
	involving human subjects" made by Council for International Organizations of Medical Sciences,				
	this clinical research wa	this clinical research was reviewed by the institutional review board (IRB) of Guang'anmen			
	Hospital of China Acader	my of Chinese Medic	al Sciences. And the protocol and informed consent		
	of this research were appr	oved.			
	Please conduct this clinic	al study following the	GCP principles and the study protocol approved by		
	our IRB. The health and r	ights of the subjects s	should be protected throughout the whole study.		
	An application should be	submitted if the char	nge of the site PI, or any modification of the study		
	protocol, informed consent, or the recruitment files are made.				
	The report of the severe adverse events (SAE) should be submitted in time if any SAE or any other				
	un-anticipated adverse ev	ent, which will affect	the risk-reward ratio of this study, occurs.		
	Researchers should submit the report of study progress before one month of the deadline according				
	to the frequency of ethical review. A summary report of the study progressshould be submitted to				
	the IRB of the leading ce	enter. In any condition	n which will greatly affect the progress of the study		
	or increase the potential r	risk of the subjects, a	written report should be submitted by the applicant		
	to IRB.	•	-		
	A protocol violation report should be submitted by the applicant/monitor/researcher in the				
	following conditions: 1) conditions that violate the study protocol: subjects who did not meet the				
			ding to the exclusion criteria were wrongly included		

$Institutional\ Review\ Board\ Documentation\ of\ Hengyang\ Hospital\ affiliated\ to Hunan\ University\ of\ Chinese\ Medicine\ (EC_AF_2013001)$

	in the study; subjects do not withdraw from the study when he/she meet the rules of withdrawal;
	incorrect treatment or dose was given; prohibited combined medicine was used; 2) conditions that
	violate GCP principle: subjects' rights and health are badly affected; the science of study was
	badly affected.
	A final report should be submitted when the study is completely done or stopped prematurely.
Validity Period	From Feb 19, 2013 to Feb 18, 2014
Contact	Jun Xie, +86 07348137737
Director Signature	
	IRB of Hengyang Hospital Affiliated toHunan University of Chinese Medicine (Seal)
	Date: Feb 19, 2013

Page 1

Version No. 1.00 / **Version Date:** 20130211

湖北省中医院伦理委员会

Ethics Committee of Hubei Province Hospital of Traditional Chinese Medicine

伦理审查批件

Ethics Review Approval

		1 1		
批件号	HBZY2013-C007-01			
项目名称	电针治疗女性单纯性压力性尿失禁有效性和安全性多中心随机对照			
	验			
申办者	中国中医科学院广安门医			
研究单位	中国中医科学院广安门医院、北京中医药大学东直门医院、四川大学华西医院、中国中医科学院西苑医院、湖南中医药大学附属衡阳医院、湖南省中医院、上海中医药大学附属岳阳中西医结合医院、天津中医药大学第一附属医院、陕西省中医医院、江苏省中医院、山西中医学院中西医结合医院、湖北省中医院			
主要研究者	周仲瑜 主任医师			
审查类别	初始审查	审查方式	会议审查	
审查日期	2013-01-23	审查地点	湖北省中医院伦理办会议室	
审查委员	涂远超、文建华、刘建忠、郭艳红、费兰波、程业刚、王小琴、高文 喜、周忠明、胡晓雪、石艳红、吴胜利			
批准文件	临床研究方案版本号/日期: VERSION1. 0_20121106/2012-11-06; 受试者知情同意书版本号/日期: V1. 0/2012-11-06。			
	审查	意见		

根据卫生部《涉及人的生物医学研究伦理审查办法(试行)》(2007)、SFDA《药物临床 试验质量管理规范 (2003)》、《医疗器械临床试验规定 (2004)》、WMA 《赫尔辛基宣言》和 CIOMS《人体生物医学研究国际道德指南》的伦理原则,经本伦理委员会审查,同意按所批 准的临床研究方案、知情同意书开展该项研究。

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。 研究开始前,请申请人完成临床试验注册。

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,请申请人及时提交严重不良事件报告。

请按照伦理委员会规定的年度/定期跟踪审查频率,申请人在截止日期前1个月提交研 究进展报告。

出现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/监查员/研究者提交违背方案报告。

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。

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湖北省中医院伦理委员会(盖章)

湖北省中医院伦理委员会

Ethics Committee of Hubei Province Hospital of Traditional Chinese Medicine

会议签到表

Sign-in Sheet of Meeting

项目名称	①肾力欣颗粒 II 期临床试验②红花黄色素注射液 II b 期临床试验③ZONCARE-S9 型全数字彩超临床验证④灯盏丹芪胶囊III期临床试验⑤臭氧综合治疗仪临床验证⑥连花急支片III期临床试验⑦电针治疗女性尿失禁⑧扶阳罐疗法⑨龙牡壮骨颗粒临床试验
会议日期	2013年1月23日

姓名	性别	专业背景	签名
涂远超	男	心血管内科	ME
巴元明	男	中医肾病	
刘建忠	男	中医儿科	3/3/2
文建华	男	中医内科	5 PM
郭艳红	女	行政管理	& Patern
费兰波	女	中医针灸	A Shi Z
程业刚	男	中西医结合肾病	K Frys
王小琴	女	中医内科	2013
高文喜	男	中医外科	125
周忠明	男	妇产科	A A A
胡晓雪	女	药学	- Why
吴胜利	男	律师	TMAN .
石艳红	女	社区警务	石巷(2
张 馨	女	中西医结合临床	连
陈学军	男	科研管理	1835

请假

Ethics Committee of Hubei Province Hospital of Traditional Chinese Medicine Ethics Review Approval

Approval No.	HBZY2013-C007-01			
Project Title	The Effect and Safety of Electro-acupuncture for Women with Pure Stress			
	Urinary Incontinence: a Multicenter, Randomized Controlled Trial			
Bidder	Guang'anmen Hospital of China	Academy of Chinese	Medical Sciences	
Clinical Sites	Guang'anmen Hospital of Chin	na Academy of Ch	inese Medical Sciences,	
	Dongzhimen Hospital of Beijing	University of Chine	se Medicine, West China	
	Hospital of Sichuan University, X	Kiyuan Hospital of Cl	nina Academy of Chinese	
	Medical Sciences, Hengyang	Hospital of Hunan	University of Chinese	
	Medicine, the Second Affiliated	d Hospital of Huna	n University of Chinese	
	Medicine, Yueyang Hospital of	Integrated Tradition	nal Chinese and Western	
	Medicine of Shanghai Universit	y of Traditional Chi	inese Medicine, the First	
	Affiliated Hospital of Tianjin	University of Chin	nese Medicine, Shaanxi	
	Province Hospital of TCM, Jiangsu Province Hospital of TCM, Shanxi			
	Hospital of Integrated Chinese and Western Medicine, Hubei Province Hospital			
	of TCM			
Site PI	Zhongyu Zhou, Chief physician			
Review Attribute	Initial Review	Review Method	Review Conference	
Review Date	Jan 23, 2013	Review Place	IRB Conference Room	
Review Committee	Yuanchao Tu, Jianhua Wen, Jian	zhong Liu, Yanhong	Guo, Lanbo Fei, Yegang	
	Cheng, Xiaoqin Wang, Wenxi G	ao, Zhongming Zho	u, Xiaoxue Hu, Yanhong	
	Shi, Shengli Wu			
Approved Files	Study Protocol: Version No. VERSION1.0_20121106, Date: Nov 6, 2012			
	Informed Consent: Version No. V1.0, Date: Nov 6, 2012			
	Review Comments			
According to the "ethical review methods for biomedical study involving human subjects (trial)"				
(2007) issued by the M	(2007) issued by the Ministry of Health, the "Good Clinical Practice" (2003), "Provisions for Clinical			
Trials of Medical De	Trials of Medical Device" (2004) issued by SFDA, WMA "Declaration of Helsinki", and CIOMS			

"International ethical guidelines for biomedical research involving human subjects", through the review of this IRB, the study protocol and informed consent were approved to perform.

Please conform to the GCP principle, and conform to the study protocol approved by this IRB. The health and rights of the subjects should be protected throughout the study. The study should be registered online before its start.

An application should be submitted if any change of the site PI, or any modification of the study protocol, informed consent, or recruitment files are made. A severe adverse events (SAE) report should be submitted in time if any SAE occurs during the study.

The researcher/applicant should submit the study progress report one month before the deadline in reference to the annual/periodical review frequency of our IRB.

A report of protocol deviation should be submitted by the applicant/monitor/researcher when the following conditions occur: 1) conditions that violate the study protocol: subjects, who do not meet the inclusion criteria, or should be excluded according to the exclusion criteria, were included in the study; subjects do not withdraw from the study when he/she meets the rules of withdrawal; incorrect treatment or dose was given; prohibited combine medicine was used; 2) conditions that violate GCP principle: subjects' rights and health are badly affected; the science of study was badly affected.

A paused/terminated report should be submitted if the study is paused or terminated prematurely.

A concluding report should be submitted if the study finished completely.

Review Frequency	12 months	
Validity Period	12 months	
Contact	Xin Zhang / Xuejun Chen	Phone: +86 027 88920956
Director Signature	Yuanchao Tu	

Ethics Committee of Hubei Province Hospital of TCM (Seal)

Date: Jan 30, 2013

Ethics Committee of Hubei Province Hospital of Traditional Chinese Medicine Sign-in Sheet of Meeting

Project Title	The Effect and Safety of Electro-acupuncture for Women with Pure Stress Urinary
	Incontinence: a Multicenter, Randomized Controlled Trial
Meeting Date	Jan 23, 2013

Name	Gender	Major	Signature
Yuanchao Tu	Male	Cadiovascular Medicine	
Yuanming Ba	Male	Nephropathy of TCM	
Jianzhong Liu	Male	Pediatrics of TCM	
Jianhua Wen	Male	Traditional Chinese Medicine	
Yanhong Guo	Female	Administration	
Lanbo Fei	Female	Acupuncture and Moxibustion	
Yegang Cheng	Male	Nephropathy of Integrated Chinese and Western Medicine	
Xiaoqin Wang	Female	Traditional Chinese Medicine	
Wenxi Gao	Male	Surgery of TCM	
Zhongming Zhou	Male	Gynaecology and Obstetrics	
Xiaoxue Hu	Female	Pharmacy	
Shengli Wu	Male	Lawyer	
Yanhong Shi	Female	Community Policing	
Xin Zhang	Female	Clinical Integrated Chinese and Western Medicine	
Xuejun Chen	Male	Scientific Research Management	

伦理审查批件

批件号	2013NL-013-04				
项目名称	电针治疗女性单纯性压力性尿失禁有效性和安全性多中心随机对照试验				
项目来源	"十二五"国家科技支撑计划				
研究单位	江苏省中医院, 中国中医科学院广安门医院				
主要研究者	孙建华				
审查类别	复审申请	审查方式	快速审查		
审查日期	2013 年 06 月 25 日 审查地点				
审查委员	殷立平				
	复审申请				
审查文件	技术合作合同				

审查意见

根据卫生部《涉及人的生物医学研究伦理审查办法(试行)》(2007)、SFDA《药物临床试验质量管理规范(2003)》、《医疗器械临床试验规定(2004)》、WMA《赫尔辛基宣言》和CIOMS《人体生物医学研究国际道德指南》的伦理原则,经本伦理委员会审查,同意按所批准的临床研究方案、知情同意书、招募材料开展本项研究。

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。研究开始前,请申请人完成临床试验注册。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,请申请人及时提交严重不良事件报告;紧急报告之后,尽快提交详细的严重不良事件随访报告。请按照伦理委员会规定的年度/定期跟踪审查频率,申请人在截止日期前 1 个月提交研究进展报告;申办者应当向组长单位伦理委员会提交各中心研究进展的汇总报告;当出现任何可能显著影响试验进行、或增加受试者危险的情况时,请申请人及时向伦理委员会提交书面报告。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/监查员/研究者提交违背方案报告。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。本项临床试验应当在批准之日起一年内实施,逾期未实施的,本批件自行废止。

请于 2014 年 06 月 25 日前 1 个月提交研究进展报告
12 个月
吴静 31618
TUSUKU +
南京中医药大学附属医院(江苏省中医院)伦理委员会(盖章)
2013年06月25日

Institutional Review Board of Jiangsu Province Hospital of Traditional Chinese Medicine

Ethics Review Approval

Approval No.	2013NL-013-04						
Project Title	The Effect and Safety of Electro-acupuncture for Women with Pure Stress Urinary						
	Incontinence: a Multicenter, Randor	Incontinence: a Multicenter, Randomized Controlled Trial					
Project Sponsor	the 12 th Five-year Plan of the National Key Technology Support Program by the						
	Ministry of Science and Technology	of the People's Repul	blic of China				
Clinical Site	Jiangsu Province Hospital of TCM, Guang'anmen Hospital						
Site PI	Jianhua Sun						
Review Attribute	Review Application	Review Method	Fast Review				
Review Date	Jun 25, 2013 Review Place						
Review Commissioner	Liping Yin						
Approved Files	Review Application						
	Technological Cooperation Contract	,	Technological Cooperation Contract				

Review Comments

According to the principles of the "ethical review methods for biomedical study involving human subjects (trial)" (2007) issued by the Ministry of Health, SFDA "Good Clinical Practice" (2003), SFDA "Provisions for Clinical Trials of Medical Device" (2004), WMA "Declaration of Helsinki" and CIOMS "international ethical guidelines for biomedical research involving human subjects", through the review of our IRB, the study protocol, informed consent, and the recruitment files were approved.

Please conform to the principle of the GCP, and the study protocol approved by this IRB. Please protect the health and rights of the subjects. This clinical trial should be registered online by the applicant/PI before its start. A revision review application should be submitted if any modification of the protocol, informed consent, or recruitment files, or a change of the site PI were made. A severe adverse event (SAE) report should submitted in time if any SAE occurs, and a follow-up SAE report should also be submitted in time after that. The report of the study progress should be submitted one month before the deadline according to the review frequency. Summary report should be submitted by the applicant/site PI to the IRB of the leading site. If any condition that will influence the progress of the study or increase the risk of the subjects occurs, a written report should be submitted to the IRB by the applicant/site PI. A protocol deviation report should be submitted by the applicant/monitor/researcher in the following conditions: 1) conditions that violate the study protocol: subjects who did not meet the inclusion criteria, or should be excluded according to the exclusion criteria were wrongly included in the study; subjects do not withdraw from the study when he/she meet the rules of withdrawal; incorrect treatment or dose was given; prohibited combined medicine was used; 2) conditions that violate GCP principle: subjects' rights and health are badly affected; the science of study was badly affected. The applicant or researcher should submit the paused/terminated report if the study is paused or terminated prematurely. A conclusion report should be handed in if the study is finished completely. The study should be performed within one year after this approval; otherwise, this approval will be abolished.

Review Frequency	Please submit the study progress report 1 month before Jun 25, 2014				
Validity	12 Months				
Contact	Jing Wu	Phone: +86 025 86617141-31618			

Institutional Review Board of Jiangsu Province Hospital of Traditional Chinese Medicine

Director Signature	
Institutional Review	Institutional Review Board of Jiangsu Province Hospital of TCM
Board	(the Affiliated Hospital of Nanjing University of Chinese Medicine) (SEAL)
Date	Jun 25, 2013

Attachment 11: Ethical approval of Shanxi Province Hospital of TCM

陕西省中医医院伦理委员会 临床研究伦理审查批件

					-	(2013) 1	7甲市一人0	40 5
项目名称	针灸疗效国际多中心临床评价研究							
	(电针治疗女性单纯性压力性尿失禁有效性和安全性多中心随机对照试验)							
申请单位	陕西省中医医院针灸科							
项目来源	"十二	五"国家科技	批准文号	-	课	题编号:	2012BAI	24B01
	支撑计划							
承担研究	针灸科		主要研究	2者	苏	司生	职称	主任医师
任务科室								
会议时间	2013年	02月01日	会议地点	Ĭ.	院	会议室	审查	会议审查
							方式	
审查文件	1、临历	末研究方案(周	反本号: VE	RSION	1.0_201	21106);		
	2、例封	报告表(版本号:	VERSION 1	.0_2012	201109)			
	3、知1	青同意书;						
	4、研3	究者专业履历及	支 专业科室	人员配名	备、设备	设施情况;		
	5、研究	究者手册;					有口	无■
	6、其作	也伦理委员会对		目的决定	Ĕ;		有■	无口
审查内容	研究者的资格: 符合要求■ 不符合要:				要求□	求□		
	人员配备: 符合			要求■ 不符合要求□				
	设备条件: 符合			予要求■ 不符合要求□				
	知情同意书: 符合要求■					不符合图	要求□	
	获取知	情同意书的方法	去:恰当■			不恰当日		
	研究方	案:	符合要	求■		不符合里	要求□	
	受试者	因参加临床试验	验 有有效	抢救措	施■	无有效技	仓救措施口	
	发生不	良反应或意外:	有补偿	规定■		无补偿规	见定口	
审查意见	同意	作必要的修改	位后同意	作必要	更的修正	后重审	不同意	终止或暂停已
								批准的研究
	3人	5		0			0	0
出席人数	应到:	9人	实到: 9人		回避:	1人(投票	请假: C	人
					时)			
审批意见: 组	经审查该	项目临床试验	方案符合要	求,请对	知情同	意书进行修	8改,具体	为:在"知情同
意书.知情告	知页"中	明确告知受试	者,对照组	为"安	慰对照	台疗"。做出	出上述修改	收后,同意开展临
床研究。	13							
主任委员签等	字: 1(7	772	会议记录	者签字	: tvi)·d).	联系电话	舌: 029-87251691

Institutional Review Board of Shaanxi Province of Traditional Chinese Medicine

Ethics Review Approval

Project Title	The Effect and Safety of Electro-acupuncture for Women with Pure Stress Urinary Incontinence: a								
	Multicente	Multicenter, Randomized Controlled Trial							
Applicant Site	Acupuncture and Moxibustion Department, Shanxi Province Hospital of TCM								
Project Sponsor	National I	National Key Technology R&D Program during the Twelfth Project 2012BAI24B0						2012BAI24B01	
	Five-year I	Plan Period of Ch	ina					No.	
Department	Acupunctu	re and	Site PI		Tongshe	ng Su		Title	Chief Physician
	Moxibustic	on Department							
Meeting Date	Feb 1, 2013	3	Meetin	g Place	Confere	nce		Review	Meeting Revies
					Room	of	the	Type	
					Hospital				
Review Files	1. study pro	otocol (VERSION	N1.0_201	21106);					
	2. Case Re	port Form (VERS	SION1.0_	2012110	9);				
	3. Informed	d Consent;							
	4. Research	4. Researchers' CV, Personnel Allocation, Equipment and Facility;							
	5. Researchers' Handbook Yes □ No ■								
	6. Decision	by Other IRB		Yes	No □				
Review Contents	Researchers' Qualification: Meet Requirement Not				Meet Requi	rement \square			
	Personnel A	Allocation:]	Meet Re	quirement		Not	Meet Requi	rement \square
	Equipment	and Facility:]	Meet Re	quirement		Not	Meet Requi	rement \square
	Informed C	Consent:]	Meet Re	quirement		Not	Meet Requi	rement \square
	Informed C	Consent Obtained:	: .	Appropriate ■ Not Appropriate □					
	Study Prote	ocol]	Meet Requirement ■ Not Meet Requirement □			rement \square		
	Subjects' A	E or Accident:	,	With Eff	ective En	nergen	icy N	leasures	
			,	Without	Effective	Emer	gency	Measures	
			,	With Co	mpensatio	n 🔳			
			,	Without	Compensa	ation			
Review Comments	approved	approved after r	evision	review	after revis	ion	not	approved	Terminated/Paused
	3	5		0			0		0

Institutional Review Board of Shaanxi Province of Traditional Chinese Medicine

Attendance People	Anticipated: 9	Participated: 9	Avoided: 1	Left: 0			
Approval Comments: The study protocol of this trial conforms to the requirements, but the informed consent need							
s to be revised. Subje	s to be revised. Subjects should be informed that the control group uses plocebo treatment. The study will be appr						
oved after revision.	oved after revision.						
Director Signature:		Meeting Recorder:		+86 029 87251691			
Guangyang Wei		Yifei	Zhao				
Date: Feb 1, 2013 Dat		e: Feb 1, 2013					

伦理审查批件

	, –	T 1 T 110 1 1				
项目名称	电针治疗女性单约	屯性压力性尿失禁有效	性和安全性多中心随机对照试验			
项目编号	2012EC007 项目来源 "十二五"国家科技支撑计划					
牵头单位	中国中医科学院广安门医院					
申办者(如有)	山西中医学院中西	西医结合医院	6			
主要研究者	王杰 高素云 赵文兵					
审查类别	初始审查 审查方式 会议审查					
审查日期	2013.2.25	审查地点	医院五楼会议室			
审查委员	樊东升,王毅东,	高继宁, 蔺涛, 刘红	[玲,田成瑛,闫荔			
批准文件	研究方案(VERS	ION1.0_20121106) ,\$	和情同意书(V1.0)			
审查意见	研究方案(VERSION1.0_20121106),知情同意书(V1.0) 根据卫生部《涉及人的生物医学研究伦理审查办法(执行)》、国家药品食品监督管理局《药物临床试验质量管理规范》、《医疗器械临床试验规定》、《药物临床试验伦理审查工作指导原则》国家中医药管理局《中医药临床研究伦理审查管理规范》以及《赫尔辛集宣言》和国际医学科学组织委员会颁布的《人体生物医学研究国际道德指南》的伦理原则,经本伦理委员会审查,同意按所批准的临床研究方案、知情同意书、招募材料开展本项研究。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。 如发生严重不良事件以及影响研究风险收益比的非预期不良事件,请申请人及时提交严重不良事件报告。 请按照伦理委员会规定的粘度/定期跟踪审查频率,申请人在截止日期前1个月提交研究进展报告;定期向牵头单位伦理委员会提交研究进展的汇总报告。					
有效期	2013年3月1日-	2014年2月28日				
联系人与联系电话	蔺涛 0351-2621:	527				
主任委员签字			地 重医结合。			
		山西中医学院中西医	结合医院伦理委员会(医务科代章)			

日期: 2013年02月25日

Institutional Review Board of Shanxi Hospital of Integrated Traditional and Western Medicine

Ethics review Approval

Project Title	The Effect and Safety of Electro-acupuncture for Women with Pure Stress Urinary				
	Incontinence: a Multicenter, Randomized Controlled Trial				
Approval No.	2012EC007	Project Sponsor	the 12 th Five-year Plan of the National		
			Key Technology Support Program by		
			the Ministry of Science and Technology		
			of the People's Republic of China		
Leading Organization	Guang'anmen Hospita	l of China Academy o	f Chinese Medical Sciences		
Applicant (if any)	Shanxi Hospital of Inte	egrated Traditional and	d Western Medicine		
Site PI	Jie Wang, Suyun Gao,	Wenbing Zhao			
Review Attribute	Initial Review	Review Methods	Review conference		
Review data	Feb 25, 2013	Review Place	Conference Room, the 5 th Floor of the		
			Administrative Building of the Hospital		
Review Committee	Dongsheng Fan, Yidong Wang, Jining Gao, Tao Lin, Hongling Liu, Chengying Tian,				
	Li Yan				
Approved Files	Study Protocol (VERSION 1.0_20121106), Informed Consent (V1.0)				
Review Comments	According to "ethical r	review methods for bi	omedical study involving human subjects"		
	issued by the Ministry of Health, "Good Clinical Practice", "Provisions for Clinical				
	Trials of Medical Device" and "Guidelines for Ethical Review Work of Drug Clinical				
	Trials" issued by Sta	ate Food and Drug	Administration (SFDA) of the People's		
	Republic of China, "r	management specifica	tions for ethical review of TCM clinical		
	studies" issued by Stat	te Administration of	Traditional Chinese Medicine, Declaration		
	of Helsinki, and "Inte	rnational ethical guid	delines for biomedical research involving		
	human subjects" mad	de by Council for	International Organizations of Medical		
	Sciences, through the	review of this IRB, t	the study protocol, informed consent, and		
	recruitment files of this	s research were approv	ved.		
	Please conduct this cli	nical study following	the GCP principles and the study protocol		
	approved by IRB. The	health and rights of the	he subjects should be protected throughout		
	the whole study.				

Institutional Review Board of Shanxi Hospital of Integrated Traditional and Western Medicine

	An application should be submitted if there are major revisions in principle					
	investigator, study protocol, informed consent, or the recruitment files.					
	A report of the severe adverse events (SAE) should be submitted in time if any SAE or					
	any other un-anticipated adverse event, which will affect the risk-reward ratio of the					
	study, occurs.					
	Researchers should submit report of the study progress one month before the deadline					
	according to the ethical review frequency. A summary report of the study progress					
	should be submitted to the IRB of the leading site.					
	A final report should be submitted when the study is completely done or stopped					
	prematurely.					
Validity period	From Mar 1, 2013 to Feb 28, 2014					
Contact and Phone	Tao Lin +86 0351 2621527					
Director Signature	Dongsheng Fan					
	IRB of Shanxi Hospital of Integrated Traditional and Western Medicine (SEAL)					
	Date: Feb 25, 2013					