

ADDING KEY STUDY PERSONNEL

Please list key personnel being added to the study below (Investigator, Sub-Investigator, Lead Study Coordinator, Research Specialist) who are responsible for the design, conduct or reporting of this research study. Attach the following documents unless they are already on file in Hatton for each person: CV, CITI Completion Certificate, and Medical License. Contact Kelly Blackwell@trihealth.com if you need to confirm documents are in database.

Name & Degree	Role	Tasks	CITI Expiration	Other Training (optional)	Employer
Hame & Degree	(PI, Sub-I, etc.)	(see key	Date	GCP, FDA Info sheets or	Lingityei
	(11, 300 1, etc.)	below)	Date	the Belmont Report;	
		Delow)		HSR Seminar web-based	
				training; HSR Training by	
				Sponsor/CRO	
				sponsor/CRO	

- 1. Obtain Informed Consent
- 2. Reviewing Medical History
- 3. Perform Physical Exams
- 4. Reviewing for Inclusion/Exclusion
- 5. Drug Dispensing

- 6. Drug Accountability
- 7. Ongoing AE Assessment
- 8. Update/Maintain IRB docs
- 9. Data Analysis
- 10.Other -

Version: 1-23-19