

CBR SUBMISSION WORKBOOK

Version 04/01/19

The CBR Submission Process consists of completing the appropriate sections of this workbook and attaching specific study documents. Please see below for a list of submission types, required documents for each submission and workbook forms applicable for each submission type. The Bookmark feature to the left will navigate you through the various sections of the Workbook. Blue underline indicates that you will be taken directly to the form or link referenced.

Please note it is required that a CBR Submission Workbook be completed for ALL CBR submissions. The Billing Plan lists all protocol activities: 1) billable services (such as labs and radiology scans that are CPT-coded services billed through the UAB Health System) and 2) non-billable services (such as informed consent and concomitant medications that are study services without CPT codes). Each study activity from the protocol Schedule of Events/ Study Schema is required to be listed under the corresponding header in the Billing Plan. Please use the attached FAQs to assist with completing this Workbook.

CBR SUBMISSION CHECKLIST FOR EACH SUBMISSION TYPE

FEASIBILITY FEE SUBMISSION

Complete the forms specified below and E-mail the CBR Submission Workbook to fap@uab.edu by using the *Submit CBR Workbook to CBR* button on the CTAW

Form page.

CBR Submission Workbook- complete the [CBR Submission Form](#);

[CTAW Form](#).

FULL SUBMISSION

Complete the forms specified below in the CBR Submission Workbook and submit any applicable additional documents via E-mail to fap@uab.edu by using the *Submit CBR Workbook to CBR* button on the Billing Plan page.

CBR Submission Workbook, at a minimum complete the [CBR Submission Form](#) and [Billing Plan\(s\)](#);

Including additional Billing Plan spreadsheets is done by clicking the "+" button at the bottom of the Excel document, then copy and pasting the Billing Plan template into the new spreadsheets.

Study Protocol, final version. The UAB IRB Human Subjects Protocol (HSP) can satisfy this requirement if a protocol is not available;

Draft Consent(s);

Manual of Procedures, if available;

Sponsor Draft Clinical Trial Agreement, if available;

Sponsor Draft Budget, if available;

Sponsor Lab Manual (If UAB Laboratories will be doing any special processing/shipping for the study).

If applicable, please submit as indicated below:

UAB Laboratory Use - The [Lab Questionnaire](#) in the CBR Submission Workbook must be completed, if the UAB Hospital Laboratories are involved in the study (which includes TKC 2nd Floor Lab, Whitaker Lab, Outreach Lab and CRU blood draws sent to UAB Labs);

Flow Cytometry Use - The [Flow Cytometry Form](#) in the CBR Submission Workbook must be completed if the UAB Flow Cytometry lab will be involved in the study;

Clinical Research Unit (CRU) Use - The CRU award letter, when available.

*****If the study involves Radiology procedures/services, please contact Radiology Research at radresearch@uabmc.edu.***

DEVICE TRIAL SUBMISSION

(Includes all devices including, but not limited to: Cat. A, Cat. B, Post Market, Pre-Market, 510K Summary and Non-Significant Risk Devices)

Complete the forms specified below in the CBR Submission Workbook and submit any applicable documents via E-mail to fap@uab.edu by using the *Submit CBR Workbook to CBR* button on the Billing Plan page.

CBR Submission Workbook, at a minimum complete the [CBR Submission Form](#), [Device Form](#) and [Billing Plan\(s\)](#);

Including additional Billing Plan spreadsheets is done by clicking the "+" button at the bottom of the Excel document, then copy and pasting the Billing Plan template into the new spreadsheets.

Study Protocol, final version. The UAB IRB Human Subjects Protocol (HSP) can satisfy this requirement if a protocol is not available;

Draft Consent(s);

Manual of Procedures (If available);

The Department of Health and Human Services (FDA) memo indicating device type and device #;

Draft Sponsor Clinical Trial Agreement;

Draft Sponsor Budget;

Reimbursement manual/Implant procedure coding from the sponsor (if available);

Sponsor Lab Manual (If UAB Laboratories will be doing any special processing/shipping for the study).

If applicable, please submit as indicated below:

UAB Laboratory Use - The [Lab Questionnaire](#) in the CBR Submission Workbook must be completed, if the UAB Hospital Laboratories are involved in the study (which includes TKC 2nd Floor Lab, Whitaker Lab, Outreach Lab and CRU blood draws sent to UAB Labs);

Flow Cytometry Use - The [Flow Cytometry Form](#) in the CBR Submission Workbook must be completed if the UAB Flow Cytometry lab will be involved in the study;

Clinical Research Unit (CRU) Use - The CRU award letter, when available.

*****If the study involves Radiology procedures/services, please contact Radiology Research at radresearch@uabmc.edu.***

AMENDMENT SUBMISSION

Complete the forms specified below and E-mail with the additional required documents to fap@uab.edu by using the *Submit CBR Workbook to CBR* button on the Billing Plan page. Note that Amendments should be submitted for CBR review *ONLY* if there are certain changes:

1) Additions/deletions made involving clinical billables; 2) Change in clinical billable service location or 3) Change in PI.

CBR Submission Workbook, at a minimum complete the [CBR Submission Form](#) and [Billing Plan\(s\)](#);

The amended Billing Plan(s) should be the last FAP-Approved Billing Plan(s) with the addition of colored highlight indicating ALL changes being made by this amendment (if changes involve billable activities). Including additional Billing Plan spreadsheets is done by clicking the "+" button at the bottom of the Excel document, then copy and pasting the Billing Plan template into the new spreadsheet.

NOTE: Please create a new Billing Plan spreadsheet for EVERY new amended Billing Plan (each arm/cohort) related to this amendment and any future amendments. All study Billing Plans should be in one Excel document in chronological order from the initial FAP-Approved Billing Plan(s) through to the current amendment in the CBR Submission Workbook.

Amended study protocol, if study changes are in the protocol document (UNTRACKED VERSION);

Summary of Changes (SOC) Document. If amendment changes are not specific protocol changes, then provide a complete listing in your E-mail submission to CBR (i.e. Change in PI or change in billing designation of existing activity at existing visit, etc.).

If applicable, please submit as indicated below:

UAB Laboratory Use - The [Lab Questionnaire](#) in the CBR Submission Workbook must be completed: 1) If there are NEW UAB Hospital lab services added OR 2) Previous labs occurring at new time points that were not reflected in the last FAP-Approved Billing Plan. The UAB Hospital Laboratories include: TKC 2nd Floor Lab, Whitaker Lab, Outreach Lab and CRU blood draws sent to UAB Labs;

Flow Cytometry Use - The [Flow Cytometry Form](#) in the CBR Submission Workbook must be completed: 1) If there are NEW flow cytometry lab services added OR 2) Previous flow cytometry labs occurring at new time points that were not reflected in the last FAP-Approved Billing Plan;

Clinical Research Unit (CRU) Use - If the CRU award letter was amended as a result of this amendment, then CBR needs a copy of this letter as soon as possible.

*****If the study amendment adds Radiology procedures/services not previously performed in the study, please contact Radiology Research at radresearch@uabmc.edu.***

UAB CLINICAL BILLING REVIEW (CBR) SUBMISSION FORM

Project Title:

Protocol Brief Title/Short Title:

Protocol #:

Sponsor Name (Funding Source):

Sponsor/Funding Type:

Federal

Investigator-Initiated

Academic Collaboration

Other:

Industry

Cooperative Group

Principal Investigator (PI):

PI E-mail: (johndoe@uab.edu)

Assigned Study Coordinator:

Assigned Study Coordinator E-mail: (johndoe@uab.edu)

Department/Division:

Submitter:

Submitter E-mail: (johndoe@uab.edu)

Submitter Phone Number:

Clinical Services will be provided in what setting? Mark all that apply.

Hospital Inpatient Outpatient CRU Inpatient
CRU Outpatient

Anticipated Study Start Date: (MM/DD/YYYY)

Estimated number of participants to be enrolled at UAB: (numeric response)

IRB # assigned to this study:

Is this study registered on ClinicalTrials.gov currently or will it be in the future?

Yes No If Yes, please provide the NCT number assigned to this study or

indicate "PENDING" if not yet received:

Is this study conducted under an IND or Device application reviewed by the FDA?

Yes No If Yes, please provide the FDA# (IND#, IDE#, K#, P#, etc.)

assigned to this study or indicate "PENDING" if not yet received:

Has this study been granted an exemption from being required to have an FDA IND?

Yes No

If Yes, please provide documentation to CBR confirming this exemption.

Please select the correct submission type below, include all required documents, and submit to fap@uab.edu via the buttons on one of the last two pages of this Workbook. CBR review will begin once ALL required documents are received.

SUBMISSION TYPE:

Previous FAP# assigned for a Feasibility Fee Submission and/or Full Submission associated with this study:

LAB QUESTIONNAIRE
HOSPITAL LAB RESEARCH PROTOCOL REVIEW QUESTIONNAIRE

(Use this form when research activities involve the use of UAB Hospital Lab Systems.)

1. How many participants do you plan to enroll in this study?

2. What is the planned study start date and study close date?

3. Are you using TKC or UAB Hospital/Outreach Laboratories for resulting local labs? If so, please designate all that apply.

TKC UAB Hospital Lab Both

4. Will you require venipuncture services through TKC or UAB Hospital Labs?

Yes No

5. Are ALL the labs that are resulted at the TKC/UAB Hospital Lab billed to research study?

Yes Some labs are billed to study No - None are billed to study

COMPLETE INFORMATION BELOW ONLY FOR CENTRAL (SPONSOR-SPECIFIC) LAB TESTING/PROCESSING/SHIPPING FOR THIS STUDY:

6. What lab facility will be used?

TKC UAB Hospital Labs CRU
CCTS

7. Will the study require venipuncture, processing, shipping or storage through TKC/UAB Hospital Labs? Mark all that apply.

Venipuncture Processing Shipping Storage

8. Will the study require routine lab testing?

Yes No

9. Will the study require after hours, weekends or holiday processing?

Yes No

10. Will the study require complexity testing (such as buffy coat)?

Yes No

11. Are stool specimens involved?

Yes No

FLOW CYTOMETRY FORM
UAB HOSPITAL LABS - FLOW CYTOMETRY LAB
PRELIMINARY RESEARCH LAB TESTING QUESTIONNAIRE

* This form must be completed and submitted no later than 2 weeks before research protocol testing begins.

1, Cluster Designations (CD) to be reported (Please specify particular subsets. CD3+CD4 + % of lymphocytes and CD3+CD8+ % of lymphocytes, not just CD3, CD4, CD8)

NOTE: In order to perform Flow Cytometry, results from CBC/Diff are required.

2. Estimated patient volume:

Number of Subjects:

Number of Specimens Per Subject:

Duration of Study (Months specimens will be received):

3. Attach protocol and reference the protocol page(s) specific to Flow Cytometry:

DEVICE FORM
Submission Form for Hospital Review of Device Trials

Purpose: UAB Hospital, The Kirklin Clinic, and other UAB Medicine clinical facilities support the conduct of Device trials in their facilities. To ensure UAB Medicine facilities appropriately submit claims for these services, please furnish the following information to the UAB Clinical Billing Review Office (FAP).

1. Type of device (attach FDA letter):

Category A	Category B
510K Summary	Non-Significant Risk
FDA-Approved	Wearable Device
Post-Market Appr. - Carotid Stenting	Pre-Market Approval
Other	

2. Name of Device:

3. Specify the venue/location where the procedure to implant the device will be performed:

TKC	UAB Hospital	UAB Highlands
Children's Hospital		
Other		

4. Provide the specific procedure area for your response in Question # 3 (i.e. OR, HVC, etc.):

5. Financial arrangements for the device and related procedure (5a-f):

a. Is the sponsor furnishing/providing the device at no charge?

Yes No

If No, what is the amount being charged?

b. Provide the name, number and E-mail for the sponsor contact responsible for negotiation of the device purchase agreement and/or the price for the device.

c. Who is paying for the procedure to implant the device?

Sponsor	Insurance (SOC)
Other	

d. What is the volume of patients expected to participate in the trial at UAB?

e. Specify the name of the procedure(s) related to the device implant:

f. Provide a list of patients who have had this procedure performed in the last 6-9 months. This includes names, MRNs and dates of service when the procedure was performed.

CLINICAL TRIALS ACTIVITY WORKSHEET (CTAW) FORM

The CTAW Form Template is attached to this document. Once you have completed the CTAW Form for this study submission, please save the new document and use the buttons below to:

- 1) Upload CTAW Form and any additional documents.
- 2) Submit this study submission for CBR review.

BILLING PLAN

The Billing Plan Template is attached to this document. Including additional Billing Plan spreadsheets is done by clicking the "+" button at the bottom of the Excel document, then copy and pasting the Billing Plan template into the new spreadsheet. Once you have completed the Billing Plan(s) for this study submission, please save the new document and use the buttons below to:

- 1) Upload Billing Plan(s) and any required documents.
- 2) Submit this study submission for CBR review.