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Introduction

Welcome to the Vanderbilt University Medical Center’s Institutional Review Board’s new electronic submission system called DISCOVR-e (the Data Integrated Study Console of Vanderbilt’s Research Enterprise). Phase I allowed researchers to access basic study information, as well as their human subjects training/continuing education status. Phase II allowed researchers to complete an IRB submission online, including the ability to use electronic signatures. Every type of submission previously sent in by hard copy (e.g., new study, continuing review, adverse event, etc.) could be submitted with a few clicks of your mouse.

Another benefit of the DISCOVR-e portal allowed researchers to view their approved documents online, including Committee Action Letters (CALs), Final Approval Letters (FALs), and approved and date stamped consent forms.

Phase III of this project includes an intuitive investigator dashboard and the implementation of an IRB application wizard that incorporates supplemental form information like vulnerable populations in research, repositories, radiological procedures for research, and waiver into one cohesive document.

Future phases may include additional features and capabilities as the system grows and users provide the development team with feedback and suggestions. The IRB plans to continuously monitor the system to make enhancements and improvements where needed.

Overview

The DISCOVR-e portal is a web-based system. You can log in and submit documents or check the status of a previous submission anywhere in the world where you have a connection to the Internet. This section of the manual will give you an overview of the system and provide highlights of its capabilities.

To access the system, please visit the Vanderbilt IRB website at https://www.vumc.org/irb/ and click on the DISCOVR-e link on the right-hand section of the page. When prompted, enter your VUnetID and e-password. That’s all it takes!!

After logging in, you will see your Investigator Dashboard. This is your home page which shows all of your outstanding action items and approved studies. The following pages will provide step-by-step screenshots for creating and responding to outstanding submissions, as well as navigation tips for the site. We plan to provide video tutorials following the rollout as a secondary training tool for the research community so check back often!
Log into the System

You can access the system by going to the IRB website, https://www.vumc.org/irb/ and clicking on the DISCOVR-E link.

You can also access the system by typing the following internet address into your browser: https://irb.mc.vanderbilt.edu/. You may want to save this address in your list of favorites/bookmarks. This will bring you to the Login page. To enter DISCOVR-e, enter your VUnetID and e-password.
Investigator Dashboard Overview

This is your Investigator Dashboard. When you log in, this is the first screen you will see. From here, you can access all of your studies and view submissions that require action from you.

The different colored buttons across the top of the page show items requiring action from you.
Notice the number in the top, right corner of each button. This number indicates how many items are requiring that particular action. The first button (green) is the Site visit request button. The number in the corner indicates how many site visits have been requested by the IRB.

The second button (yellow) is the pre-review requests button. The number in the corner indicates how many submissions require a pre-review response. Clicking this button will allow you to access these studies.

The next button (blue) is the Submissions Requiring My Signature button. The number in the corner indicates how many submissions require your signature before moving forward. Clicking this button will allow you to access these studies.

The next button (red) is the Submissions Requiring a CAL Response button. The number in the corner indicates how many submissions require a response to a CAL. Clicking this button will allow you to access these studies.
The final button (purple) is the Draft Submissions button. The number in the corner indicates how many submissions have been started but have not been completed or sent to the IRB. Clicking this button will allow you to access these studies.

Lists of approved studies are located below the buttons for quick access. My studies is the default view on the dashboard. This view shows studies in which you are listed as PI, Study Contact, or Faculty Advisor.

Studies listing me as KSP will display only studies in which you are listed as other KSP (not PI, Study Contact, or Faculty Advisor).
These lists are automatically sorted by Study Expiration. If a study is within 8 weeks of expiration, it will appear highlighted in red. Lists can be sorted by Study Title, IRB Number, PI, and Study Contact by clicking on the heading.

You can also search for studies in these lists by entering study identifiers such as PI or Study Contact name, parts of the study title, or IRB number into the search box on the right.
The views accessed on the Dashboard can also be accessed at any point via drop down menus. These drop down menus can be accessed on any screen, allowing you to access submissions without returning to your dashboard.

Clicking on Dashboard will return you to your Investigator Dashboard.

Clicking Submissions will display a drop down menu. This menu holds items corresponding with the buttons on the dashboard, as well as other study views.

Drafts corresponds with the purple Drafts button.
Requiring my signature corresponds with the blue Submissions requiring my signature button.

Pre-review requests corresponds with the yellow Pre-review requests button.

CAL responses corresponds with the red Submissions requiring a CAL response button.

This dropdown also holds views for Submissions in route and Submissions I’ve signed. Clicking Submissions in route will display a list of submissions pending signature from another person.
Clicking Submissions I’ve signed will display a list of all submissions signed by you.

Clicking Submissions not yet approved will display a list of all submissions you have created and where you are listed as PI, study contact, or faculty advisor that have been submitted to the IRB and have been signed, but have not yet received approval.

Clicking Studies will display another drop down menu. This menu holds items corresponding with the studies lists on the Dashboard, as well as Inactive studies.
My Studies corresponds with the My Studies button.

Studies listing me as KSP corresponds with the Studies listing me as KSP button

Clicking Inactive will display a list of your inactive studies.
IRB Training expiration can be easily viewed on the Investigator Dashboard using the gauge displaying days until expiration.

This view displays your IRB Training status and expiration date.
If you submitted a training certificate after (date), your history will contain the certificate. Clicking on the document link will allow you to download the certificate.

To look up a training certificate for another member of KSP, enter the name or VUNET ID into the Look up another user textbox. Their IRB Training Summary will appear and will have the same features (expiration date, downloadable certificate, etc.).
New study submission are the only submissions created from the Investigator Dashboard. Clicking the green Create a new study button will begin the process of creating that new study submission.

Throughout the website and application, you will find small black circle icons with an ‘i’ inside. These are information buttons. Hovering your mouse over these icons will display helpful information about that item. For example, hovering over this icon next to the Approved Studies heading displays information regarding the use of the Approved Studies view.
Submitting a New Study

Submitting a new study through the DISCOVR-E portal does not require any special training. The following guide is a basic set of instructions and illustrations to help you in the submission process. If you have not logged into the system yet, please follow the instructions provided above.

Click on the “Create a New Study” button on the right side of the page to start the process.

After selecting the “Create a New Study” button, you will be prompted to begin your new study submission in eSMART. This feature provides a central location to view the status of your study’s approval/regulatory requirements. To proceed in eSMART, click the “Start in eSMART” button. To bypass this feature, click the “Continue New Submission” button.
You will begin your new study submission by typing in a title for the study. Next, enter the Principal Investigator (PI) for the study. You can search for the Investigator by first and/or last name or even do a partial name search. Click the name of the PI you want to use. If you are not able to find the Investigator in the database, save your work and contact the IRB at 322-2918 for assistance.

Indicate whether or not the Principal Investigator is a student. If the answer is “yes,” you will be prompted to identify a Faculty Advisor. Selecting a Faculty Advisor follows the same process as choosing a Principal Investigator.

If applicable, select a Study Contact by choosing the appropriate contact person from the database. The process for indicating a study contact is the same as outlined previously for selecting a Principal Investigator or Faculty Advisor.

All new study submissions require a department chair and/or a division chief signature. The process for indicating this individual is the same as previously outlined. If you experience trouble identifying the department chair and/or a division chief for your research, save your work and contact the IRB at 322-2918 for assistance.

Click on the “Create New Study” button at the bottom of the page to advance in the new study creation process.
Selecting and Managing Key Study Personnel

Under "My studies", Click on the title of study you wish to add KSP.

NOTE: If a KSP’s requirement for annual VU IRB Human Subjects Training is not current, the individual may not be added to the KSP listing at this time. Those individuals may find information on how to update training at https://www.vumc.org/irb/education-and-training.
The PI, Study Coordinator, and Faculty Advisor, if applicable, identified in the “Info” tab will be the first to populate into the Key Study Personnel (KSP) tab.

Under the “Current KSP” tab, begin typing the name, email, or VUnetID in the “Add another” search field. You will be required to enter credentials, pager number, role in project and whether the KSP will be accessing Protected Health Information. These fields must be completed in order to advance to the subsequent sections of the Application Wizard. Repeat this process until you have identified all of the individuals that will help in the conduct of your research.

Hit the “Save KSP’s” button to save your changes.

To remove a person from the list of KSP, click the “trash” icon next to the VUnetID of that person’s name. The individual will be removed from your KSP listing.
Click the “restore” icon to restore a person’s name. If you do not have any other KSP to add/delete, please click the “Save KSPs” button and proceed to the next section.

The history tab allows you to see the date KSP were added/removed. You can also see who added/deleted KSP.
Adding a Funding Source

Indicate whether the study is funded by external support, VICTR funding support/use of VICTR facilities, internal funds or not funded.

Next, specify the study's funding source by clicking the "Add Funding Source" button.

To add a funding source for your study, select the funding type within the dropdown.
Add the funder for your research and select "Accept".

Your funding source will be found under "Funding Sources". You may click the "pencil" icon to edit your funding source or the "trashcan" icon to delete it.
Once all funding-related questions have been completed and you are ready to proceed with the creation of your new study submission, click the “Save and Continue” button at the bottom of the form.
Completing the Performance sites tab

The Performance Sites tab will hold all of the information about other sites involved in the study that are:

1) Participating sites in a Single IRB Study for which Vanderbilt is the reviewing IRB, and  
2) Performance sites for which the Vanderbilt PI is responsible that are either engaged or not engaged in research.

*Once a study is approved, this information is managed differently. Please see the Site Add section of this manual for further instructions on how to manage performance sites after initial study approval.

If the study is a Single IRB Study for which Vanderbilt is the reviewing IRB, change the answer to the question in the IRB of Record box to Yes.
This will open the Single IRB Performance Sites table.

To add a Single IRB Performance Site, click Add a Site.
Begin typing the name of the site in the box labeled Site.

As you begin typing, options of sites that are already in the system will appear. Select the site from these options.

If the site you would like to add is not included in the drop down list, move to the section below Single IRB Performance Sites labeled Requested Sites. In this text box, include the name of the site.
Indicate which site is the lead site for the study by clicking the check box next to that site on the Single IRB Performance Sites table.

<table>
<thead>
<tr>
<th>Site</th>
<th>Lead Site</th>
</tr>
</thead>
<tbody>
<tr>
<td>Johns Hopkins University School of Medicine</td>
<td></td>
</tr>
<tr>
<td>Duke University Health System</td>
<td></td>
</tr>
<tr>
<td>University of Alabama at Birmingham</td>
<td></td>
</tr>
<tr>
<td>Vanderbilt University Medical Center</td>
<td></td>
</tr>
<tr>
<td>Vanderbilt University</td>
<td></td>
</tr>
</tbody>
</table>

Note: If Vanderbilt University Medical Center or Vanderbilt University are participating in the study, they **WILL** need to be included in this performance sites table.

To remove a site from this table, click the Remove button.
To add a Coordinating Center, change the response for the coordinating center question from No to Yes and enter the appropriate information in the boxes that appear.

To add Performance Sites for which the Vanderbilt PI is responsible, click Add a Site under the Performance Sites section.

Enter the name of the site
Indicate if the site is engaged or not engaged in research

And indicate if the approval letter or letter of cooperation for that site is attached with this submission or if the letter is pending and will be submitted at a later date

To remove a site once you have added it to the table, simply click the Remove button next to that site.
Completing Your “Wizard” Application

You will begin your “Wizard” application by answering the questions found in each application tab. Your response(s) will prompt additional selections to populate regarding your submission. As you work through your submission, you will notice a progress bar building across the top of the page.

You can click any of application headings on the left side of your screen to go back and view or edit your submission.

Once all of the tabs appropriate for your research are complete, you have finished your “Wizard” application. Incomplete submissions can be saved and returned to later. These submissions can be accessed via the drafts submission button on your Investigator Dashboard. **Note: The submission portal will not allow an incomplete submission to be submitted or routed for signature.**

You may choose to download an electronic version of the IRB Application by clicking the “Download Application” button as shown in the example above.

To proceed to the next section of the submission process, click the “Continue” button.
Document Uploads

The Documents Uploads section is where you will attach/upload additional study documents such as consent forms, protocols, questionnaires, and any other study related documents or materials. Uploading documents is easy! You may upload/attach documents from locations saved on your computer by dragging and dropping the item(s) into the proper field or by browsing your computer for the appropriate file and attaching the document the standard way. To utilize the drag and drop feature, click the “Drag and Drop” button to activate it. Next, select the document you wish to upload and drag it to the proper field and release the document. Multiple uploads may be dragged and dropped at a time if they will be uploaded to the same field. Otherwise, attachments must be uploaded one at a time. As you load additional items, a list will build showing what has been attached to your submission.

To upload/attach a document the standard way, click the “Standard” button to activate it. Click the “Choose File” button under the “Attachment” section to search your computer for the item you would like to attach. Select the item and then click the “Open” button. The item you selected will be automatically attached to your submission for review. Next, indicate the type of study document by clicking the one of the choice from the dropdown menu. Once the type of study has been selected, click “Upload” to attach the document to your submission. Attachments should be uploaded one at a time. If you have a group of consent forms, you may zip them and attach the zip file as one attachment. However, we ask that you do not provide a zip file for all of your study documents.
If you would like to delete an attachment, click the box next to item you wish to remove then click the “Delete Selected Files” button.

To continue with submission process, click the “Continue” button.

Your submission is now ready to be submitted. Click the “Submit” button to route your submission for the appropriate signatures.

As other signatures are obtained, the submission is automatically routed through the submission portal and ultimately ends its journey at the IRB. Once it arrives at the IRB, the submission will be assigned an IRB number and routed to one of the teams for review.
How to Retract a Submission

You may choose to retract a submission if you wish to make additional edits, or if you wish to delete the submission all together. Please note, you can only retract submissions that have not been formally submitted to the IRB.

Only once a submission has been routed for additional signatures, such as those from a department chair and/or a division chief signature, shall it be retracted. To retract the submission after it has been signed off on by the PI, select the “Submissions” dropdown menu at the top of the screen and click the “Submissions in Route” button.

Locate the submission you wish to retract by selecting the study title of that submission.
Click the “Retract Submission” button and then select “OK” on the confirmation prompt to continue with the submission retraction process.

Your submission should now be located in your “Drafts Submissions” view within your Investigator’s Dashboard.
If you need to revise any information within your submission, navigate to the appropriate tab of your submission and update where necessary.

If you would like to delete your submission, simply click the “Delete” button.

After making your edits, select the “Save and Continue” button to save your changes and to advance you back to the “Submit” tab. Click the “Submit” button to re-route your submission for the appropriate signatures.
How to Respond to a Pre-review

The Principal Investigator and Study Contact will receive an email when a pre-review request has been received. The link in this email will lead to the submission. These submissions can also be accessed on the dashboard. The pre-review requests button will contain a counter listing the number of items with pre-review requested changes.

The pre-review requests are sorted by two categories: *My Submissions* (studies in which you are the PI or Study Contact) and *Submission for studies listing me as KSP*.
Select the study title for the submission you wish to respond to. The pre-review tab in the study submission will list general issues and comments. A summary of changes requested to the IRB application will also appear.

Select the application tab to respond to pre-review suggestions. All sections of the application with suggested changes will be highlighted. Select each section to view the specific items with change requests.
The pre-review comment will appear below the item. To revise an answer to align with the request, simply select the appropriate response or revise the text in the dialogue box as necessary. As all pre-review comments are suggestions, please add a comment providing a rationale for any requested changes not made.

If revisions are necessary to study documents, attach any documents with pre-review requested changes in the document uploads tab. Please ensure that all changes are tracked. Documents can be uploaded by dragging and dropping the document into the corresponding box, or using the standard upload method to find the document on your computer.
Once all pre-review suggestions have been responded to, return to the pre-review tab. Review the requested changes and ensure that all necessary information has been included. Select the green button at the bottom of the tab to submit the pre-review response.

A dialogue box will appear giving you an opportunity to include comments to the analyst. Including comments is optional. To complete the pre-review response, select the blue button to re-submit to the analyst.

The pre-review response is now submitted. The submission will be processed and moved forward for review by the IRB.
Submitting an Amendment to an Approved Study

All amendments will be submitted using the same submission process. The administrative amendment and standard amendment forms will no longer be utilized. Changes to key study personnel no longer require the submission of an amendment.

To initiate an amendment submission, select from the study in your dashboard and select the study that requires an amendment.

From the study page, click “Create Submission.”
This will open the study page. Using the *Create Submission* tab on the study page, select to create a new amendment submission.

![Create Submission tab](image)

Begin by completing the main page of the amendment. Save your changes at the bottom of the page before moving forward. Changes that are not saved before moving to a new tab may be lost.

![Main page amendment](image)

This tab is also used to change the Principal Investigator, Study Coordinator, or Faculty Advisor. This change must be made as an amendment after the individuals are added as Key Study Personnel.
Completing the Amendment Tab

Check the boxes for each document being revised or change being made as a part of this amendment. Additional questions will appear based on these selections. Please answer each item and save to continue the submission.

Making Changes to the IRB Application

In the IRB application wizard, review each tab and make changes as necessary. The changes made will be tracked under each item. Since the application builds based on the information input, some new questions may appear as you make revisions. Include answers to each new question that appears and save your changes before moving on to the next section of the application.
Attaching new or revised Documents

Please track changes to all currently approved study documents that are affected by the amendment. This will aid the review process. In the Document Uploads tab, upload any new or revised study documents by dragging and dropping each document into the corresponding category. The Other category is available for any documents that do not fit a specific category. Alternatively, you can change the upload method to a standard selection method in the right corner of this tab.

The Submit tab will notify you if any items are incomplete. This is a good time to review each section of the submission and ensure that all necessary changes have been made. When you are ready to move forward, select the submit button.
The amendment submission is now ready for the Principal Investigator’s signature. If you are the Principal Investigator, a page will appear listing the responsibilities of the PI. Please review these responsibilities. At the bottom of the page, sign the document by inputting your Vanderbilt password.

If the submission is being made by key study personnel, the amendment will route for the PI’s signature. The PI will receive an email notifying him or her that a submission is pending signature. The link in this email can be used to access the submission and signatory page. This submission can also be accessed for signature under the Submissions Awaiting my Signature item on the dashboard.

The amendment submission will now be routed for signature to the Faculty Advisor if applicable for your study. Each of these individuals will have an alert on his or her dashboard to notify that a submission is pending signature, in addition to receiving an email notification. This item can be used to access the submission, review the amendment, and sign at the bottom of the page.
Retracting the Amendment Prior to Submission

If any signatories (PI, advisor, chair, or chief) would like to make changes before the submission is sent to the IRB for review, the amendment can be placed in draft mode by retracting it on the main page of the amendment. After any changes are made, the amendment can be submitted by signing at the bottom of the signatory page.

The Amendment is now submitted. You can track the progress of the amendment by accessing your submissions on your main Wizard page. You will receive a notification if a pre-review is necessary. Please see the Respond to a Pre-Review section of the manual for assistance in making your response.

Submitting an Amendment to Add Non-Single IRB Performance Sites

This will function the same way the amendment usually does. On the Amendment Form click Other, and in the box to provide the rationale for the amendment, include the Site Name, if the site is engaged or not-engaged in research, and if the approval letter is attached with the amendment or if it is pending.

To remove a site, indicate the site in this box.
Submitting a Continuing Review

Submit a continuing review for continuing studies or to close a study with the IRB. To start a continuing review submission, access your approved studies and select the appropriate study. Note that any studies with approval expiring in the next 8 weeks will be highlighted red.

From the study page, click “Create Submission.”

On the Create Submission tab, select to create a Continuing Review.
Select the **Continuing Review** tab to complete the Continuing Review wizard application. In the first item, select the study type. The corresponding continuing review application will begin to build based on this item.

The continuing review application for standard/expedited studies and repositories will ask for the current status of the research. Additional questions will appear based on this information. Provide an answer to each question and save your answers. Incomplete submissions can be saved and returned to later. These submissions can be accessed via the drafts submission button on your dashboard. The system will not allow an incomplete submission to be submitted or routed for signature.
Attach any necessary documents (i.e. Consent forms, Progress reports) for the continuing review in the Document Uploads tab. Submit both a stamped and clean copy of all consent documents for review and approval.

The Reviews tab allow you to review the currently approved study information. To make changes to this information, please see the amendment submission section of this manual.

Review all tabs for completeness and submit the continuing review in the Submit tab. The submission will now route for signature before being sent to the IRB for review.

The Continuing review has now been routed for signature. The IRB will receive the submission after the Principal Investigator signs the submission. Submissions that are awaiting signature (submissions in route) can be located in the submissions drop down menu at the top the screen.
Submitting an Adverse Event

On the Dashboard, Approved Studies can be found in the list at the bottom of the screen. Approved studies for which you are the PI, Study Coordinator, or Faculty Advisor are found on the My Studies tab. This is the default view for the dashboard. Studies for which you are listed as KSP are found on the Studies listing me as KSP tab.

If you see the study that the adverse event is related to, click the study title to go to the study page. If you do not see the appropriate study, you can use the “search” field by typing the IRB number, sponsor number, part of the title, or another relevant piece of study information.
From the study page, click “Create Submission.”

Click Adverse Event. Clicking this button will create the Adverse Event submission.
Complete the first form and click Save and Continue. This will move you to the Adverse Event tab. Click Save will save your information but will keep you on the Main tab.

On the Adverse Event tab, you will complete additional Adverse Event questions. Required items are marked with *must provide value*. Please complete all of the requested items on the form. The Wizard system will not allow incomplete forms to be submitted.

The following types of events are required to be reported to the IRB. If the event does not fit the categories shown, the event is not reportable to the IRB. However, the event may need to be reported to the sponsor. If you have any questions, please call the IRB (615-322-2918) or check with your sponsor.

```
<table>
<thead>
<tr>
<th>Question</th>
<th>Yes</th>
<th>No</th>
</tr>
</thead>
<tbody>
<tr>
<td>Was this an event that requires prompt reporting to the sponsor in accordance with the protocol (e.g., serious adverse events)?</td>
<td></td>
<td></td>
</tr>
<tr>
<td>* must provide value</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Unanticipated? (An event is &quot;unanticipated&quot; when it was unforeseeable at the time of its occurrence)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>* must provide value</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Serious? (An event is &quot;serious&quot; if it adversely alters the risk/benefit relationship of the research)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>* must provide value</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Related? (An event is &quot;related&quot; if it is likely to have been caused by the research procedures)</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
```
In this section, please indicate the Participant ID# or initials. **Please do not include a name or any other personal identifiers.**

<table>
<thead>
<tr>
<th>Participant Identifier:</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
</tr>
<tr>
<td><strong>NOTE:</strong> DO NOT include a name or ANY personal identifiers.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Participant Age:</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Identify Drug/Biologic/Device/Treatment/Intervention:</th>
</tr>
</thead>
<tbody>
<tr>
<td>* must provide value</td>
</tr>
</tbody>
</table>

Provide a detailed description of the event/problem including the timing of study treatment, dosing, or intervention. Also remember to include start and stop dates of relevant research interventions. Indicate the timing of the event/problem in relation to the research intervention. For example, 1) Research drug was started at 10:00; 2) Participant began wheezing at 10:15; 3) Drug stopped at 10:16; 4) Participant continued to have labored breathing and was treated with inhalers. 5) Respirations relaxed; 6) Breathing sounds clear by 11:00 and participant denied further complaints with breathing. 7) The sponsor was notified of the event/problem and the participant was withdrawn from further participation in the study.

<table>
<thead>
<tr>
<th>List 1-4 keywords describing the event/problem:</th>
</tr>
</thead>
<tbody>
<tr>
<td>(e.g., loss of confidence, nausea, vomiting)</td>
</tr>
</tbody>
</table>

Provide a description of the event/problem including the timing of the study treatment, dosing, or intervention with start and stop dates of relevant research interventions.

| * must provide value                          |

**This study (choose one):**
- has a Data and Safety Monitoring Committee Board (DSMC/DSMB) or Data Safety Monitor (DSM);
- does not have a Data and Safety Monitoring Committee Board (DSMC/DSMB) or Data Safety Monitor (DSM);
- unknown;
Once all required questions are complete, click the Save button at the bottom of the page.

![Image of a form with questions and options to select]

After saving, if you need to submit additional documents, click the Document Uploads tab at the top of the page. If no other documents need to be submitted, click the Submit tab.

![Image of a form with options to upload documents]
On the Document Uploads tab, documents can be submitted with either the Drag and Drop option, or the Standard option.

When all documents have been uploaded, click the Continue button at the bottom of the page. This button will bring you to the Submit tab.
If the Adverse Event form is not complete, you will see this error message on the Submit tab. Click the Adverse Event tab to complete the missing information, click save, and then click the Submit tab again.

Once the required information is complete, you will be able to click the Submit button on the Submit tab.

After a signature from the PI, the submission is automatically routed through the Wizard and ultimately ends its journey at the IRB where it will be sent to a team to start the review process.
Submitting a Non-Compliance with the Protocol

On the Dashboard, Approved Studies can be found in the list at the bottom of the screen. Approved studies for which you are the PI, Study Coordinator, or Faculty Advisor are found on the My Studies tab. This is the default view for the dashboard. Studies for which you are listed as KSP are found on the Studies listing me as KSP tab.

If you see the study that the non-compliance with the protocol is related to, click the study title to go to the study page. If you cannot find the appropriate study, you can use the “search” field by typing the IRB number, sponsor number, part of the title, or another relevant piece of study information.
From the study page, click “Create Submission.”

Click Non-compliance with Protocol. This will create the non-compliance with the protocol submission.
Enter the description of the Non-compliance and the date of the event or problem. Click Save.

Once you have saved the Main tab, click the Non-Compliance tab to continue.
Complete all questions on the Non-Compliance form. The Wizard will not allow you to submit if a question is not answered. Once all questions have been answered, Click Save at the bottom of the page.

If you have additional documents to upload, click the Document Uploads tab. If you have no additional documents to upload, click the Submit tab.
On the Document Uploads tab, upload additional documents using either the Drag and Drop or Standard method.

Once all documents are uploaded, Click Continue.
On the Submit tab, if the required information is not complete, you will get an error message. Click on the Non-Compliance tab to answer incomplete questions.

Once the form is complete, click the Submit button.

After a signature from the PI, the submission is automatically routed through the Wizard and ultimately ends its journey at the IRB where it will be sent to a team to start the review process.
Submitting a Response to a Committee Action Letter (CAL)

On the dashboard, the Submissions Requiring a CAL Response button will display a number indicating the number of outstanding items awaiting a response from you.

Clicking this button will show you a list of these outstanding items.

If the title is too long, the title will be truncated. Hover the cursor over the truncated title to display the entire title.
Select the appropriate response item.

From the Review Screen there are three options to download the CAL:
1. Click the Download CAL button above the letter
2. Click the download button in the PDF viewer

3. Click the Download CAL link at the bottom of the page under Submission Reviews
Once the CAL is downloaded, review the changes and concerns expressed by the Committee in the letter. Changes requested within the Wizard can be addressed by clicking the tab corresponding with the concern.

Please limit your changes to only those requested by the Committee. If the IRB has not requested any specific changes within the Wizard, simply click the Document Uploads tab, which is the tab where you can upload your cover letter to address each change requested by the Committee and any revised documents.
Please be sure to update the revision date and track all changes on any revised documents. Documents can be submitted with either the Drag and Drop option, or the Standard option.

When all documents have been uploaded, click the Continue button at the bottom of the page. This button will bring you to the Submit tab.
When you are sure every concern listed in the CAL has been addressed, click the Submit button.

Once the CAL has been submitted, it will be routed for signature by the PI. If you need assistance, please do not hesitate to contact the IRB. Each team has a Regulatory Compliance Analyst that has been involved with the development of the Wizard system. These analysts are there to help you with any issues or difficulties you may encounter.
Assign an NCT number and/or Business Officer

The “Info” tab for your approved study has been designed to allow you to enter information regarding your National Clinical Trial registry number (if you are required to register your study) or assign a Business Officer for your study. The Business Officer is responsible for the financial aspects of the study with regard to payment of IRB invoices. If you need to add/update either item, just follow the step-by-step instructions below:

On the Dashboard, Approved Studies can be found in the list at the bottom of the screen. Approved studies for which you are the PI, Study Coordinator, or Faculty Advisor are found on the My Studies tab. This is the default view for the dashboard. Studies for which you are listed as KSP are found on the Studies listing me as KSP tab.

If you see the study that the non-compliance with the protocol is related to, click the study title to go to the study page. If you cannot find the appropriate study, you can use the "search" field by typing the IRB number, sponsor number, part of the title, or another relevant piece of study information.
From the study page, be sure you are on the Main tab.

To include a ClinicalTrials.gov registry number, enter the NCT ID in the box provided and click save. When the page has saved, a message will appear on the top of the page indicating the save was successful.
To add a Billing Contracts/Grants Manager, enter the name in the box provided and click save. When the page has saved, a message will appear on the top of the page indicating the save was successful.
Managing Performance Sites on a Single IRB study where Vanderbilt is the IRB of Record

Once a new study is approved, the study team can manage performance sites from the study page.

On the Dashboard, Approved Studies can be found in the list at the bottom of the screen. Approved studies for which you are the PI, Study Coordinator, or Faculty Advisor are found on the My Studies tab. This is the default view for the dashboard. Studies for which you are listed as KSP are found on the Studies listing me as KSP tab.

If you cannot find the appropriate study, you can use the “search” field by typing the IRB number, sponsor number, part of the title, or another relevant piece of study information.

From the study page, click the Performance Sites tab.
On this tab, you can see the status of all sites that have been added to the study. You can also begin the process of adding sites and remove sites from this page.

Studies that have been submitted with a Site Add submission and are approved will appear in Green and will have a status of Approved.
Approved sites will also have an approval date in the Approval Date column.

<table>
<thead>
<tr>
<th>Site</th>
<th>Agreement</th>
<th>LOI</th>
<th>Local Context</th>
<th>Reliance Decision</th>
<th>Approval Date</th>
<th>Status</th>
</tr>
</thead>
<tbody>
<tr>
<td>Alzheimer Disease Center</td>
<td>SMART IRB</td>
<td>Yes</td>
<td>3/3 surveys complete</td>
<td>Yes</td>
<td>8/20/2019</td>
<td>Approved</td>
</tr>
<tr>
<td>Detroit Medical Center</td>
<td>SMART IRB</td>
<td>No</td>
<td>3/3 surveys complete</td>
<td>Yes</td>
<td>--</td>
<td>Study</td>
</tr>
<tr>
<td>Women &amp; Infants Hospital</td>
<td>SMART IRB</td>
<td>No</td>
<td>3/3 surveys complete</td>
<td>Yes</td>
<td>8/20/2019</td>
<td>Approved</td>
</tr>
<tr>
<td>Billings Clinic</td>
<td>SMART IRB</td>
<td>No</td>
<td>2/3 surveys complete</td>
<td>No</td>
<td>--</td>
<td>Removed</td>
</tr>
</tbody>
</table>

Sites that have been added to the table, but have not been marked as Ready to be Added by IRB staff will appear in white. These sites have the ability to be removed by the study team by clicking the Remove button.
If there is an outstanding document preventing the site from being ready to be added, the Local Context column will indicate how many out of the 3 required surveys have been completed. Clicking on the arrow will display a drop down menu which will indicate which surveys have been completed and which have not. The surveys with a check mark next to them have been completed, and those with a blank box have not yet been completed.

To add a new site to this table, click Add a Site.
You will be prompted to include the site name. If the site does not appear in the drop down list, add the site in the Requested Sites box. After entering the site name in either the Single IRB Performance Sites table, or the Requested Sites box, click Save Sites. This will then appear for the IRB Analysts to begin gathering information for that site.

Once an analyst has marked a site ready to be added, it will appear in blue with a status of Ready.
Submitting a Site Addition for Single IRB Studies

Once you have sites ready to be added as indicated on the Performance Sites table, click Create Submission.

Select Site Addition.
You will be prompted to select the sites to be added with this submission. Only the sites that have a status of Ready will appear in this section. Select the site(s) to be added by selecting the box next to the site name. Click save and continue when the sites have been selected to move forward to the Document Uploads tab.

The system will create a section for documents to be uploaded for each site. In the Document Uploads tab you can 1) expand or collapse each site section individually by clicking on the green bar with the site name in it, 2) expand all sections by clicking Expand All, and 3) collapse all sections by clicking Collapse all.
Once a site is expanded, it will have its own drag and drop section to attach documents.

To use the standard document uploads feature, click Standard.
Each site will appear in the Site drop down menu. Select the appropriate site for the document you are uploading.

Once all documents are attached, click Continue to move forward to the Submit tab.
After clicking submit, the Site Addition submission will be ready for signature and routing to the IRB.

Your submission is ready to be submitted. Click the submit button to start the signature process. Once all required parties have signed off on the submission, it will be sent to the IRB for review.
Glossary of Terms

Committee Action Letter (CAL) – A letter from the IRB that needs a response from the Principal Investigator

DISCOVR-E - Data Integrated Study Console of Vanderbilt’s Research Enterprise

E-Submission – An electronic study submission sent to the IRB for review

Final Approval Letter (FAL) – A letter from the IRB stating that a submission has received approval.

KSP (Key Study Personnel) – People responsible for helping with the conduct of a study

Log-In – Use your VUnetID and password to enter the DISCOVR-E system

My Studies – The view within DISCOVR-E where you can view studies where you are listed as the PI, Faculty Advisor or Study Contact

Principal Investigator – The individual responsible for the conduct of a study

Portal – Another name for the DISCOVR-E system

Studies listing me as KSP – The tab within DISCOVR-E where you can view studies in which you are listed as KSP

View IRB Training – The dashboard in DISCOVR-E where you can view your current IRB Training Status and when that training will expire

Wizard – The computer programming that takes you step by step through DISCOVR-E