Introduction

The Experiment Management System is used for the scheduling and management of research participants and the studies they participate in. Participants, researchers, principal investigators, and instructors all use the system for their respective purposes. As a researcher, you can set up your studies in the system, schedule the sessions (timeslots) when participants may participate, and grant or revoke credit after the session. All of this is handled through a simple web-based interface that you can access at any time, from any popular web browser.

System Basics

In the system, you create studies. Each study may have a number of timeslots, which are the times when you plan to run the study. Participants sign up for the timeslots by viewing a list of studies and available timeslots. You grant or revoke credit to participants after the session occurs.

Getting Started

The system works best if you use any popular web browser that is less than 2 years old, like Internet Explorer, Firefox, and Safari. It will work with other web browsers, and with older versions of popular web browsers, however the layout may not be as clean. No functionality will be lost by using an older web browser.

Ask your system administrator if you need help with installing or using a web browser. This documentation assumes you have a basic knowledge of how to use the web. On this system, it is not necessary to use the Back button. You can always use the toolbar on the top to navigate to anywhere on the site.

Logging In

Your administrator will provide you with a username and password to login to the site, as well as the URL (web address).
Your login (also known as a session) will expire after a certain period of inactivity, usually 20. This is done for security purposes. If this happens, you can always log in again. When you are done using the system, it is better to explicitly log out, to prevent any problems that may arise if someone uses your computer before the session expires.

**Retrieving a Lost Password**

If you have forgotten or do not have your password, then you may choose to have your password emailed to you. You will see an option on the main login page. Your password
will be emailed after you submit the form, and should arrive in your email box momentarily. If you provided an alternate email address (see the Email Address Options section of this documentation), it will be sent there. Otherwise, it will be sent to your main email address, which is derived from your user ID. If you requested that the system email a password to you, and it has not arrived after 30 minutes, then check in your email program’s junk mail folder in case the email was delivered there.

**Logging Out**

When you are done using the system, choose Logout from the top toolbar to log out. You are now logged out. It is always a good security practice to close all your web browser windows as well, especially if you are using a computer that is shared by others.

**Changing Your Password and Other Information**

If you would like to change your password or other information about yourself, choose My Profile from the top toolbar. If you would like to change your password, type your new password (twice, for confirmation) in the provided boxes. If you would not like to change your password, simply leave these boxes empty.

If you change your password, please be sure to select a password you do not use on any other systems or websites. This is good computing practice, and especially important as in some cases, your password may be sent over email.

![Figure 3 - Updating Your Profile](image)

It is recommended you provide your phone number and office location, as most human subject committees require that this information be made available to research participants. If you are a researcher, this contact information will be displayed to participants when they view information about the study. If you are a principal investigator, this contact information will be displayed if a participant explicitly chooses to view it (since the researcher is the primary point of contact for a study).
Researchers may also choose to receive a daily reminder (by email) with information about all of their study sessions scheduled for the following day.

**Email Address Options**

There are certain events in the system which will cause an email notification to be sent to you. Most often, these are notifications that a participant has signed up or cancelled their sign-up for your studies, but there are a few other cases where it may be used as well. The email address is also displayed to the participant when they view information about the study, in case they need to contact you with questions.

You will be required to provide an email address (which will be listed as “Email Address” instead of “Alternate Email Address”) and all emails to you will go to that address.

**Working with Studies**

Most of your time on the system will be spent, not surprisingly, using the study-related features of the system. Be sure to read this section closely, in its entirety, as there are special features and situations you should be aware of.

**Two-Part Studies**

You may create a two-part study in the system. Often, these are studies involving memory research, where the participant must return a specified number of days after the first session. When creating a study, you may specify the day range for the second part of the study (e.g. 7 to 10 days after the first part). Participants are required to sign up for both sessions at the same time, to reduce the chance they will forget to sign up for the second part. Each part of a two-part study may have a different credit value and duration.

You may specify that the second part of the study must be scheduled to take place at the exact same time as the first part (on a different date), or at any time on the dates that are the specified number of days after the first part.

You should ensure there are enough available timeslots for both parts of the study, or participants will be prevented from signing up for either part. Participants may cancel either part of their sign-up if necessary. If they cancel the first part, the second part is automatically cancelled as well. If they cancel only the second part and the first part has already occurred, and they would like to participate in the second part later, you will need to manually sign them up for the second part.

If you grant a no-show for the first part of a two-part study, the second part of that participant’s sign-up will not be cancelled automatically, but you will be reminded of the situation in case you would like to cancel the second part. The cancellation is not automatic as there are some situations where automatic cancellation is not desirable.
**Adding a Study**

Do not add a study to the system. Any studies added to the system that were not approved by the HREAP will be deleted.

**Updating a Study**

You may update any of your studies at any time. To do so, choose My Studies from the top toolbar, and you will see a list of your studies. Click on the desired study, and choose the Change Study Information link. See the table below for information regarding each field. The changes you make will take effect immediately after they are saved.

If you need to change the credit value for a study, and there is no option to do so, this means the study already has at least one participant signed up for it. You cannot change the credit value when a study is in this situation because there is no easy way to handle past credits for the same study (e.g. should old credit grants for the same study be adjusted to reflect the new credit value, or kept the same?). If the study is nearing the end of its run, and variable credit granting is enabled, then the easiest solution is to grant the new credit value to participants who sign up in the future. If you prefer that the credit value is changed for the entire study, contact the administrator, who can make the change for you under certain conditions (which depend on the credit status of existing sign-ups for the study).

<table>
<thead>
<tr>
<th>Field</th>
<th>Explanation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Study Name</td>
<td>Assigned by Linda. DO NOT CHANGE.</td>
</tr>
<tr>
<td>Brief Abstract</td>
<td>Approved by ethics. DO NOT CHANGE WITHOUT APPROVAL.</td>
</tr>
<tr>
<td>Eligibility Requirements</td>
<td>Approved by ethics. DO NOT CHANGE WITHOUT APPROVAL.</td>
</tr>
<tr>
<td>Pre-Requisites</td>
<td>If there are studies a participant must participate in before participating in your study, choose them here. You may select multiple studies, and on most systems, you hold down the Ctrl key and click the desired studies. You may specify that participants must have participated in all of the studies you specify, or at least one of the studies specified. The system will handle enforcement of the pre-requisites in a strict or lenient fashion depending on how your system is configured. In strict enforcement mode, the participant must have received credit for (participated in) the pre-requisite studies. In</td>
</tr>
</tbody>
</table>
lenient enforcement mode, the participant must only be scheduled to participate in the pre-requisite studies (it is assumed they will go on to complete the pre-requisite studies). You can ask your administrator how this is configured, if it is of concern. If your system is in lenient enforcement mode, and a participant cancels a necessary pre-requisite for you study (they are warned of this situation), and you have configured your study so that the researcher will receive notifications of cancellations or sign-ups, then the researcher will receive notification of the pre-requisite problem and can contact the participant if necessary.

If there is a long list of studies for this setting, an Enlarge List button will appear. You can click this to make the list of studies larger and thus easier to click on.

<table>
<thead>
<tr>
<th>Disqualifiers</th>
</tr>
</thead>
<tbody>
<tr>
<td>If there are any studies a participant must not have participated in, please select them here. You may select multiple studies. The system will handle enforcements of the restriction, during the sign-up process.</td>
</tr>
<tr>
<td>If a study has some other study listed as a disqualifier, and a participant signs up for this study, then they will be prevented from signing up for the disqualifier study.</td>
</tr>
<tr>
<td>If there is a long list of studies for this setting, an Enlarge List button will appear. You can click this to make the list of studies larger and thus easier to click on.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Duration</th>
</tr>
</thead>
<tbody>
<tr>
<td>The amount of time, in minutes, that each study session will take. If you are setting up a 2-part study, then this setting applies to the first part of the study.</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Timeslot Usage Limit</th>
</tr>
</thead>
<tbody>
<tr>
<td>Depending on how your system is configured, you may see an item that specifies the maximum number of study session hours available to this study. This value is set by the administrator, and only the administrator can adjust it. To determine the current session usage for a</td>
</tr>
</tbody>
</table>
study, go to the Add A Timeslot page for
the study, and the usage will be listed there.

<table>
<thead>
<tr>
<th>Preparation</th>
<th>Enter any advanced preparation a participant must do here (e.g. “do not eat 2 hours before session”). If there are no preparation requirements, leave this field as-is.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Invitation Code (USED RARELY)</td>
<td>If you would like to have a special sign-up password for this study, enter it here. This is known as an invitation code, and applies just for this study. Participants must know the invitation code to sign up for this study. This is often used in cases where the researcher wants to personally select participants, so the researcher only provides the invitation code to the desired participants. Invitation codes are not case sensitive. If you do not need an invitation code, leave this field blank.</td>
</tr>
<tr>
<td>Credits</td>
<td>Enter the number of credits or compensation for the study. A value of 0 is acceptable, and may be desired in cases where the study is part of a set of studies, where only the final study is credit-earning. If the study has a credit value, the credit value specified must be evenly divisible by the credit increment specified. If you are setting up a 2-part study, this is the value for the first part of the study. After a study has sign-ups, you may not change the credit value of the study. However, the administrator can change the credit value, in certain situations. A study may not be changed between a study for credit and for payment, after it has been created.</td>
</tr>
<tr>
<td>FOR TWO PART STUDIES:</td>
<td></td>
</tr>
<tr>
<td>Credits, Part 2</td>
<td>Enter the number of credits for part 2 of the study, if this is a two-part study (the value is ignored otherwise). A value of 0 is acceptable, and may be desired in cases</td>
</tr>
</tbody>
</table>
where the study is part of a set of studies, where only the final study is credit-earning. If the study has a credit value, the credit value specified must be evenly divisible by the credit increment specified.

<table>
<thead>
<tr>
<th>Part 2 Duration</th>
<th>The amount of time, in minutes, that part 2 of the study will take.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Part 2 Scheduling Range</td>
<td>Specify the number of days (as a range) after part 1 is scheduled, that part 2 should be scheduled. This setting only applies to two-part studies. The range may be the same value (e.g. “between 7 and 7 days”) if desired, but must be a whole number. See “Two-Part Studies” for more information.</td>
</tr>
<tr>
<td>Part 2 Scheduling Leniency</td>
<td>In some cases, you may want to ensure that the participant schedules the second part of the study to take place at exactly the same time (on a different date) as the first part. If so, choose Yes for this option. If there is some flexibility so they can sign up for any time within the Part 2 Scheduling range, choose No for this option.</td>
</tr>
</tbody>
</table>

**OTHER INFORMATION:**

| Active Study? | Select Yes if this study is in progress. You must select Yes and the study must be Approved if you want the study to show up to participants so they can sign up for it. The reason to select No is if the study is being kept for historical purposes, but should not show up to participants on the list of studies they may sign up for. Often, this is done so the system can enforce pre-requisites, where the inactive study is a pre-requisite for an active study. |
| Should the Researcher receive an email notification when a participant signs up or cancels? | If set to Yes, the researcher for this study will receive an email notification whenever a participant signs up, or cancels their sign-up, for this study. The email notification will be sent to an email address based on the information the researcher has provided. See the Email Address Options section of this documentation for more information on how the email address is determined. Emails will contain the first 50 characters of the study name as part of the |
subject line, to make it easy to sort the emails with an email program that supports filtering based on subject line.

Emails are sent to all researchers specified for the study, unless a specific researcher is assigned to the timeslot that the email notification is being sent about. See Timeslots Linked to Specific Researchers for more information.

<table>
<thead>
<tr>
<th>Researchers at Timeslot-Level</th>
<th>If set to Yes, it will be possible (but not required) to assign a specific researcher (from the list of researchers for the study) to a timeslot. If set to No, then it is assumed that all researchers (assigned to the study) are responsible for all timeslots. See Timeslots Linked to a Specific Researcher for more information.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Can a participant sign up for this study more than once?</td>
<td>If you would like to allow participant to sign up (and receive credit) for your study more than once (at different times), choose Yes. Otherwise, choose No. If No is chosen, participants may only sign up for the study more than once if they previously failed to show up for the study (a no-show).</td>
</tr>
<tr>
<td>Private Comments</td>
<td>This is an optional area where you may enter any comments or notes about the study, which are only visible to the researchers for this study, and not to participants. The maximum length of this field is 3,000 characters.</td>
</tr>
<tr>
<td>Participant Sign-Up Deadline</td>
<td>Enter the deadline before the study is to occur that the participant may sign up, in whole hours.</td>
</tr>
</tbody>
</table>

**Deleting a Study**

Do not ever delete studies from the Sona System. If you are done collecting data from a study, you should deactivate it, but do not delete it.

**Timeslot Usage Summary**

The timeslot usage summary is available when viewing your study. This gives some basic information about timeslot utilization in the past and in the future, as well as some basic no-show information. It also gives information on timeslots for the study by location (if
the study is not an online survey study or external web study), and by researcher (if the study is configured to allow researchers to be assigned to specific timeslots).

For credit studies, the system also provides a summary of how many credits were granted. This summary accurately computes credit usage, taking into account any variable credit grants (if Variable Credit Granting is enabled in System Settings), where some participants may have received credit in a different amount than the study’s listed credit value.

If timeslot usage limits are enabled, the system will provide an estimate of how many timeslots can be added. Note that if the study is a two-part study, it will estimate based on allocating all the limit to the first part or the second part (both estimates are provided), however in practice it’s more likely a researcher will want to add timeslots to both parts of the study, so this should be taken into account when viewing these estimates, especially if the first part and second part of a study have a different duration.

**Bulk Mail Summary**

The system tracks whenever any type of bulk email is sent (by a user) related to the study. This includes contacting those who have already signed up for the study. This information is kept for 6 months, and it is tracked to ensure that all users follow generally accepted Internet practices for responsible use of email. The administrator also has access to this information.

**Prescreen Participation Restrictions**

If enabled on your system, the system might contain an online prescreen that participants may (or must, depending on your system configuration) complete. You may place participation restrictions on your study based on prescreen responses. Participants are unaware that such restrictions are placed on the study. These restrictions are never listed to them. If they do not qualify to participate in a study because they do not meet the prescreen participation restrictions, then the study will simply not be listed to them. This is important to note – participants never know why a study was or was not listed to them, because they are unaware of the prescreen restrictions.

You may restrict a study on any question or questions on the prescreen that allowed for a multiple-choice answer where only one choice could be selected. You may also restrict a study based on a computed section sum or average score for a participant, if the prescreen was set up in such a manner. You may restrict to one choice or many choices for any question. If you restrict on multiple questions, it is the same as a logical “AND.” For example, if you setup the prescreen restrictions so that participants must have answered “Yes” to a “Do you wear glasses?” question and “Blue” or “Grey” to “What color are your eyes?”, then they must meet both requirements to participate. In other words, only participants who wear glasses and have either blue or grey eyes are eligible. There is no support for a logical “OR” restriction across multiple questions. The restrictions are inclusive, which means that if you select a choice as a restriction, then participants must have answered at least one of the choices selected for each question that is part of the
restriction in order to see and participate in the study, as opposed to exclusive where checking the choice as a restriction would exclude them from participation.

Figure 4 - Prescreen Restrictions Question Selection

To set participation restrictions, view (do not choose edit) your study and choose View/Modify Restrictions. You will see a list of eligible questions which you may use for your restrictions. If the study already has some restrictions, those will be checked, and you will see how many participants currently meet the restrictions. Choose the questions you would like to restrict upon (and keep the existing checked restrictions checked, unless you want to remove that restriction), and click on the Set Restrictions button. On the subsequent page, you can select each value that is acceptable for each question you have chosen. Once you have selected all the acceptable values, save your changes and they will take effect immediately. It is important to note that if you change the restrictions, it will not remove the study sign-ups for participants who qualified under the previous set of restrictions. For this reason, you should probably decide on your restrictions before making the study available to participants.

If you have restriction requirements where you would like to restrict participation to a percentage of the population (for instance, the responses that were chosen by the top 25% of people), but you are not sure which responses meet this requirement, you can use the prescreen response analysis feature to determine the valid responses. See Prescreen Response Analysis for more information. You may also use Analyzing Prescreen Responses to get an idea of how many participants are potential candidates for participation in your study, based on a specified set of restrictions.
Figure 5 - Prescreen Response Restrictions

Viewing Your Studies

To view your studies (and not the studies of others), choose the My Studies option on the top toolbar. The system will list all your studies in alphabetical order by study name, grouped by studies that are active, then inactive studies.

Figure 6 - Your Studies
**Participant Study View**
If you would like to see how your study appears when participants view it, find your study and choose the Participant Study View option. This will show exactly how the study appears to participants, with the exception that when a participant views a study, next to each pre-requisite and disqualifier study (for a study) is listed a status indicator about whether they have met that requirement. In Participant Study View, the pre-requisite and disqualifier studies are listed, but there is no status indicator next to each study in the list.

If for some reason you think your study is not visible to participants, it may be due to various restrictions you have set on the study, like prescreen participation restrictions, such that few (or none) of the participants in the pool qualify. You can ask the administrator to use the Check Study Configuration tool (available to them when they view your study) to provide advice on why your study may or may not be visible to participants. Administrators there also have an option to type in a specific participant to see if that participant would qualify for your study.

**Viewing Other Studies**
To view all studies that are visible to participants, choose the All Studies option from the top toolbar.

You will see a list first of all Active studies. These studies will show up to participants on the list of available studies. The next group of studies (if there are any) is Inactive studies. These will not show up on the list of available studies (to participants), but participants can access information about these individual studies on links from the page with their progress (if they participated in the study) or if another study has the Inactive study listed as a pre-requisite or disqualifier.

**Working with Timeslots (Sessions)**
Timeslots (also referred to as Sessions) are the available times when a participant may participate in the study. Timeslots allow you to specify a date, time, location, maximum number of participants, and researcher for a session.

**Timeslot Usage Restrictions**
If enabled on your system, you may find there is a limit to the amount of time available for scheduling timeslots. This usage is computed by adding up all the past timeslots where credit was granted, and then adding all timeslots in the future, regardless of credit status. You may find that the usage goes down over time, as time progresses and timeslots that were in the future had no participants signing up for them. The usage and limit is listed whenever you add a timeslot, if usage restrictions apply. It may also be listed when you view your profile, depending on how your system is configured.
**Timeslots Linked to Specific Researchers**

You will also have an option to link timeslots to a specific researcher. This is done primarily for organization purposes, and has no effect on who can view and modify the study, or any timeslots for that study.

This feature is useful when there are a number of researchers running a study, and researchers are responsible for running specific timeslots. If a timeslot has a specific researchers linked to it, then only that researcher will be listed as the contact point when a participant receives any emails related to their participation in that timeslot. Finally, only the researcher connected to that timeslot receives related notification emails, such as participant cancellation notification, and reminder emails (assuming such emails are enabled).

It is also possible to have some timeslots where a specific researcher is linked to them, and others where all researchers (who are assigned to the study) are responsible for the timeslot. It is not possible to link more than one, but not all of the researchers (for the study), to a specific timeslot. The options are to either link one researcher to the timeslot, or all of them.

If a researcher is removed from a study, then any timeslots that were linked to them for that study will be changed so all researchers (for the study) are now responsible for those timeslots.

To use this feature, the system must be configured to allow multiple researchers per study. Then, the study itself must be configured to allow researchers to be linked to specific timeslots. Finally, the study must have more than one researcher connected to it.

**Creating Timeslots**

To add a timeslot for a study, you must first choose the study that you would like to add a timeslot for. To view your studies, choose the My Studies option on the top toolbar. Click on the desired study, and choose the Timeslots choice.

You will see a list of any existing timeslots, and the Add A Timeslot option at the bottom of the page. Click on Add A Timeslot.
The following table lists the information you may enter about a timeslot, along with an explanation. All fields are required.

<table>
<thead>
<tr>
<th>Field</th>
<th>Explanation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Date</td>
<td>The date for the timeslot.</td>
</tr>
<tr>
<td>Start Time</td>
<td>The time for the timeslot. A sample time will be provided. If you want to</td>
</tr>
<tr>
<td></td>
<td>change the time, please use the same format as the time you see presented.</td>
</tr>
<tr>
<td></td>
<td>Note in particular how “a.m.” and “p.m.” are handled (if such a format is</td>
</tr>
<tr>
<td></td>
<td>enabled on your system).</td>
</tr>
<tr>
<td>End Time</td>
<td>The time when the timeslot will end. This is computed automatically based on</td>
</tr>
<tr>
<td></td>
<td>the duration you entered when you set up the study.</td>
</tr>
<tr>
<td># of Participants</td>
<td>The number of participants for this timeslot. This limit is not visible to</td>
</tr>
<tr>
<td></td>
<td>participants. They will only see whether the timeslot is full or not. The</td>
</tr>
<tr>
<td></td>
<td>maximum number is 999.</td>
</tr>
<tr>
<td>Location</td>
<td>The physical location where the study will take place, for this timeslot.</td>
</tr>
<tr>
<td></td>
<td>It will be automatically filled with the location of the previous timeslot,</td>
</tr>
<tr>
<td></td>
<td>when available, to ease in data entry.</td>
</tr>
<tr>
<td></td>
<td>Depending on how your system is</td>
</tr>
</tbody>
</table>
configured, you may see a list of pre-configured locations. You may choose any of those locations and click on View Schedule to see the schedule for a location. The system will automatically prevent you from adding a timeslot using a location that is already in use at the time you try to schedule the timeslot. If you do not see the location in the list that you plan to use, you can simply type in the location in the text field below it.

The location field does not apply for web-based studies.

| Researcher | The researcher assigned to this specific timeslot. The list will contain a list of all researchers for the study. Choose ALL if all researchers (for the study) should be assigned to this timeslot. See Timeslots Linked to Specific Researchers for more information. |

To ease data entry, the system will automatically fill in the date, time, and location based on the ending time of the last timeslot for this study. If applicable, your current timeslot usage will be listed, and you will be prevented from adding a timeslot that would exceed your timeslot usage time limit. A convenient calendar is provided next to the form, and you can click on any date and that date will be transferred to the form.

If you add a timeslot such that there is another timeslot (for any study) that occurs in the same time, at the same location, you will receive a warning (but the addition will be allowed). If you add a timeslot that will take place outside of normal hours (for example, at 1:00am), the system will provide a warning but will allow it to be scheduled. The system allows adding timeslots to a study that is not available to participants (not active or not approved), but it will give a warning because participants are not able to sign up for the timeslot.

**Creating Multiple Timeslots**

If you would like to add multiple timeslots at once, choose the Add Multiple Timeslots link. You may choose to add a specified number of timeslots, or copy the timeslots from another week to a specified week. If you choose to copy, the system will copy the time, location, and number of participants for the specified week to the desired week, for each day of that week (starting with Monday).

If you choose to create a specified number of timeslots, you can choose the number of timeslots you would like to add, the start time and date, and the amount of time between
each timeslot (to allow for breaks). You also may specify that timeslots that would occur outside normal business hours be shifted to the next business day, and specify when business hours occur. The system considers Monday-Friday to be business days.

On the subsequent page, you may change any of it to deal with special cases. Timeslots that you attempt to add, that either have errors or would result in exceeding the timeslot time usage limit, will not be added. This feature is not available for web-based (online) studies, as web-based studies rarely have more than one timeslot.

If you would do not want to add a specific timeslot that is listed, choose No in the Add This Timeslot? Column.

![Figure 8 - Creating Multiple Timeslots](image)

**Modifying and Deleting Timeslots**

To modify or delete a timeslot for a study, you must first choose the study that you would like to deal with. To view your studies, choose the My Studies link from the top toolbar. Choose the Timeslots option in the timeslots column for the desired study. You will see a list of all recent timeslots. Recent timeslots in the past with no participants signed up will not be displayed. To work with timeslots more than a few days old and to see all timeslots, you will see a link to view all timeslots for the study. Select the timeslot you would like to deal with, and click the Modify button.

If the timeslot has no participants signed up for it, you will see a Delete button. You may not delete a timeslot that has participants signed up for it. If you would like to delete the timeslot, click the Delete button, and you will see a confirmation page. Choose Delete again to delete the timeslot.

If you would like to modify the timeslot, modify the desired information and click the Update button just below the timeslot information. It should be noted that participants will not be notified (by email) of any changes you make to the timeslot, so you should contact them if information needs to be passed on to them (a link is provided on the same page to do so). If you change the date or time of the timeslot, you will be warned that this was changed in case the change was unintended. You may not update the size of the
timeslot (number of participants) to a value lower than the current number of participants signed up for the timeslot. Generally, researchers only update timeslots with sign-ups to update the location, if it was not available when the timeslot was originally created.

If the study (or researcher) is subject to timeslot time usage restrictions, the system will enforce them and prevent you from increasing the number of participants in a timeslot if that would result in exceeding the timeslot usage limit.

**Timeslot Change Tracking**
The system automatically tracks certain changes that occur with a timeslot, including any time key information about the timeslot (date, time, etc.) is changed, as well as any time a manual sign-up or cancellation is performed (i.e., not a sign-up or cancellation done by the participant). This information is tracked for the last 3 months of changes for each timeslot.

To view this information, choose the View Timeslot Modification Log when viewing a timeslot, and you will see this information.

**Deleting Multiple Timeslots**
If you would like to delete multiple timeslots at once, you may do that as well. Such a feature is only available for timeslots which have no participants signed up. To do so, select the desired experiment and choose Timeslots. At the bottom of the Timeslots page, you will see a Delete Multiple Timeslots option. The option may not appear in certain cases where such an option is not available because of a lack of available timeslots to delete.

After going to that page, you will see a list of timeslots eligible for deletion. Choose the timeslots you would like to delete, and choose Delete Selected Timeslots to proceed. If you would like to delete all empty timeslots, there is a Select All option at the bottom of this page that will automatically select all timeslots listed on the page for deletion. Click the Reset button to revert the effect of choosing the Select All option.

The system routinely deletes all empty timeslots more than 3 months old to preserve database space.
Figure 9 - Delete Multiple Timeslots

**Manual Sign-Up**

If enabled on your system, you may manually sign up participants for your study. There are a number of situations where this is desirable. If the participant happens to show up for a timeslot they were not signed up for, and you elect to let them participate, you can sign them up on the spot for the timeslot. The participant in many cases cannot sign up on their own in this situation, because the sign-up deadline has passed. You may also sign up a participant for a study that has already occurred, if necessary.

Also, a manual sign-up overrides any restrictions you have placed on the study (e.g. prerequisites), though you will be warned if you are overriding any restrictions. You may not sign up a participant for the same timeslot that they are already signed up for. You are allowed to sign them up for a study even if they are already signed up for a different timeslot for that same study, though you will receive a warning in this case. You may not sign up a participant for a study if it would cause them to exceed their maximum credit limit. If it is necessary to do so, please ask the administrator to do this, as they are allowed to do a manual sign-up even when it will violate maximum credit earning limits.

You also may not sign up a participant whose account is Limited, if your study is not a research alternative study, as those participants are ineligible for your study (the administrator can still do this).

If the system is configured as such, the participant will receive a confirmation email when you sign them up for a study. In that case, you are also given the option to enter comments to be included in this email that may better explain to the participant why they were signed up. If you are signing up a participant for a timeslot more than one year old, a confirmation will *not* be sent despite the system configuration. This is to make it easier
when transitioning from an existing system, as you may sign up old participants for the purposes of preventing them from signing up for the same study again in Sona. You may only sign up participants for your own study.

To sign up a participant for a timeslot, you must first find the desired study and timeslot. To view your studies, choose the My Studies option from the top toolbar. Click on Timeslots for the desired study, then select the timeslot you would like to deal with, and click the Modify button.

At the bottom of the page, you will see a Manual Sign-Up option, if it is enabled. Type in the participant’s User ID (you may have to ask them for this) and click Sign Up. If enabled, you may also choose to sign up a participant using their unique ID code. You may also have the choice to enter their last name and choose from a list of participants. In all cases, after submitting the form, you will see a confirmation page that also lists any restrictions on the study. Choose Sign Up to complete the sign-up.

![Figure 10 - Manual Sign-Up Confirmation](image)

If you are subject to timeslot time usage restrictions, the system will enforce them and prevent you from signing up a participant in the timeslot if that would result in exceeding your timeslot usage limit.

If you are doing a manual sign-up for a two-part study, you must do a manual sign-up for each part separately. The system will overlook the scheduling range restrictions as well.

You cannot use the manual sign-up feature for online survey studies, because the sign-up for the study is integrated with the administration of the survey.

The manual sign-up feature will not appear for a researcher if the study requires approval by the administrator and it has not yet been approved. This is to ensure sign-ups cannot occur for a study that has not yet been approved, since research should not take place prior to approval.
**Manual Cancellation**

If enabled on your system, you may have the opportunity to cancel a participant’s sign-up. You may only cancel sign-ups that are in a No Action Taken state. To cancel a sign-up, find the desired timeslot and participant, and click Cancel next to their name. The participant will be sent an email about the cancellation (and who performed it), along with a confirmation code, and their sign-up will be immediately cancelled. The administrator may also receive a copy of this cancellation email, depending on how the system is configured.

You may cancel all participants for the same timeslot at one time, when applicable. The option will appear below the list of signups, in cases where there are two or more participants signed up for the timeslot who are eligible for cancellation (No Action Taken state).

![Figure 11 - Manual Cancellation](image)

**Viewing the Participant List**

To view the list of participants who have signed up for your study, you must first select the study and timeslot you wish to see. To view your studies, choose the My Studies option from the top toolbar. Click on the Edit link in the timeslots column for the desired study, then select the timeslot you would like to see, and click the Modify button.

The list of participants, along with their email addresses, will be listed. If ID codes are enabled, you will only see an ID code and no name or email address for each participant, and the list will be sorted by ID code.
At the completion of a session, you should promptly deal with the participants, in the system, to ensure proper credit grants. The reason for the prompt handling of this situation is in the event your study is a pre-requisite for another study, and a few other situations. You do not want to hold up other studies that are waiting on your response to the study you just ran.
To grant or revoke credit for a timeslot, you must first find the desired study and timeslot. To view your studies, choose the My Studies option from the top toolbar. Click on the Edit link in the timeslots column for the desired study, then select the timeslot you would like to see, and click the Modify button.

You will see a list of participants, identified either by name or ID code. If the participant properly participated in the study, click the Credit Granted button next to their name (this text may appear as Participated if the study is set up for payment).

If the participant did not appear for the timeslot, you may choose to mark their no-show as excused or unexcused. Depending on how your system is configured, an unexcused no-show may result in a penalty being assessed for the participant (the system will compute this automatically), or their privileges to use the system may be restricted. You should ask your administrator for guidelines about when to grant an excused no-show or an unexcused no-show. Generally, excused no-shows are granted for extenuating circumstances, like if the participant was involved in a car accident on their way to the appointment. An unexcused no-show is generally used when the participant did not show up and had no reasonable excuse. For most schools, the majority of no-shows are unexcused and are due to carelessness on the part of participants.

Depending on how your system is configured, you may see an option to grant a credit value that is different from the standard credit grant. This is useful when you want to grant a participant a lower credit value because they left the study early (if they deserve a lower credit grant), or a higher credit value if the study ran longer than expected. The default value that is selected is the study’s standard credit value. If this is enabled, then you may also grant 0 credits. This is useful if you do not want to grant credits to the participant, but you also want to prevent them from participating in the study again. If a participant is granted 0 credits, and the study is set to prevent duplicate sign-ups, then the participant will not be able to sign up for that study again.

If desired, enter any comments about the session in the Comments section (generally, this is used to indicate the reason for denying credit). Participants will see anything you enter in the Comments section for their sign-up, and these comments will be included in the email sent to participants when a credit grant/revocation occurs, if notification emails are enabled on your system.

Click on the Update Sign-Ups button at the bottom of the list of sign-ups to save your changes. Credit will be granted or a penalty assessed as necessary. The participant(s) will be emailed about this if the system is configured in such a manner.

It is not recommended to leave any sign-up for a timeslot that has occurred in the “No Action Taken” stage. This is a credit “limbo” and the system will warn you upon your next login about the offending timeslot that has not been dealt with properly. Note that if Manual Cancellation is enabled and you would like to cancel a participant’s sign-up, the sign-up must be in No Action Taken state.
Depending on how your system is configured, the system may automatically grant credit to participants for timeslots that are more than an administrator-specified number of hours old, and where the researcher has taken no action. You can always change the automatic credit grant later if it was in error. The automatic credit grant takes place once a day, usually overnight. Your administrator can let you know if such a feature is enabled on your system.

If you need to do a simple credit grant across many timeslots, see the Uncredited Timeslots section which offers such a feature.

**Batch Credit Granting**

Do not use this feature.

**Emailing Participants**

If you wish to contact participants in a particular timeslot for any reason, you may click on the Contact link that will appear next to each participant’s name (or ID code) to contact an individual participant. To email the group of participants for a particular timeslot, click the Contact All Participants choice at the bottom of the Modify Timeslot page for that timeslot.

You will be taken to a page where you can fill out a message that the system will send to the selected participants. The message is auto-filled with some basic information about the study, so participants are aware of which study you are referring to. You may remove this information if desired. You may choose to receive a copy of the email that you send.

Participants will receive a reminder about upcoming studies the day before they are scheduled to participate.

Figure 14 - Contacting Participants

Do not contact participants outside of the bounds of your ethics approval.
**Viewing Uncredited Timeslots**

When you login to the system, you will receive a warning if you have any timeslots that are more than 2 days old and haven’t been dealt with. You may view a list of all timeslots that have not been dealt with by choosing the View Uncredited Timeslots option from the My Studies page. The default view will show in-person studies with timeslots in the past, as well as all uncredited timeslots for online studies. Timeslots for online studies, including those in the future, are always considered in need of a response. See the Web-Based (Online) Studies section of this documentation for more information.

If you would like to do a simple credit grant (standard credit grant, no comments), you may do so directly from this page. Select the desired sign-ups/timeslots, and then choose Grant Credits. The action may take a short time to complete, so please be patient while the credit grants are processed.

If you need to do something more complex, like mark a no-show, add comments, or perform a special credit grant with a non-standard credit amount, you can easily click on the timeslot’s date and time, and go directly to that timeslot.

In cases where a study has timeslots linked to specific researchers, you will see the warning only for timeslots that are specifically linked to you, or to everyone in the study (i.e., not timeslots linked to someone else in the study). However, when you view uncredited timeslots, you will see all uncredited timeslots for your studies, even if someone else is linked to one of the timeslots for your study. This is done to make it easier to give your fellow researchers (for your studies) assistance in dealing with uncredited timeslots.

![Figure 15 - Uncredited Timeslots](Image)
Analyzing Prescreen Responses

If online prescreening is enabled on your system, then you might also have the opportunity to analyze prescreen responses in aggregate or as raw data. Choose the Prescreen Results option from the top menu bar. You can then select which question you would like to analyze, and whether you would like to see summary data or raw data (in CSV format) for the selected question. The raw data will identify each participant only by a unique ID code, not by their name, for privacy reasons. If for some reason you need the participants’ real names, ask the Administrator to run the same analysis, as they can also pull the real names with their report. This gives you access to all prescreen data across all participants in the system.

If you would like to analyze the prescreen data for just those who participated in your study, select the Download Prescreen Responses after clicking on your study. See Viewing Prescreen Responses in this documentation for further information.

Figure 16 - Prescreen Response Analysis

Prescreen Qualification Analysis

If you would like to get an idea of how many participants meet a set of requirements (for help in setting prescreen restrictions on your study), use the Prescreen Qualification Analysis link from the Prescreen Responses page. Using this feature, you can select multiple questions (only questions that qualify for study participation restrictions are listed), and then the desired responses for those questions, and you will see how many participants meet that criteria.

If enabled, you may also contact participants and invite them to participate in any of your studies. See the Invite Qualified Participants to a Study section in this documentation for more information on how this works. The functionality is the same as the functionality described in that section, though a few options may be unavailable when not inviting directly from a study, because those options do not apply. Be sure to include information
about how to sign up for the study in your communication to them, as a direct link to the study is not provided in the email.

**Frequently Asked Questions (FAQ)**

*I want a participant to participate in an upcoming session, but the system says it is too late for them to sign up. What do I do?*

If enabled, you can perform a manual sign-up. See the Manual Sign-Up section of this documentation. If not enabled, your administrator can still perform a manual sign-up.

*Where are email notifications to me sent?*

Email notifications (e.g., sign-up notices) are sent to either an address derived from your user ID or your alternate email address. See the Email Address Options section of this documentation for more information.

*How do I deal with dyads?*

A dyad is a study which requires a pair of people to participate, but often the second participant is not a “real” participant, but rather a colleague of the researcher who is “colluding” with the researcher as part of the study itself.

You do not need to deal with dyads in the system itself. Participants cannot see how many people have signed up for a timeslot, nor how many spaces are available for a timeslot. So, your “fake” participant can just act like a real participant and the real participant will be unaware of this.

*I have finished running my study. What should I do?*

So it does not clutter the list of studies for participants, you should make the study Inactive. See the Updating a Study section of this documentation for more information.

*Who has access to my studies?*

All users can see the information about your studies and the available timeslots. Administrators, the principal investigator (if applicable) and the researchers for the study are the only people who can see who has signed up, and modify the study.

**Regulatory Compliance Guidelines**

**Introduction**

This software complies with all major regulations governing human subject research and privacy of data stored online. The system complies with both HIPAA and Common Rule for customers in the United States. For customers in Canada, it complies with the Personal Information Protection and Electronic Documents Act as well as the Tri-Council
Statement. For customers in the European Union or in countries that follow OECD rules, it complies with OECD privacy rules and the European Union Directive of Data Protection. Your organization may or may not need to comply with the relevant regulations. Your subject pool administrator can advise you on this situation.

Even if you are not required to comply, compliance is still a good idea, as protecting sensitive data is always a good thing. Compliance in the context of this system is as simple as reading the remaining paragraphs of this section (that apply to your organization) and following the guidelines contained therein. The remaining compliance issues involving software, privacy and electronic data storage are all handled automatically by the software. You should still consult with your IRB or organization to learn about additional compliance rules you must follow outside of use of this software (the handling of the data you collect during your study would be one example).

Some regulations (particularly the US HIPAA regulations) are focused primarily on health data. You may think the system does not store confidential health data (in HIPAA terms, it is called PHI -- Protected Health Information), but depending on how your organization uses the software, there may very well be confidential data in the system. Consider the case of a study that requires that a participant come from a family that has a history of mental illness. Merely knowing who signed up for that study can be considered confidential because that type of information should not be revealed to the public. It may turn out that your studies are not of such a nature, but even more benign situations, like a study that requires that participants be regular contact lens wearers, can be construed as confidential information. Organizations typically err on the side of caution given the criminal and civil penalties for violation of these types of regulations.

**Data Handling and Security Guidelines**

In your role, you have access to your studies and you can see who has signed up for those studies. You may also have access to prescreen responses. Because of these privileges, you should follow these simple guidelines:

- **Secure Your Account.** Use a password that is difficult to guess. The most secure passwords contain a combination of letters and numbers, do not spell a real word, and are at least 8 characters long. Your university IT department can provide you with assistance on choosing a secure password.

- **Secure Your Work Area.** If you are logged into the system and you leave your computer, you should logout of the system or use a password lock on your computer. Ask your network administrator for help with setting up a password lock.

- **Handle Paper Documents Carefully.** Any printouts from the system should be kept reasonably secure. Store them in desk drawer out of the public view. Documents you decide to discard should be shredded if possible.