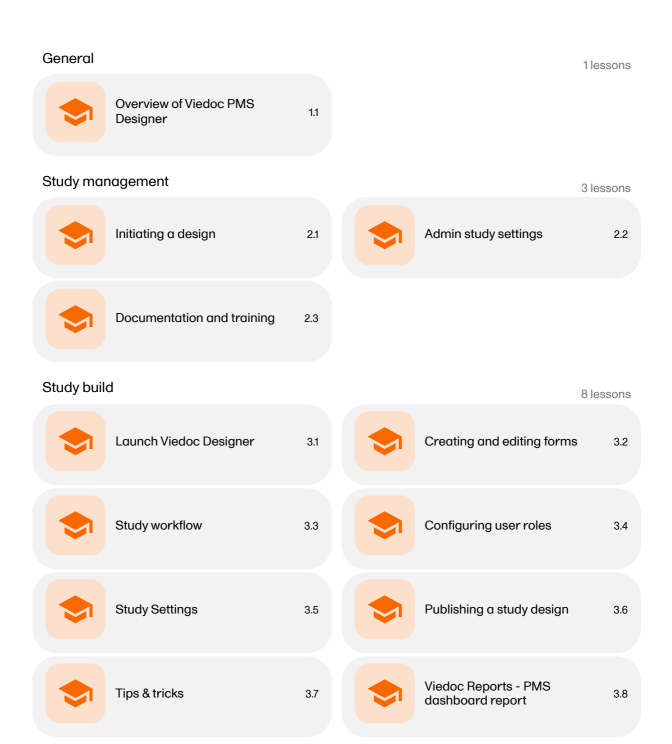
viedoc learning"

Viedoc PMS Designer User Guide

13 Lessons 13 from Viedoc System





4.1



Overview of Viedoc PMS Designer

Overview of Viedoc PMS Designer

Published by Viedoc System 2022-12-14

- 1. Introduction to Viedoc PMS
- 2. Clinic side versus sponsor side
- 3. Booklets
- 4. The send/receive/return process for handling booklets (Kaifu)

1 Introduction to Viedoc PMS

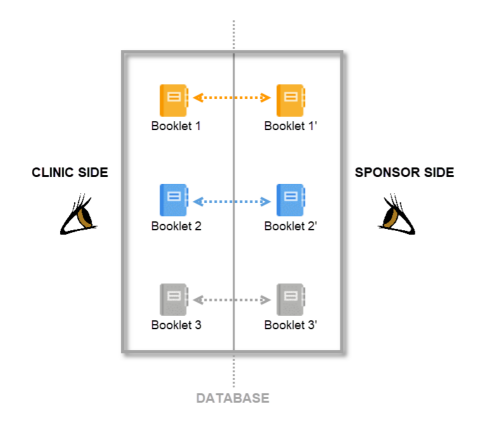
Viedoc PMS is the product on the Viedoc platform that can be used for Japanese Post Marketing Surveillance (PMS) studies. It fulfills all requirements of a PMS study, supports collection of data in booklets, and supports the process of sharing data between site (clinic) and sponsor via the submit-receive-return *Kaifu* function.

PMS studies are built in Viedoc Designer. Viedoc Designer is where you perform the technical part of a study build, either from scratch or by importing a design from a previous project. A design consists of the study forms, the booklets workflow, study roles, and other configurations and settings, as described further in this curriculum. The following sections describe the specific steps essential to build and design a PMS study.

For more information on how to design a study, see Viedoc Designer User Guide.

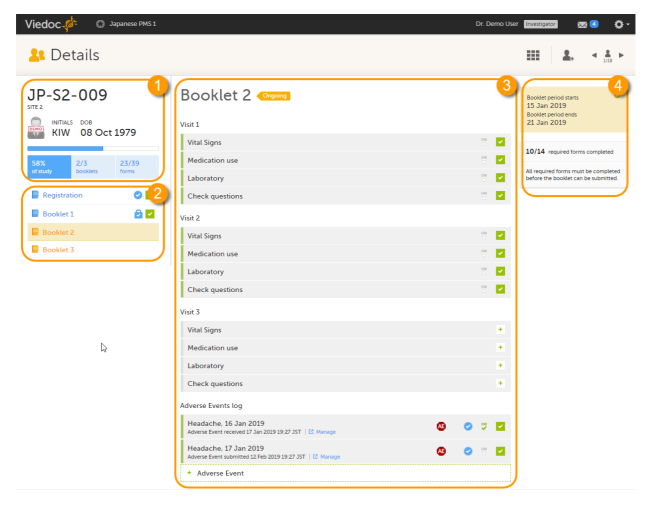
2 Clinic side versus sponsor side

In Viedoc PMS, the database is shared by two different sides, a clinic side and a sponsor side. The database contains two versions of each data set, one version that is displayed to the clinic side users and one version that is displayed to the sponsor side users. On the clinic side, typically the Investigator enters subject data, while on the sponsor side the Data Manager typically reviews the data and archives (freezes) the data.



3 Booklets

By mirroring the data collection and review process via booklets, Viedoc matches the workflow of a Japanese <u>PMS</u> study. A booklet can be seen as a compilation of data being collected during a specific period of time rather than during a specific event date, which is more typical in clinical trials.



- 1.Subject details
- 2.Overview of booklets
- 3.Content of selected booklet
- 4.Details of selected booklet

4

The send/receive/return process for handling booklets (Kaifu)

Sending and receiving data on request is a fundamental requirement for a Japanese PMS study. Viedoc PMS offers support for sending and receiving booklets between site and sponsor, a process referred to as Kaifu. In the Kaifu process, the clinic user chooses when to share data with the sponsor and the sponsor side user chooses when to receive the data. It should be noted that the sponsor side user does not have access to any data entered in a booklet until the booklet has been shared by the clinic through the submit function, and a receive action has been actively performed by a user on the sponsor side.

For more information, see Overview of the submit-receive-return process

For more information about PMS operations for Clinic side and Sponsor side users, please see the following User Guides:

Viedoc PMS User Guide for Clinic side users

Viedoc PMS User Guide or Sponsor side users



Initiating a design

Initiating a design

Published by Viedoc System 2022-12-19

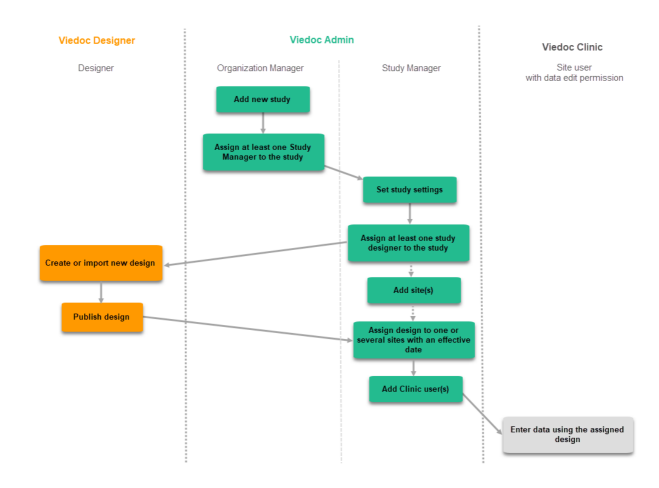
- 1. Configuration workflow for the first study design version
- 2. Building a PMS study
- 3. Adding a new PMS study

1

Configuration workflow for the first study design version

When creating and configuring a PMS study in Viedoc for the first time, you must perform the following steps:

- 1. In Viedoc Admin, the Organization Administrator creates a new PMS study and invites a Study Manager.
- 2. In Viedoc Admin, the Study Manager sets the non-version-controlled common settings and invites a Study Designer. For more information, see <u>Managing users (for Org Admin)</u>
- 3. In Viedoc Designer, the Study Designer creates the first version of the version-controlled settings in the study design and makes the study design available to the Study Manager by publishing the design to Viedoc Admin.
- 4. In Viedoc Admin, the Study Manager creates the site(s) and assigns the first design version to the site(s)
- 5.The Clinic side user can start entering data in Viedoc Clinic using the assigned design.



2 Building a PMS study

To build a new PMS study, you must first add a PMS study to the Viedoc platform and then invite a Study Designer who will build the study in Viedoc Designer.

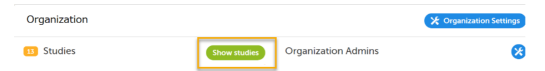
The following sections describe the steps needed when adding a PMS study. For more information on setting up a study, see Viedoc Admin User Guide, <u>Adding a new study</u>.

3 Adding a new PMS study

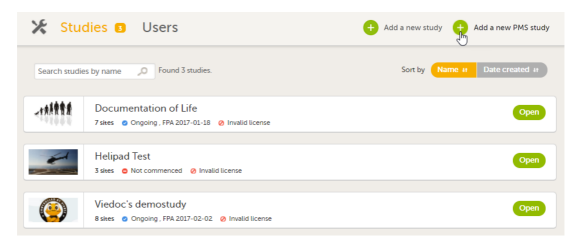
Notel This step is performed by the Organization Administrator

To add a new study:

1 Open Viedoc Admin and select **Show studies** for the organization to which you would like to add a study. The study overview page opens.

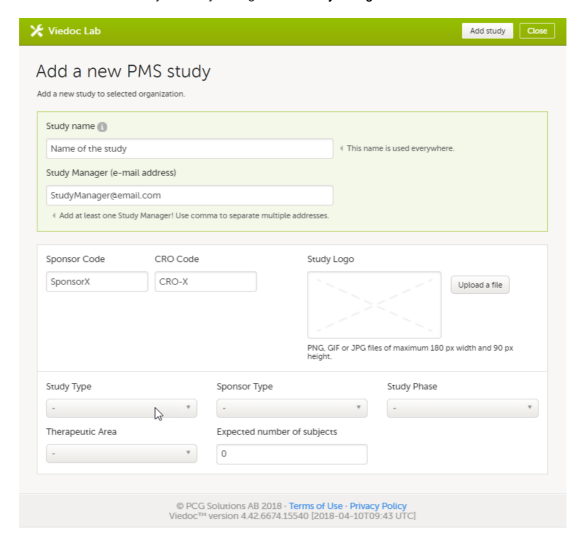


2 Select Add a new PMS study



In the Study name field, enter the name of the Study, and in the Study Manager field, enter the Study Manager's email address.

This information is mandatory. The information in the Sponsor Type and Study Type fields is optional and can be filled in later by the Study Manager under **Study settings**.



4 Select **Add PMS study**. The study will appear in the list of studies on the study overview page. An e-mail is sent to the Study Manager with an invitation to the newly created study.



Admin study settings

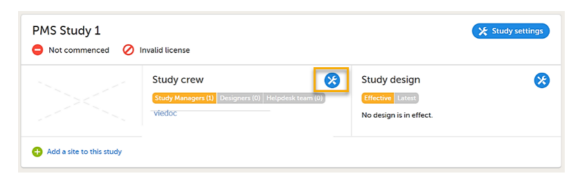
Admin study settings

Published by Viedoc System 2022-12-12

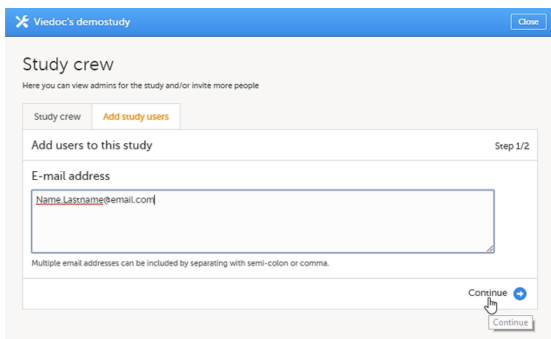
- 1. Inviting the Designer
 2. Completing the study setup
- 1 Inviting the Designer

This step is performed by the Study Manager.

- 1 In Viedoc Admin, open the PMS study to which you would like to invite the Designer
- 2 Select the toolbox icon in the Study crew field. The Study crew pop-up opens.

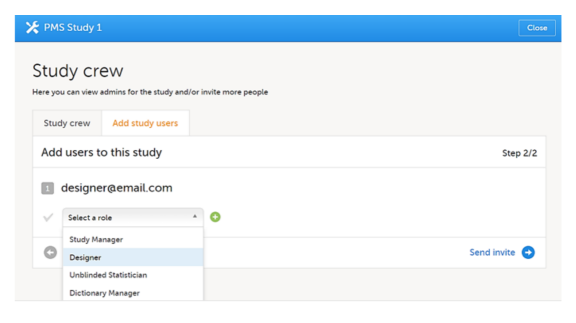


3 In the Add study users tab, enter the e-mail address of the user you would like to invite. Select Continue.



Notel You can invite multiple users at once by adding multiple email addresses in the field. Separate the email addresses with a semi-colon or comma. An invitation email will be sent to the specified e-mail addresses.

4 Select **Designer** from the dropdown menu and then select **Send Invite**. You can add multiple roles by selecting the + icon. You can remove roles by selecting the - icon.



2 Completing the study setup

These steps are performed in Viedoc Admin by the Study Manager.

To complete the study setup:

- 1 Add the study site(s).
- 2 Enter the following study details under Study settings: Sponsor Code, Contract Research Organization (CRO) code, Reference ID, Study Type, Sponsor Type, Study Phase, Therapeutic Area, Expected number of subjects.

- 3 When the Designer has published the study design, <u>assign the study design</u> to the sites in the study.
- 4 Invite users to the different <u>system roles</u> and <u>clinic roles</u>.
- 5 Open the study in Viedoc Clinic and test the study.

Note! These steps are described in more detail in the eLearning lessons under Study Management.



Documentation & training

Documentation and training

Published by Viedoc System 2022-12-14

1. Documentation & training

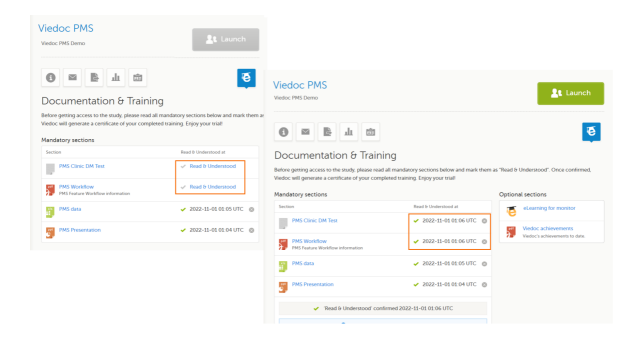
1

Documentation & training

Viedoc allows users to set up the required materials that must be read before accessing and working on a study. Training content can be in Word, Excel, PDF, Power Points, and other document file types, as well as eLearning and web pages.

Training material can be mandatory or optional, and the target audience for each lesson can be customized for each site or role. Additionally, training content can be reset to mandatory for the following booklets, for example, if case protocols and procedures change or are forgotten, during a long period between sessions.

For information about how to set up training documents, see Setting up user documentation and training.





Launch Viedoc Designer

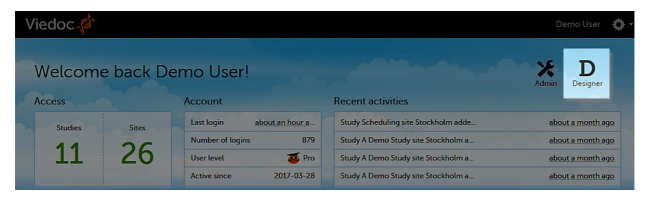
Launch Viedoc Designer

Published by Viedoc System 2022-11-15

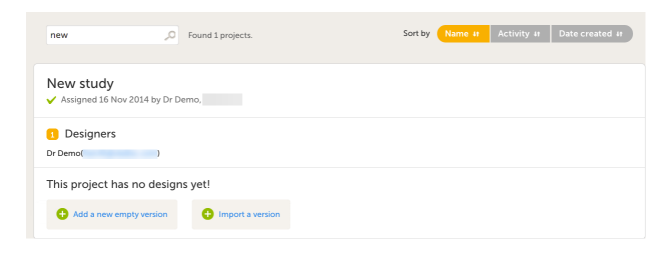
- 1. Launch Viedoc Designer
- 2. Initiating a PMS study design
- 3. Add a new empty version

1 Launch Viedoc Designer

As a Designer, you will be able to access Viedoc Designer once the Study Manager has invited you to a study and you have accepted the invitation. If you have access to Viedoc Designer, you will see the Designer icon in the top-right-hand corner of the main page after you have logged in to Viedoc.



When you select the Designer icon, Viedoc Designer opens and displays a list of all the organizations and studies you have access to as a Designer. If you have several projects, you can search for the relevant project in the Search by study name text field in the top left corner. For more information, see Overview of Viedoc Designer.



2 Initiating a PMS study design

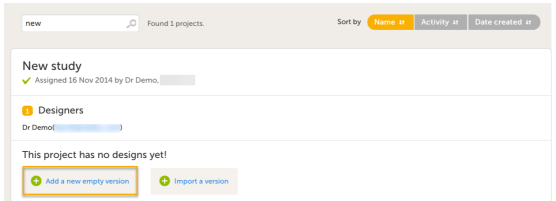
From the Projects page, you can create a new design version, and begin building and designing the study.

Design steps

- 1. Adding a new empty version
- 2. Creating and editing forms
- 3. Creating booklets and setting up the workflow
- 4. Setting booklet periods and alerts
- 5. Configuring the roles for the clinical side and sponsor side
- 6. Configuring the study settings
- 7. Configuring global design settings

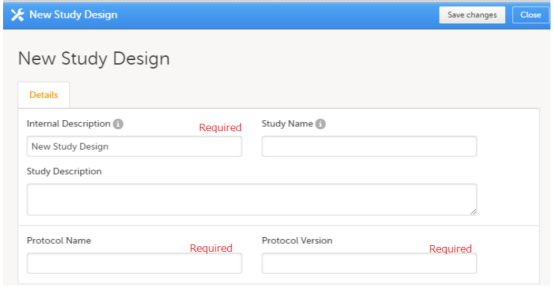
3 Add a new empty version

1 Select Add a new empty version. The New Study Design pop-up window opens.



Notel You can import an existing design using the **Import a version** option. For more information, see <u>Initiating a design</u>.

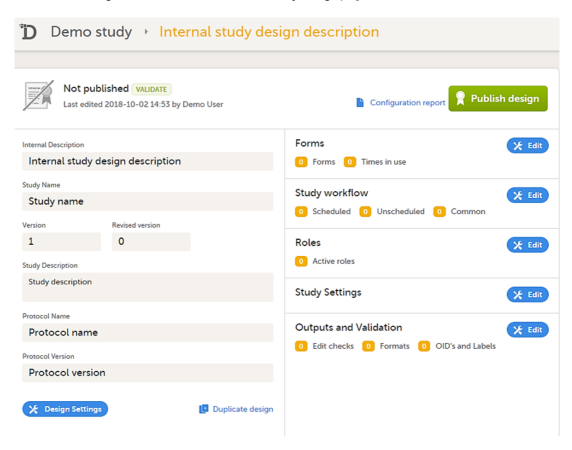
2 In the New Study Design window, fill in the required fields.



In Viedoc Clinic, when the user selects the respective study, only the Study Description will be shown.

Note! All these fields can be changed in new versions or revisions of the study design.

3 Select Save changes. You will be directed to the Study Design page.





Creating and editing forms

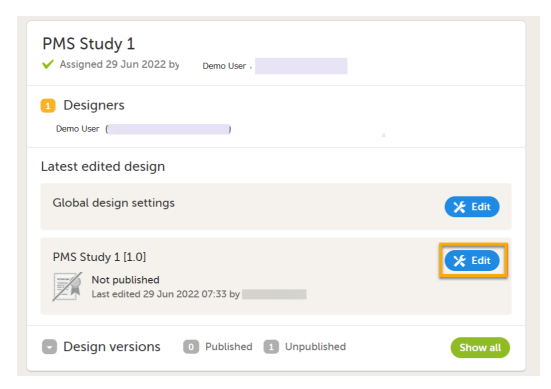
Creating and editing forms

Published by Viedoc System 2023-03-07

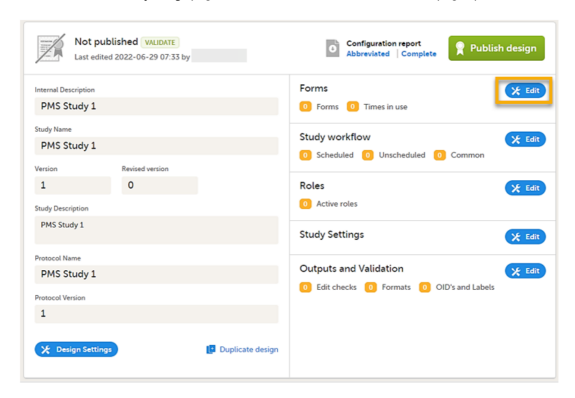
- 1. Creating and editing forms
 2. Allow a form to be copied
- 1 Creating and editing forms

To create a new form, follow the steps below:

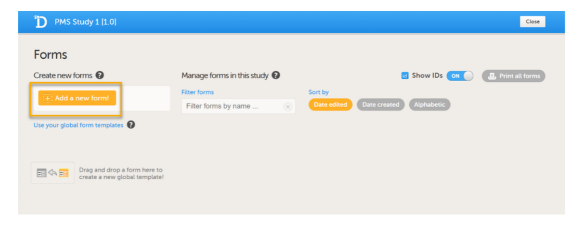
Select **Edit** in the study design name section in the **Projects** window to open the **Overview of study design** page.



2 On the Overview of study design page, select Edit in the Forms field. The Forms page opens.

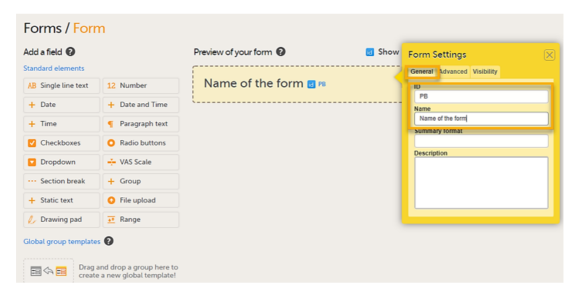


3 Select Add a new form. A new form is created, and then the Form Settings dialog opens.



In the **Form Settings** dialog on the **General** tab, enter the form ID and the name of the form. A form ID must be set for each form and it must be unique. Also, an item ID must be set for each item in the form, and the item ID must be unique within each form.

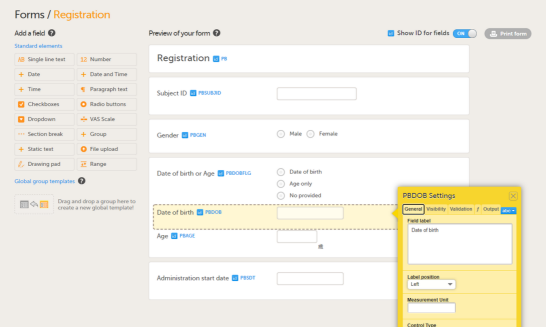
The form ID will be used to identify the form in the database and in the export output. It is also used when referring to the form in JavaScript expressions.



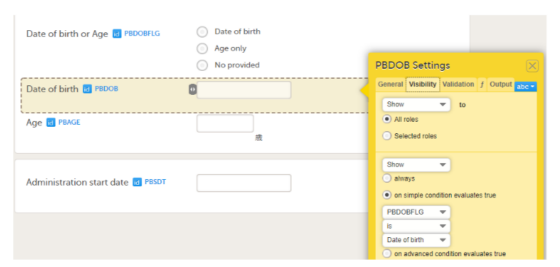
5 Select the required item type from the standard elements menu and click to place all required items on the form and set the required attributes for each.

To add an item to the form:

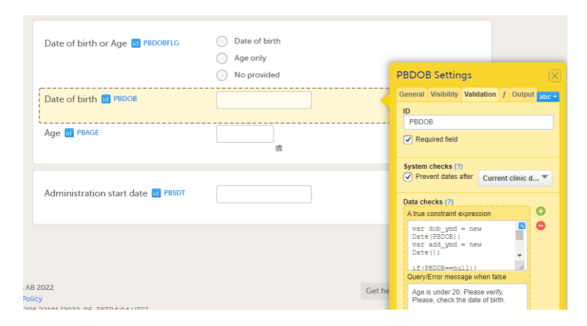
- Select one of the standard elements (items) in the left pane of the form window, or
- Drag and drop an element (item) to an existing item group.



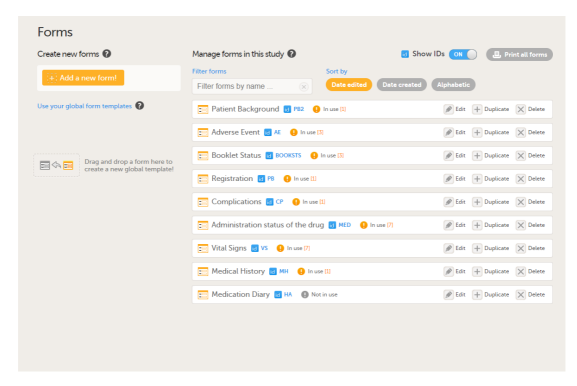
On the Visibility tab, you can set the visibility conditions of the item.



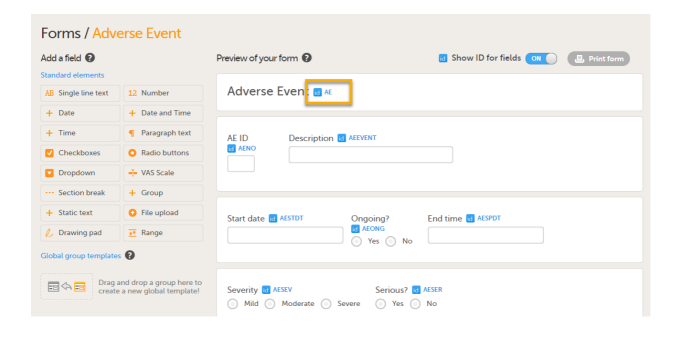
On the Validation tab, you can set the ID of the item and add data checks that validate the item.



- 6 Select **Save changes** to finish creating the form. For more information, see <u>Creating and editing forms</u> and <u>Configuring an item</u>.
- Note! For PMS studies, you should first create the initial registration form and then add all forms required for each booklet. The completed forms for the study are listed on the Forms page.

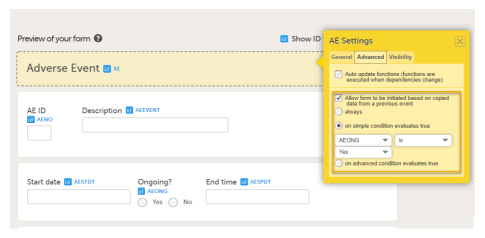


Note! For PMS studies, if you set a form ID to AE, this form becomes an Adverse Event (AE) form. When the AE form is completed and saved on the Clinic side, it can be submitted to the Sponsor side independently of the booklet. For more details about the Adverse Event form, see Adding Adverse Event forms.



2 Allow a form to be copied

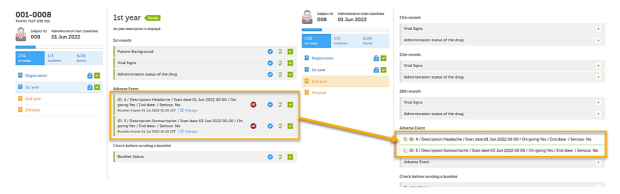
When the option Allow form to be initiated based on copied data from a previous event is selected on the Advanced tab of the form, the data can be copied from a form within one booklet to another instance of the same form within another booklet.



In the example in the figure above, the data is copied when **on simple condition evaluates true** condition and AEONG (ongoing) are selected.

This is a useful feature especially for the Adverse Event form. When an Adverse Event form is added in a booklet and the adverse event continues to the next booklet period, you can add an Adverse Event form in the next booklet with data copied from the Adverse Event form of the previous booklet.

You can add an Adverse Event form which copies data from the adverse event in the previous booklet, when ongoing is selected in the previous booklet.

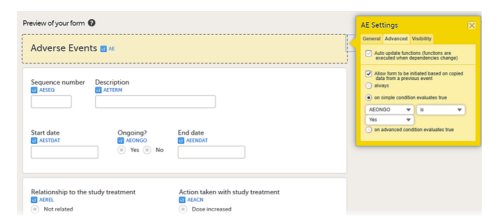


You can select the form to be copyable: always, on simple condition evaluates true, or on advanced condition evaluates true, as shown below.

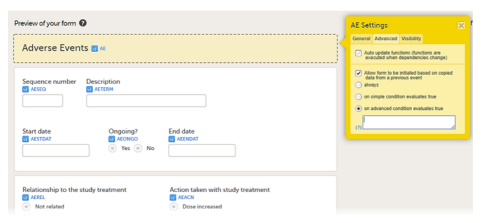
· always



• on simple condition evaluates true. From the dropdown menus, select the item in the form that the condition should be based on, select is or is not, and select the code list item to specify the condition.



• on advanced condition evaluates true. Enter an expression in JavaScript to specify the condition.



For more information, see Allow form to be copied.



Study workflow

Study workflow

Published by Viedoc System 2023-03-07

- 1. Creating booklets and setting a workflow
- 2. Scheduling booklets
- 3. Duplicating the event settings
- 4. Scheduling booklet reminders

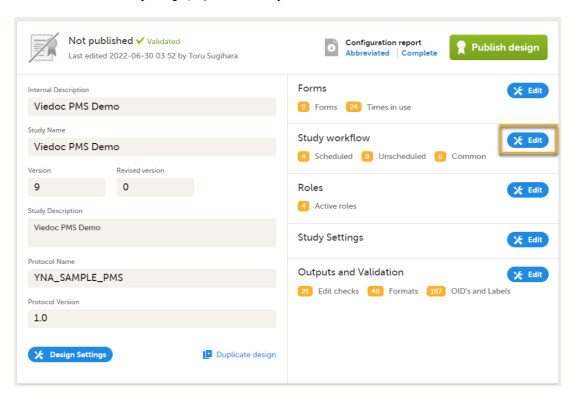
1 Creating booklets and setting a workflow

In a PMS study, adding a single event with the relevant forms to the Study workflow creates a single booklet.

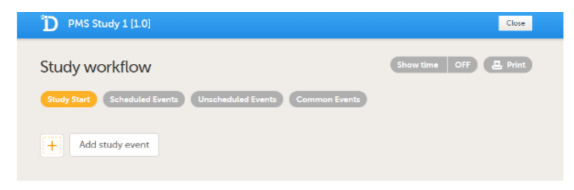
For more information about study workflows, see Study Workflow.

To create a booklet:

1 On the Overview of study design page, in the Study workflow field, select Edit.



2 Select Add study event (add a booklet) in the Study workflow window.

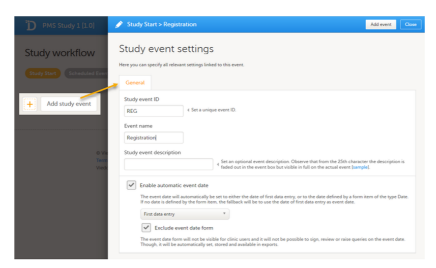


3 Study start booklet

Add a booklet (study event) on the **Study Start** page. This registers basic information whenever a new subject is added.

Select **Add study event** to open the **Study event settings** dialog and add an event. Enter the **Study event ID** and the **Event name**.

The example below shows a booklet with the **Study event ID** - REG, **Booklet (Event) Name** - Registration.



To add an activity to an event, select the (+) icon in the Registration booklet (event) field to open **Activity** settings and enter the activity ID. In the example below, the activity ID: REG_ACT01 has been added.



Add the form(s) you want to include in the **Study Start** event to the activity. In the example below, the **Registration** form has been added to the activity.



Important! Add the study start event first, add an activity and a form, and select **Save**. Make sure that these steps are performed before proceeding with adding other events to the study workflow.

Add the booklets you require in **Scheduled Events**. You can add multiple activities to each event, and the name you give to the activities can be used to separate them within the booklets (for example, the number of months from the study start, Adverse Event, as shown in the example below).



Note! To set a form as repeating, select **Allow form to repeat** for forms that can occur more than once, such as Adverse events.



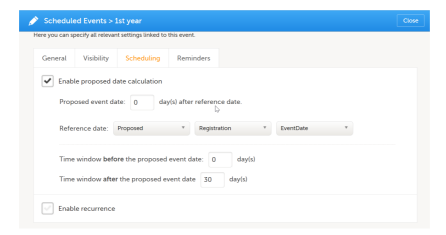
1 Select the pen icon to open the Study event settings dialog to edit the booklet schedule.



- 2 On the **Scheduling tab**, select **Enable proposed date calculation** if you want to activate calculation of a proposed date for the booklet, and configure the following:
 - Set the Proposed event date to n day(s) after reference date, in which n is the number of days between the booklet and the reference date.
 - Define the Reference date by selecting whether it should be based on the Actual or Planned or Proposed event date and select the reference event from the dropdown list.
 - Optionally: Set a time window during which the event can be initiated, by entering the number of days in the fields Time window before the Proposed event date, and in the Time window after the Proposed event date. By default, these are set to 0 days.

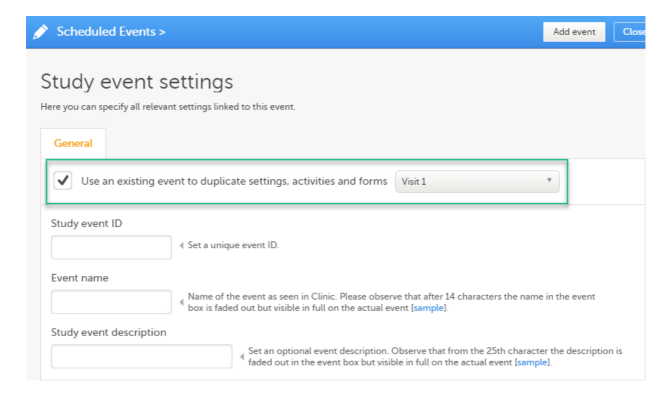
Note! If Actual or Planned is selected for the Reference date, then the scheduled date is calculated on the Reference date entered by the site. However, if the reference event has not been initiated, then the Planned date is used. And if the reference event has not been planned, then the Proposed date of the reference event is used.

In the following example, the **Registration** booklet date is the reference date, and 30 days is the time period during which you can enter data for the booklet.



3 Duplicating the event settings

When creating a new event, it is possible to start configuring it by duplicating the settings of an existing event. This is done by selecting **Use an existing event to duplicate settings, activities and forms** and selecting from the dropdown list the event the settings will be copied from:



After entering the **Study event ID, Event name**, **Study event description** and selecting **Add event**, a new event is created with the same configuration as the selected event with regards to:

- The activities and forms
- Visibility
- Scheduling
- Automatic event date
- Short/long summary format
- Source (clinic/subject initiated)

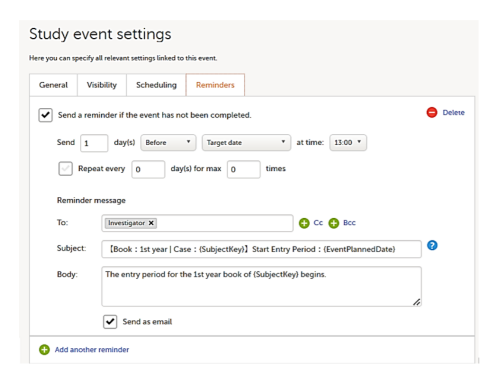
4 Scheduling booklet reminders

On the **Reminders** tab, you can set reminders according to the scheduled settings. The reminders are shown as messages in Viedoc Clinic, and, optionally, sent as emails.

Note!

- The reminders are for incomplete events, and for PMS studies, the definition of a complete event is that the booklet is not in control of the site (it is Submitted/Received/Frozen).
- For PMS studies, only roles with access to Clinic side data are available as recipients.

For more information, see Setting scheduled event reminders



Any static string and/or the following variables can be embedded in both the Subject and Body fields:

- Context form variables can be referenced directly using item ID, for example {SAE}.
- System variables. For a list of available system variables, see the section System variables in the lesson <u>Using JavaScript in Viedoc.</u>
- Other variables can be referenced using the format EventId.FormId.ItemId, for example {SCR.PATINFO.SEX}.

Note! The item values included in the message are visible for all the users with the defined roles in To:, Cc:, and Bcc: without respecting the role visibility settings.



Configuring user roles

Configuring user roles

Published by Viedoc System 2023-04-25

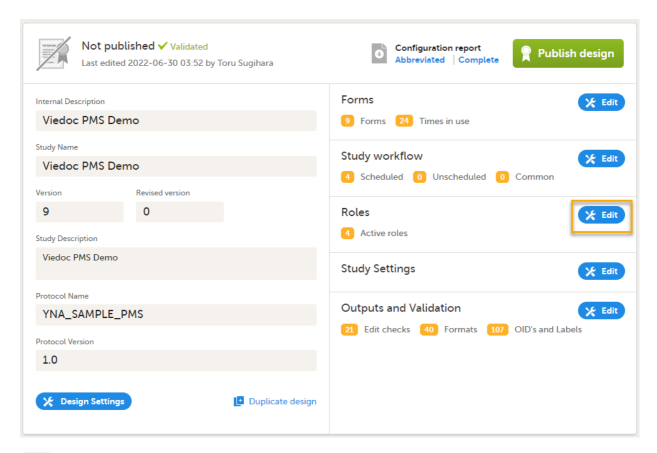
1. Configuring user roles

- 1.1 The Roles page
- 1.2 Configuring roles
- 1.3 User rights
- 1.4 Using predefined roles

1 Configuring user roles

1.1 The Roles page

Roles are configured on the <u>Roles page</u>. In Viedoc Designer, on the **Overview of study design page**, select **Edit** in the **Roles** field to open the **Roles** page.



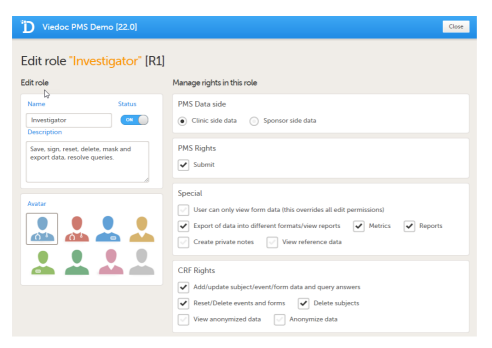
1.2 Configuring roles

The **Roles** page allows you to edit, view, add, copy, and delete user roles. Select the Pen icon to open the Edit roles menu and set the rights for each role.

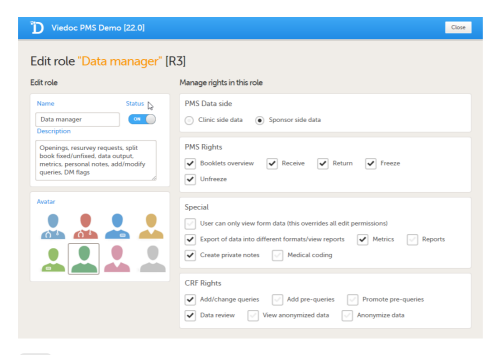


For PMS studies, there are Clinic side and Sponsor side **Edit role** pages where you can choose to edit roles for either the Clinic Side or the Sponsor side. For more information about how to edit these pages, see <u>Configuring roles</u>.

Clinic side Edit role page



Sponsor side Edit role page



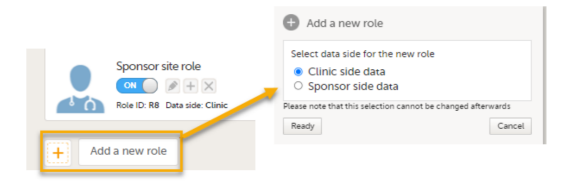
1.3 User rights

The following rights can be selected:

PMS Data side

- Clinic side data
- Sponsor side data

Notel You can only select a data side when you add a new role. The PMS Data side can NOT be changed after a role has been added



PMS rights - booklet permissions

Clinic Side:

Submit: User can submit booklets

Sponsor side:

- Booklets Overview: User can see Booklets overview on the study start page
- Receive: User can receive booklets
- Return: User can return booklets
- Freeze: User can freeze booklets
- Unfreeze: User can unfreeze booklets

Special rights - rights that give access to specific features

- User can only view form data (this overrides all edit permissions)
- Export of data into different formats/view reports
- Metrics

Reports (only visible when Metrics is selected). For export/download rights in Viedoc Reports, the user
must also have "Export of data into different formats/view reports" selected. The rights may not be
applied directly due to the 24-hour data sync in Viedoc Reports.

Note

For Demo sites to work with Viedoc Reports, a user must be invited with direct site access (not through All sites site aroup)

For Production sites, Viedoc Reports is available for all site groups (All sites and the country-specific groups).

- Create private notes
- Medical coding, and if selected (Sponsor side):
 - Perform medical coding
 - Approve medical coding
- View reference data, and if selected (Clinic side):
 - Edit reference data
 - Publish reference data

CRF rights - rights regarding adding/editing/saving data and queries

Clinic Side:

- Add/update subject/event/form data and query answers
- Reset/Delete events and forms
- Delete subjects
- View anonymized data
- Anonymize data

Sponsor Side:

- Add/Change queries
- Add pre-queries
- Promote pre-queries
- Data review
- View anonymized data
- Anonymize data

1.4 Using predefined roles

By default, a set of predefined roles is set up by the system, and it can be modified for your study. The default roles and default permissions for the Clinic side and the Sponsor side are listed in the following tables:

Role	PMS rights	Special rights	CRF rights
Investigator (Clinic side)	Submit booklets	- Export of data into different formats/view reports	- Add/update subject/event/form data and query answers - Reset/delete events and forms - Delete subjects - Anonymize data
CRC (Clinic side)		Export of data into different formats/view report	- Add/update subject/event/form data and query answers -Reset/Delete events and forms - Delete subjects
Data Manager (Sponsor side)	- Booklets overview - Receive booklots - Return booklets - Freeze booklets - Unfreeze booklets	- Export of data into different formats/view reports - Metrics - Reports - Create private notes	- Add/change queries - Data review

	I		
Role	PMS rights	Special rights	CRF rights
MR (Sponsor side)		- The user can only view form data (this overrides all edit permissions) - Export of data into different formats/view reports - Metrics - Reports	
Sponsor (Sponsor side)		- The user can only view form data (this overrides all edit permissions) - Export of data into different formats/view reports - Metrics - Reports	
Data puncher (Clinic side)	Submit booklets	Export of data into different formats/view report	 - Add/update subject/event/form data and query answers - Reset/Delete events and forms - Delete subjects
Reference Data Editor (Clinic side)		- The user can only view form data (this overrides all edit permissions) - View reference data - Edit reference data - Publish reference data	
Regulatory Inspector (Sponsor side)		- The user can only view form data (this overrides all edit permissions)	- View anonymized data



Study Settings

Study Settings

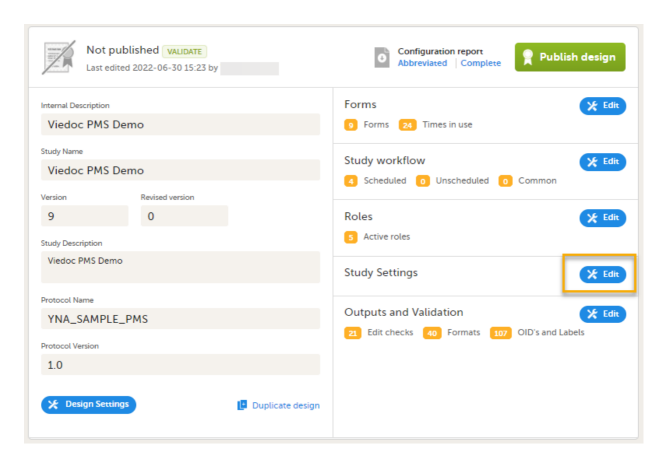
Published by Viedoc System 2023-03-07

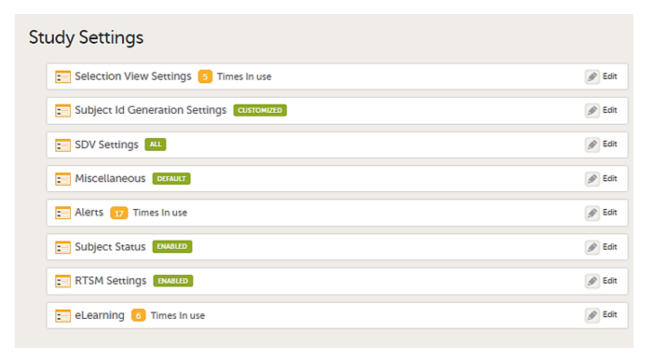
1. Introduction

- 1.1 Selection View Settings
- 1.2 Subject ID Generation Settings
- 1.3 SDV Settings
- 1.4 Miscellaneous
- 1.5 Alerts
- 1.6 Subject Status
- 1.7 RTSM Settings

1 Introduction

On the Study Settings page, you can configure other settings useful for your study.

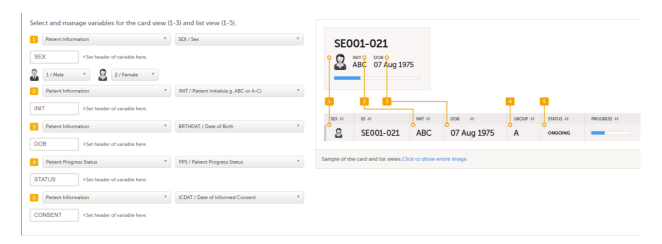




Below is a summary of each menu. For more information, select the relevant page links.

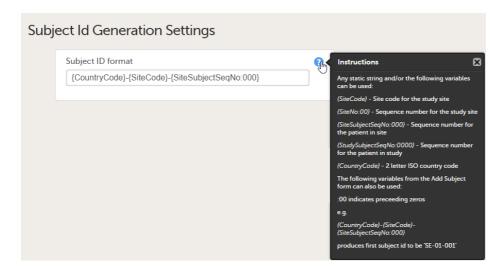
1.1 Selection View Settings

The information to be displayed on the subject card is set on the <u>Selection View Settings page</u>. The subject cards are displayed on the <u>Selection page</u> in Viedoc Clinic.



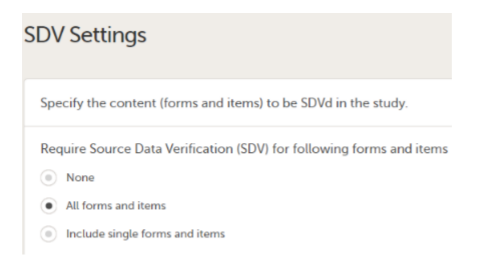
1.2 Subject ID Generation Settings

The format for the Subject ID, used to identify a subject within the system, can be configured on the <u>Subject ID</u> <u>Generation Settings page</u> under **Study Settings > Subject ID generation settings**. The Viedoc default configuration consists of the country code followed by the site ID and finally the consecutive subject ID. This can be changed by modifying the contents of the text field:



1.3 SDV Settings

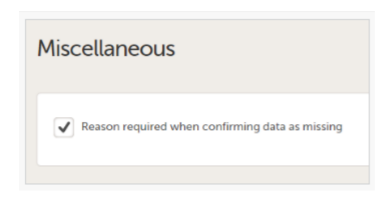
Note! Not available for PMS studies. The Source Data Verification (SDV) setting enables you to choose which forms and items to require SDV in your study.



1.4 Miscellaneous

The Miscellaneous section is for various settings that don't fit in anywhere else.

Currently you can choose to enable/disable the need for entering a reason when a field is left blank, that is, when confirming data as missing in Viedoc Clinic.



1.5 Alerts

By setting up an alert in your study you can notify users about important occurrences in the data. You can set up alerts that are issued in defined conditions (for example, in case of a Serious Adverse Event). For more information. see Alerts.

Note! For Japanese PMS studies, there is a setting where you define which type of change that will trigger the alert. For more information, see <u>Alert triggers for Japanese PMS studies</u>

1.6 Subject Status

The subject status calculations are used in Viedoc in the following places:

- Metrics in Clinic
- <u>Exporting data</u> in Microsoft Excel Open <u>XML</u>, <u>CSV</u> and <u>ODM</u>
- Viedoc Reports

In the Subject status settings, the following statuses are defined:

- Screened subjects
- Enrolled subjects
- Completed subjects
- Withdrawn subjects

For more information, see Subject status

1.7 RTSM Settings

Viedoc offers support for randomization. (Not available for PMS studies).

Subjects can be randomized using either: static randomization: randomization based on a randomized list, or dynamic randomization (Pocock and Simon): randomization based on an algorithm. For more information, see RTSM Settings.



Publishing a study design

Publishing a study design

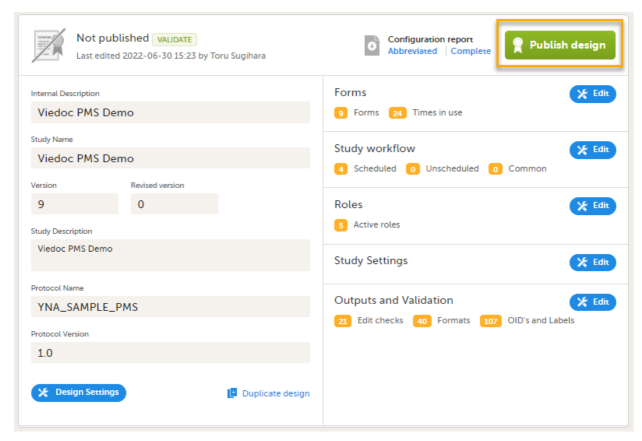
Published by Viedoc System 2022-11-15

1. Publishing a study design

1

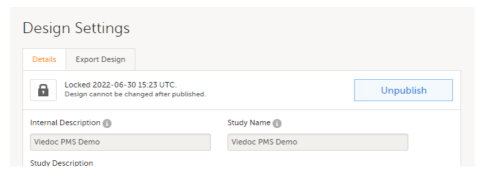
Publishing a study design

When you are ready with the design, select **Publish**. This will first validate the design, and if no errors are detected, this version will be locked for editing and published, becoming available in Viedoc Admin. All the study design settings are, however, available in view mode.



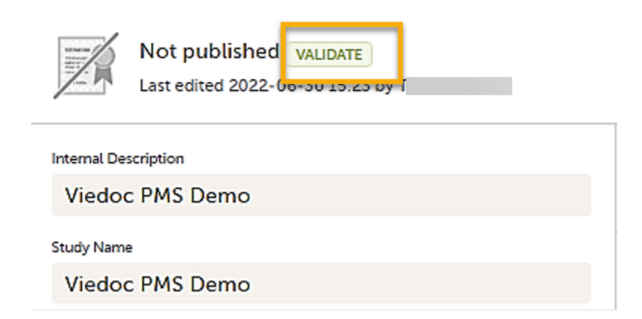
Note! A published study design can be unpublished and unlocked only if it has not yet been assigned to any site(s) in Viedoc Admin.

To unpublish a design, Go to Design Settings and select Unpublish:



For more information, see: <u>Unpublishing a study design</u>.

Note! You can validate the system to check for any study design errors by selecting the **VALIDATE** icon in the top left corner of the study design page. For more information, see <u>Validating a study design</u>.





Tips & tricks

Tips & tricks

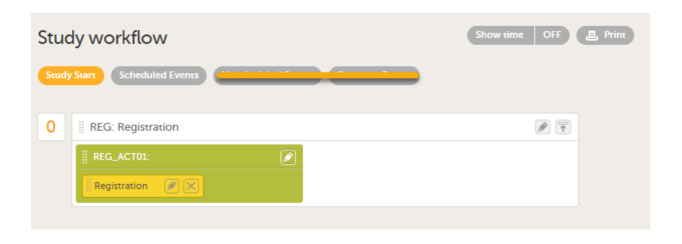
Published by Viedoc System 2022-12-14

- 1. Unscheduled and common events
- 2. Adding Adverse Event forms
- 3. Managing Adverse Events independently
- 4. Setting alerts

4.1 Alert triggers for PMS studies

1 Unscheduled and common events

Important! Unscheduled Events and Common Events are not available for PMS studies although they are visible on the **Study workflow** page.

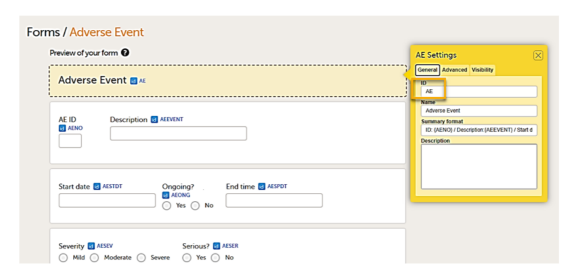


2 Adding Adverse Event forms

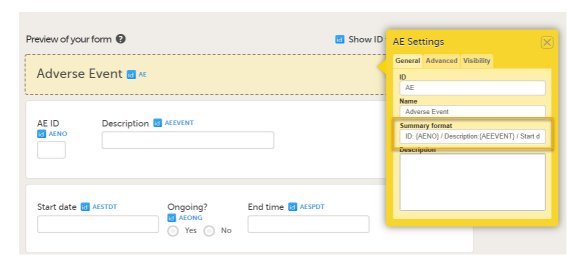
To add an Adverse Event form:

1 Create an Adverse Event form with the form ID as AE and add it as a repeating form in a booklet. A form with an ID set to AE becomes an Adverse Event (AE) form.

2 In AE Settings, on the General tab, set the ID of the AE form to AE.



3 Set the format to be displayed in Viedoc Clinic in the **Summary format** field using the IDs of the items on the form.



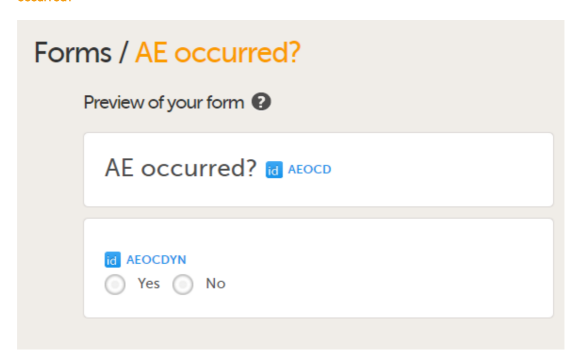
Select the pen icon on the form in the study workflow and select **Allow form to repeat**. This allows you to add multiple adverse events to a booklet.



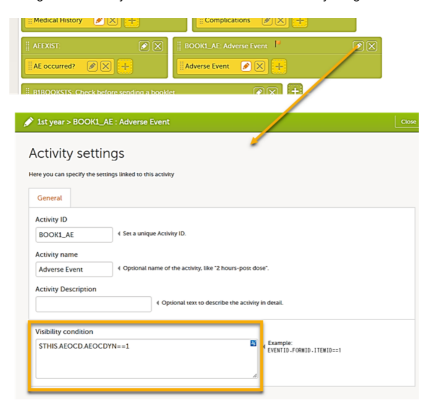
Note! All forms in the booklet, including the Adverse Event form, must be completed in order to submit the booklet. Therefore, the Adverse Event form should appear only when an adverse event occurs.

One way to implement this is described in the steps below:

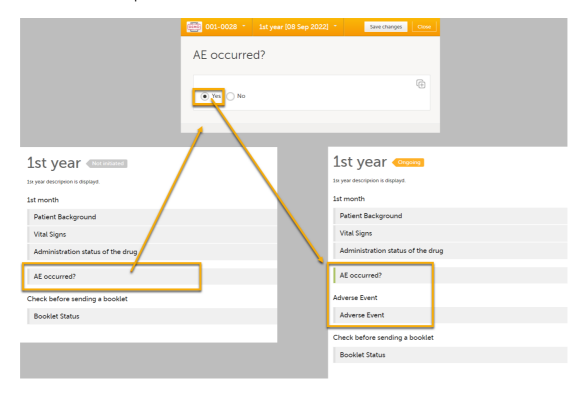
1 Add a form to check if an Adverse Event occurred during the booklet period by adding the form AE occurred?



2 Configure the visibility condition of the Adverse Event activity using the added form item value.



3 If you perform these steps, the Adverse Event form will become visible when an adverse event occurs, as shown in the example below.



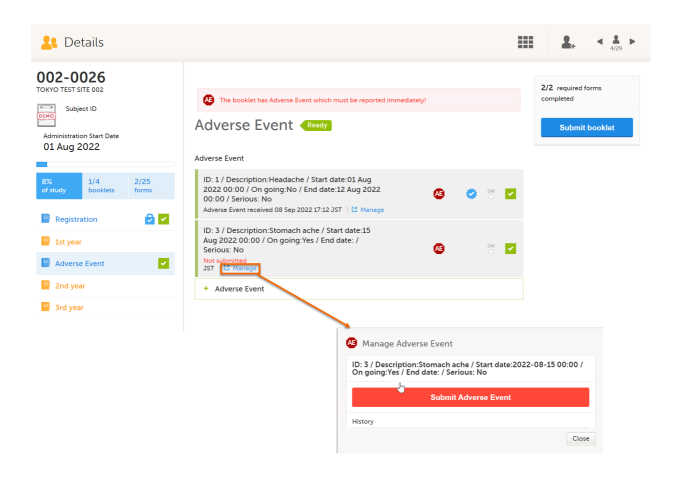
3 Managing Adverse Events independently

If you want to manage adverse events independently from other forms, you can do that by adding one booklet that contains only an AE form. You can report multiple adverse events at any date and time by selecting **Allow form to repeat**.



In Clinic, each time you add an adverse event, you can submit each adverse event individually.

You must submit by selecting **Manage** on the Adverse Event form. If you submit the booklet by selecting the **Submit** booklet button, the booklet is locked and you cannot add more adverse events.



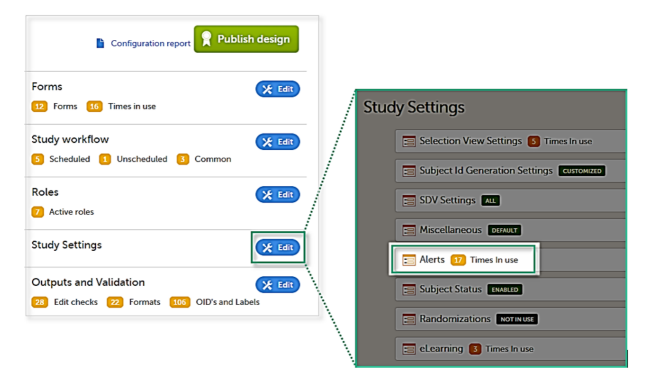
4 Setting alerts

You can add alerts to a study to notify users about important events.

Viedoc allows users to set up alerts that are issued under certain conditions (for example, when a severe adverse event occurs). It can be used to notify the Sponsor side that the Clinic side has submitted a booklet, or to notify the Clinic side that the Sponsor side has returned a booklet to request a re-investigation.

See Alerts for more information.

Setting up the alerts is done in Viedoc Designer, under Study Settings > Alerts:



The Alerts page displays a list of existing alerts (if any) and allows you to add new ones, as illustrated below:

- The Add new button that allows you to create a new alert by opening the alert details page.
- The Edit button directs you to the alert details page, where you can see or edit the alert.

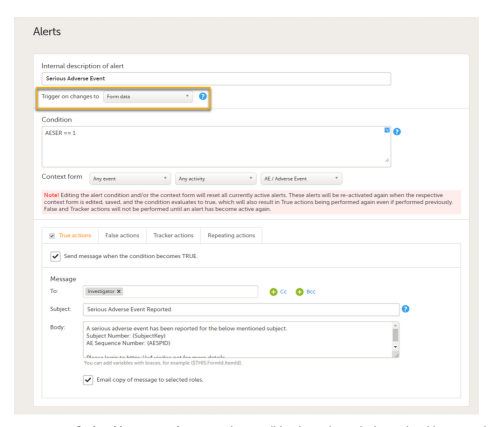


4.1 Alert triggers for PMS studies

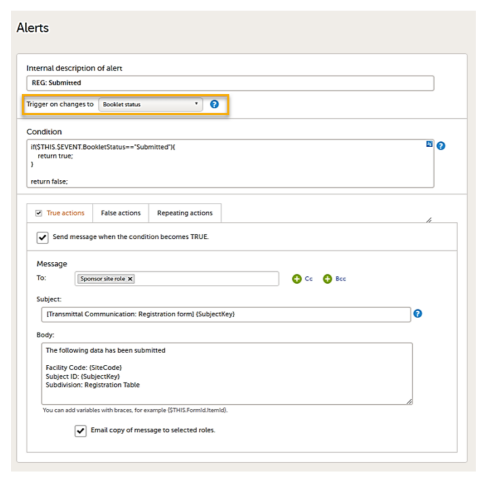
On the Alert details page, you can define which type of change will trigger the alert. There are two options:

• On **context form data** changes - the condition is evaluated when the selected context form below is saved. This option is the default for all existing alerts.

Sponsor users do not receive the message until the Sponsor side receives the booklets with settings that meet the alert condition.



• On **booklet status changes** - the condition is evaluated when a booklet status is changed. The booklet that changes its status will then be the context.





Viedoc Reports - PMS dashboard report

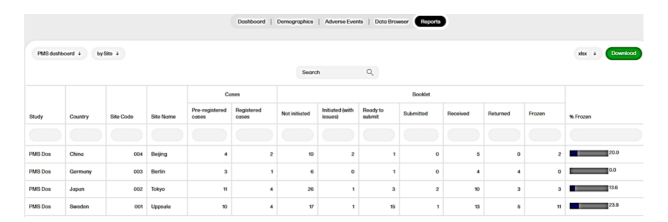
Viedoc Reports - PMS dashboard report

Published by Viedoc System 2022-11-15

1. Introduction 2. Settings

1 Introduction

The PMS dashboard report is available for Japanese PMS studies only. This report allows data to be sorted to focus on the booklet status by site, subject, booklet, booklet history, or timelapse. The **PMS Dashboard** page gives users snapshots of the registration and study progress with terms specific to Japanese PMS studies.



For more information, see PMS dashboard and Japanese PMS studies

2 Settings

The following settings are required in order to use Viedoc Reports:

In Viedoc Admin, on the Study settings page, select Enable Viedoc Reports.

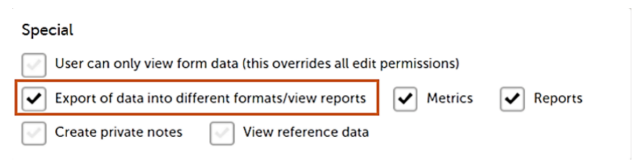


 In Viedoc Designer, select Metrics and Reports on the Roles page for the Clinic role that will be using the report.

Special

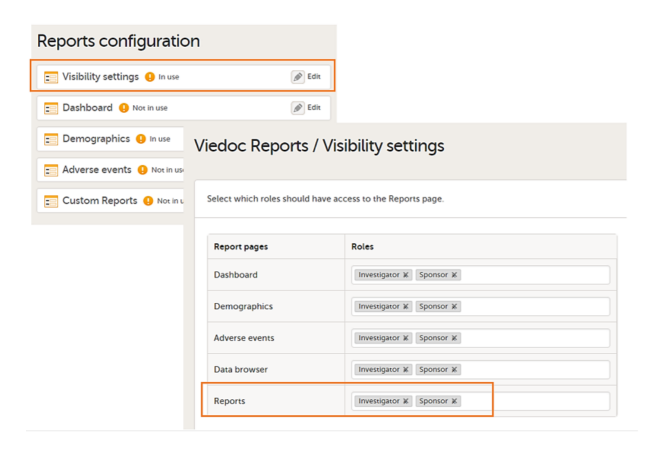
User can only view form data (this overrides all edit	permissions)		
Export of data into different formats/view reports	✓ Metrics	✓ Reports	
Create private notes View reference data			

To be able to download report files, the user also needs the permission Export of data into different formats/view reports.



Notel The export is allowed only if the export permission is applicable to all the assigned sites.

In **Global design settings > Reports configuration > Viedoc Reports / Visibility settings**, add the roles who have access to PMS Reports.





Assigning a study design to production sites

Assigning a study design to production sites

Published by Viedoc System 2023-03-07

1. Assigning a study design to production sites



Assigning a study design to production sites

When the Study Designer has added all booklets to Scheduled Events, and published the study design, the study will appear on the production environment in Viedoc Clinic.

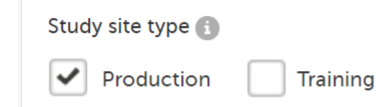
The Study Manager assigns the designs to sites in Viedoc Admin. See <u>Assigning a study design</u> for more information. When the study design is assigned to one or several sites in the study, the study is available in Viedoc Clinic and all booklets are then available for the sites to enter data.

Once a study is on the production server, it is possible to configure the sites to operate in one of the following modes:

Training (demo) mode only: does not require a license, and the data is saved on demo/training only. This is to be used for the test sites only.

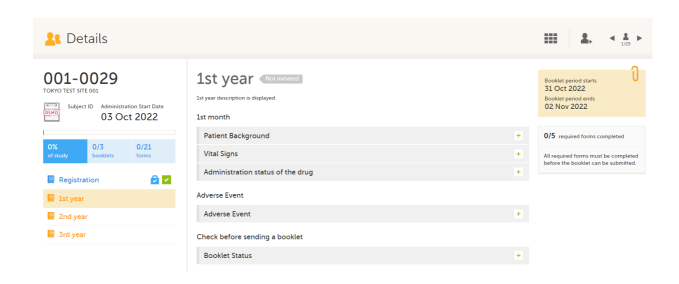


Production mode only: used for the production sites, that is, real sites where real data will be entered, not for testing purposes.



Both training (demo) and production modes: This is **not** recommended, see <u>Training(Demo) vs Production mode</u>.

Your study is now in production, and you can start work on the site.



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