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VISTA REMOTE DATA CAPTURE EORTC ELECTRONIC DATA CAPTURE SYSTEM

https://rdc.eortc.be

USER GUIDE

(FOR TRIALS WITH ELECTRONIC QUERIES)

VERSION 4.4 June, 2014

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Version history - RDC user guide for studies with electronic queries

REVISION HISTORY			
Version	Brief Description of Change	Author	Issue Date
1.0	Initial Release	Antoine Briffaux	04/12/2003
1.1	Update of RDC form approval process and screenshots	Antoine Briffaux	22/06/2004
2.0	Reorganization of document and update to RDC 2	Joke De Wever	11/03/2008
3.0	Update to RDC 3 (Queries and Data Correction)	Joke De Wever	23/04/2009
4.0	Update to RDC 4 (minor changes: save incomplete/complete + request for missing forms not applicable)	Joke De Wever	08/04/2010
4.1	Specification of automatic entry of empty decimals with 0	Caroline Gilotay	29/07/2010
4.2	Options 'Display All Queries', 'Hide Sent Queries' + option 'Print' + option 'Save as pdf'	Katrien Van den Bossche	02/11/2010
4.3	Adaptations to upgrade of software Explanations of queries color codes	Caroline Gilotay	07/06/2011
4.4	User guide imported in Confluence and document split into different pages that can be easily updated electronically in the system. New features added in the user documentation.	Marlies Dictus	June 2014

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Overview

VISTA Remote Data Capture (VISTA RDC) is used for the completion of most case report forms. These forms will be accessible from the EORTC website and have to be completed on-line (https://rdc.eortc.be). The system allows remote users to have access to the EORTC VISTA patient clinical database.

Note: Some CRF (e.g. pathology form, Quality of Life, SAE, etc.), could still have to be completed on paper. Please check for each study which forms (if any) need to be completed on paper. All other forms will have to be completed on-line.

For further information regarding the timepoints of completion of the case report forms please refer to the study specific guidelines for CRF COMPLETION.

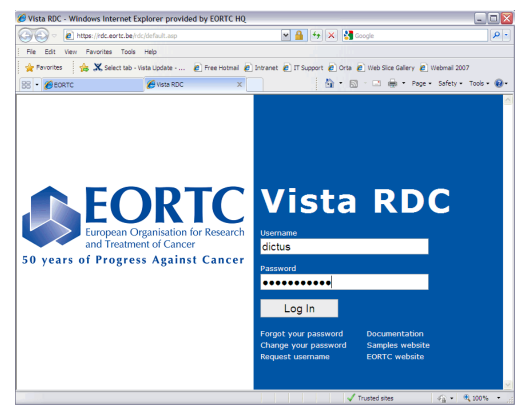
The EORTC VISTA-RDC system provides an electronic version of the CRFs and will enable users to enter and edit data on-line and, once completed, to directly transfer the data into the EORTC clinical database. Sent forms stay visible and accessible in VISTA RDC. After the data transfer, the data reported on the Case Report Forms will be reviewed and validated by the EORTC Data Manager. In case of missing or contradictory data, queries will be raised (per patient) in order to obtain consistent data. These queries will appear in VISTA RDC. Queries requesting an overdue form will result in a 'requested form' appearing on the 'blank' tab. Other queries are displayed in the 'queries' tab.

VISTA RDC is NOT used to register, enroll or randomize patients. For the registration, enrollment or randomization of your patient, please go to: h ttp://orta.eortc.be (via Internet at anytime). An ORTA user guide is available on the weblink.

Access the RDC system

Access to the EORTC VISTA RDC system is possible at the following internet address: <u>https://rdc.eortc.be</u>/ or through the EORTC website (htt p://www.eortc.org), in the 'Investigator's Area'.

On this web page, you will be prompted to enter your EORTC ORTA Username & password (identical to the one used for registration/randomization of patients).



Note: If this screen does not appear or if you have problems after login, your browser might be too old to run the program. Install an upgraded version of your internet browser. These programs are free and available for download on the Internet.

Username & password

All investigators and implicated staff in activated centers will be provided with an ORTA/RDC username and password to access the VISTA RDC system. The type of access that will be given will depend on what was reported on the "Signature and delegation of responsibilities log" document that was provided to the EORTC HQ at time of site activation. Some users might have the right to complete, approve (sign) and send forms while other users might only be authorized to complete forms, without having the possibility to approve and send them.

Important note: forms that are not approved and not sent <u>are not visible</u> to the EORTC Data Managers. It is thus important to approve and send the forms as quickly as possible after completion in order to avoid delay in data validation by EORTC.

Users who are already member of the EORTC can use their existing ORTA/RDC username and password. For the other users, ORTA/RDC usernames and passwords will be attributed using the signature log.

If needed however, a username and password can be requested on the RDC website (https://rdc.eortc.be; 'Request username'). An Account request form has to be completed with the details of the person, institution and study number. Your new username and password will be sent to you by e-mail within a few days.

Note: **Passwords expire after 90 days**. However, you don't need to connect every 3 months to change your password. When you connect with an expired password, the system will ask you to choose a new one. Changing the password periodically is a security feature of the system. Choose whatever password you like, but with **at least 6 characters**. A combination of alphanumeric characters is preferable.

In case you have forgotten your password, a link is provided to request it. Your password will be sent to your email address.

Study Identification

Once logged on, you can begin the actual RDC process. The first step is to identify the EORTC study.

The study identification screen prompts you to select a study number. Only the studies for which you have fulfilled the necessary requirements for participation will be listed in the drop-down menu.

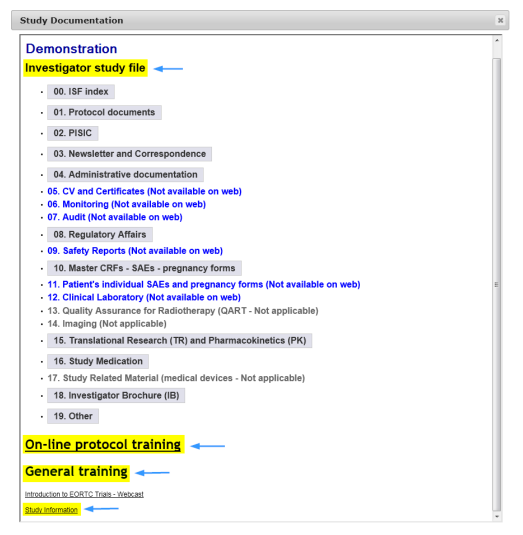


Study documentation

After selecting the study, information on the study can be retrieved by clicking on 'Documentation' in the left part of the screen.

S	tudy Demonstration	-	Manag	je Patients		
9	Study	Your	Pati	ents		
	Documentation Metrics	SeqID	Code	Birthdate	Chart	Registration
	Patients 2	1	INZ	8-Jan-1943		11-Mar-2009

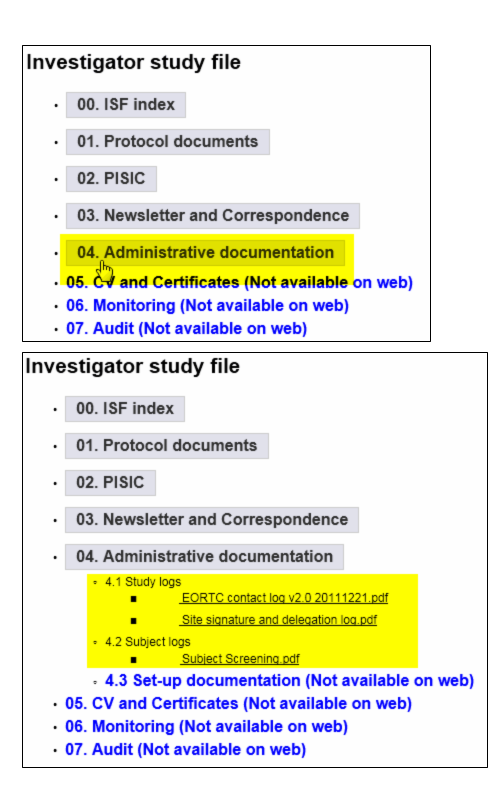
You will be redirected to the study documentation page where you can access the investigator study file, on-line protocol training (if available), general training on EORTC clinical trials activities and the study information page.



Investigator study file

This part is showing the structure and contents the investigator study file should have on site. The investigator study file consists of all the documents which should be kept at the site in the investigator study file (e.g. protocol documents, CRF completion guidelines, IB, ...). The general documents (not site specific) are directly available on-line on this study documentation page. This is the case of **blank paper forms** such as quality of life or SAE forms, **CRF completion guidelines**, which can thus be easily consulted during data entry in RDC.

By clicking on the grey bars, additional information appears and the PDF documents can be opened directly and printed for filing.



On-line protocol training

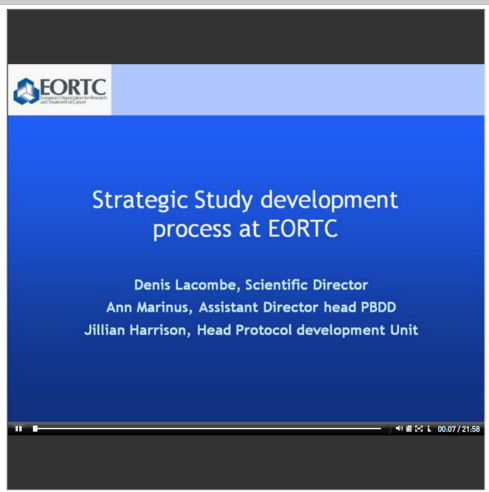
When available for the study, on-line protocol training can be followed from the study documentation page by clicking on the link 'On-line protocol training'. Different training slides can be accessed and when the training is finished, the trainee can confirm online that the training has been followed and a training certificate will be provided by email.

Study Documentatio	'n			х
European Organisati and Treatment of Car	on for Research neer			EORTC AISBL / IVZW Avenue Mounierlaan, 83/11 Bruxelles • Belgie Belgique +32 2 774 16 11
Training 1. Protocol 2. Translation 3. Drug Suppl 4. Data Mana	Biological TR SIV slides v1. Biological TR SIV slides v2.0 Yorug Supply and Handling S gement Jata Management (1) SIV sl Jata Management (2) SIV sl Jata Management SIV slide Data Management SIV slide Data Management SIV slide Data Management SIV slide	121002.pdf 0 20120406.pdf 0 20130311.pdf IV slides v1.0 20120406.pd IV slides v1.1 20130311.pd lides v1.0 20120406.pdf lides_Form specific v1.0 20 s v2.0 20130313.pdf s_Form specific v2.0 20130	<u>f</u> 120406.pdf	
Name	Firstname	Function	Email	Institution number
I confirm You will receive a trainin Thank you.	g confirmation by email.			

General training

The general training on EORTC clinical trials activities consists of a webcast training which is recorded from the one day introduction to EORTC trials. Specific presentations on EORTC's study development, study activation, patients safety, randomization and data management, ... can be viewed directly from the link.

Docun	nentation				X
	ORTC ean Organisation for Research reatment of Cancer		Brussel 1	EORTC AISBL / IVZW Avenue Mounierlaan, 83/11 200 Bruxelles • Belgie Belgieue +32 2 774 16 11	
	One	Day I	ntroduction to EORTC Tria	ls	
entatives.	The purpose of this in	troducto	members (investigators, data managers, research ry workshop is to give guidance for participating ir ve audience in October 2010.		5.
u using a l			ript to be enabled and the latest version of the Ma d please enable it now. Otherwise, please update		
	Title	Length	Content	Presenter	≡
1.	Introduction	35 mins	General Introduction, structure and activities, scientific strategy and perspectives	Francoise Meunier (Director General)	
2.	Study Development Project and Budget Protocol Review	22 mins 9 mins 8 mins	Strategic Study Development Process at EORTC Project and Budget Development Protocol Review Process / Protocol Development	Denis Lacombe (Scientific Director) Ann Marinus (Head of Project and Budget Development) Jillian Harrison	
	Study Activation	12 mins	Study Activation Timelines	Head of Protocol Development)	
3.	Trial Methodology	70 mins	Phase I, II and III trial design, randomization, selection of endpoint, sample size, statistical methods, reporting, monitoring and publication	Richard Sylvester (Senior Statistical Scientist)	-



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Study information page

The study information page contains information on the specific protocol (trial status, targeted sample size, drug, study staff, ...).

Study Documentation		×
Demonstration		•
Trial Status	Closed for recruitment	
Dates	Date of activation: 18-Jul-11 Date Step1 close: 05-Aug-13	
Data management at EORTC		
Phase	3	
Randomized trial	Yes	
Туре		
Targeted Sample size	EORTC Groups: 587 - All Groups: 587	E
Number of steps	2	
Drug	Pazopanib Blind trial medication	
Study Staff	Name (Study Coordinator) - Royal Marsden Hospital - Sutton, Surrey, Sutton Name (Statistician) - EORTC Headquarters, Brussels Name (Data Manager) - EORTC Headquarters, Brussels Name (Project Manager) - EORTC Headquarters, Brussels, Name (Clinical Research Associate) - EORTC Headquarters, Brussels, Name Clinical Research Associate) - EORTC Headquarters, Brussels Name Clinical Research Associate) - EORTC Headquarters, Brussels Name Clinical Research Associate) - EORTC Headquarters, Brussels	
Type of cancer	Lung	
Participating Groups	EORTC Lung Cancer Group(Coordinating Group)	
Protocol summary	Cancer.gov (PDQ) ClinicalTrials.gov	
NCT number	NCT01208064	
EudraCT	2010-018566-23	
Protocol documents in		Ŧ

Metrics

After selecting the study, the number of patients randomized/registered (step 1) in the study can be seen by clicking on 'metrics' in the left part of the screen.

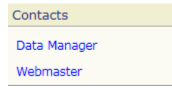
Study DemoMonitored	t 💌	Manag	e Patients	
Study	Your	Patie	ents	
Documentation Metrics	SeqID	Code	Birthdate	С
Study Metrics				
Patients				
Registered 4				
Ineligible 0				

!! Please do not take the 'ineligible' number into account, as this is not applicable for studies using randomization by ORTA.

Request for help

Would you encounter a technical problem with VISTA RDC, please contact either the EORTC webmaster or the study team. For protocol related issues you can always contact the EORTC Data Manager.

Both the webmaster and the EORTC Data Manager responsible for your trial can be directly contacted by e-mail via the following links in the left part of the screen:



Select a patient

Once you have selected the study number, a table listing the patients enrolled by your institution will appear.

The table 'Your Patients' shows the patient's SeqID allocated at the end of the registration/randomization procedure, the patient code, date of birth, chart number (not applicable anymore since 01/01/2008) and date of registration. By default, the table is sorted by Seqid. To sort by code, date of birth or date of registration, click on this field in the header of the table.

Study Demonstration	-	Manag	ge Patients			
Study	Your	Pati	ents			
Documentation Metrics	SeqID	Code	Birthdate	Chart	Registration	Click on the header to sort
Patients 2	1	INZ	8-Jan-1943		11-Mar-2009	
with requested forms	3	AGED	26-Mar-1957		11-Mar-2009	
with incomplete forms 2 with complete forms with queries		g 1 to 2	of 2 entries		Fir	rst Previous 1 Next Last

By default all the patients are shown. It is also possible to make a selection of patients based on the following criteria:

- Patients with requested forms
- Patients with incomplete forms
- Patients with complete forms
- Patients with queries

The numbers shown in the left panel are the number of patients in the selection.

Open the patient's file

To open a patient's file, you may either select the patient in the table or type in the SeqID number (followed by the Enter button) in the SeqID field on the left part of the screen.

Your Patients

SeqID	Code	Birthdate	Chart	Registration	
1	INZ	8-Jan-1943		11-Mar-2009	Select patient in the tag
2	AGE	10-Mar-1932		11-Mar-2009	
3	AGED	26-Mar-1957		11-Mar-2009	
4	HAHA	25-May-1944		11-Mar-2009	

OR

Study Demonstration	•	Manag	ge Patients		
Study	Your	Pati	ents		
Documentation Metrics	SeqID	Code	Birthdate	Chart	Registration
Patients 2 with requested forms	1	INZ AGED	8-Jan-1943 26-Mar-1957		11-Mar-2009 11-Mar-2009
with incomplete forms (2) with complete forms with queries	Showing	g 1 to 2	of 2 entries		Firs
Patient					
SeqID	- Type th	e SeqID) number		

Selecting the patient SeqID will give you access to the patient's file. This on-line file compiles all forms for the selected patient (Blank Forms,

Incomplete forms, Complete Forms and Sent forms) as well as the unresolved queries.

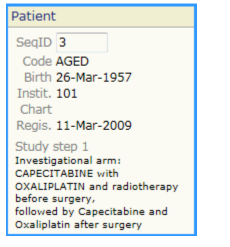
Entries shown in the table

A maximum of 25 entries is shown in the table with 'your patients'. With the buttons 'previous' and 'next' you can go the previous/next list of entries. With the buttons 'first' and 'last' you can go the first/last list of entries.

Your Patients

Patient identification

The patient's identification can be found on the left of the screen.



Code: the patient code (chosen at registration, 4 characters at maximum)

Birth: the patient's date of birth (dd/mm/yyyy)

Instit: the EORTC reference number for the institution that registered/randomized the patient

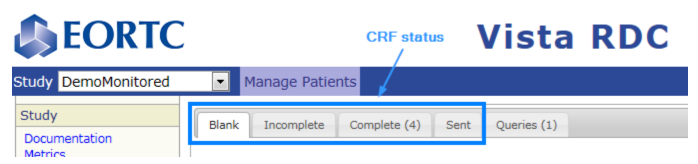
Chart: the patient's chart number (not applicable anymore since 01/01/2008)

Regis: the date the patient was registered in the trial (for the first step)

Study step 1: the treatment that was allocated during the registration/randomization of the patient for step 1 (depending on the study multiple steps can be displayed)

Forms status

Five tabs are available in the **center of the screen**. Four of them are related to the CRFs status while the last one is dedicated to the unresolved queries of the patient.



Blank forms

This tab contains 2 subsections (requested forms and other forms). If a form has not been sent in due time, it will be requested in the section '**req uested forms**'. Form code, name and reason for request will be displayed. The form can be created by clicking on 'create'. Once the form has been created, the request will disappear from the list. If a requested form cannot be completed, click on 'not applicable', enter the reason and click on 'Save'. The request will disappear.

All forms remaining available for completion are displayed in the 'other forms' section. In the 'Instructions' it is reported when the form has to be completed.

The number next to 'Blank' (in the tab) corresponds to the number of requested forms.

Blank (1)	Incomplete (2)	Complete	Sent	Queries
Requeste	ed Forms			

Code	Name			
2	ON-STUDY form	create	Please complete on study form.	Not applicable

If the form is not applicable because of death, end of study treatment, withdrawal of consent: click on 'Not applicable' and provide a comment. If the form cannot be completed for another reason (e.g.: visit not done): create the form, enter the theoretical date and explain the reason in the comment boxes at the end of the form.

Other	Forms		
Code	Name		Instructions
13	Follow-up form	create	This form has to be filled out at EACH follow-up visit (every 6 months for 5 years).
3	PACE form	create	This form should be completed before first chemoradiotherapy if patient is >= 70 years old.
4	Laboratory form	create	Please complete this form: - at baseline (pre-treatment) - weekly during the preoperative chemoradiotherapy (+ at the end) - before start of each postoperative chemotherapy cycle - at the end of last cycle

Incomplete forms

When created, a new form has an "incomplete" status. A form can be saved as 'incomplete' at any time.

Blank	Incomplete (1)	Complete (4)	Sent	Queries	(1)
Code	Name	Class. date	e Las	t modif.	
5	ADVERSE EVENT for	m 17-Sep-201	2 17-5	Sep-2012	edit - view

Complete forms

Once all required data on a form has been reported, the form should be saved as 'complete'. 'Complete' forms must be sent to the EORTC database by users that have the right to sign forms. You can edit complete forms as long as they are not yet sent.

Blank (2) Incomplete (8)	Complete (12)	Sent Qu	ueries (3)		
Code	Name		Class. date	Last modif.	Approve	
5	Hematology Form		20-Jan-2010	14-Jul-2010		edit - view
70	Adverse Event Form		13-Jan-2010	14-Jul-2010		edit - view
6	Biochemistry Form		12-Jan-2010	14-Jul-2010		edit - view
5	Hematology Form		6-Jan-2010	14-Jul-2010		edit - view
6	Biochemistry Form		6-Jan-2010	14-Jul-2010		edit - view

Sent forms

Once a form is approved and sent, it is made visible to the EORTC Data Manager. The form arrives with an on-hold status in the EORTC database. Forms that have been sent can no longer be edited; however it is possible to Create an electronic Data Correction.

In the sent tab also the sent paper forms are shown (e.g. Quality of Life forms). The forms that are entered and sent through VISTA RDC (

E	Blai	nk (15)	Incomplete Complete Sent			
		Code	Name			
	e	crg1	Randomization form			
		930lc	QoL C-30 + LC13			
		905	Health Economics Questionnaire EQ-5D			
	e	5	Vital signs form			
	e	5	Vital signs form			
	é	5	Vital signs form			
	é	5	Vital signs form			

Order forms for a structured view

In each table of all the tabs the forms can be ordered by the following:

- Form code: the reference number/letter code of the form (e.g. form '4' laboratory form/form 'LBHEM' hematology form)
- Name: the name of the form (e.g. Adverse event form)
- Class. date: the classification date of the form (this is a date on the form that is used for the identification in the system, e.g. the classification date of the Follow-Up form, is the date last known to be alive).
- Last Modif. (in tab 'Incomplete' and 'Complete'): the date on which the last modification was done
- Sent (in tab 'Sent'): the date the form was sent

To order the forms within a table according to one of the above-mentioned criteria, click on the related table headers.

Blar	k Incomplete (2)	Complete	Sent	Queries		
Co	de Name	Class	. date	Last modif.	-	Click on the header to sort
12	END of treatment	form 23-Ma	r-2011	23-Mar-2011	edit - view	
cr	1 Eligibility checklis	t 3-Mar	-2009	11-Mar-2009	edit - view	

Create a form

Click on the button 'Create' next to the form you would like to complete in the tab 'Blank'. All available RDC forms are displayed in the table 'Other forms'. <u>Instructions</u> for the timely completion of the forms are mentioned in this table. To complete a form for which a request was issued, click on the button 'Create' in the table 'Requested forms'.

Blank (1)	Incomplete (8)	Complete (2)	Sent
Request	ed Forms	1	
Code	Name		

0000	Trainie -		
6	INITIAL MEASUREMENTS	create	Please comp
tf tho f	iorm is not applicable boy	anusa of	death and

If the form is not applicable because of death, end applicable' and provide a comment.

If the form cannot be completed for another reason theoretical date and explain the reason in the comm

Other	Forms	1	
Code	Name	- ×	
11	ADVERSE EVENT FORM CTCAE V3.0	create	Please comple and at the en
14	FOLLOW UP	create	Please return
15	LATE ADVERSE EVENT	create	This form nee and according follow-up.

Uni-sequential forms

These forms can **only be created once** as they concern data collected once per patient. Once these forms have been created they will no longer be available in the 'Other forms' table (e.g. On study form, End of treatment form).

Sequential forms

These forms can be created as many times as necessary as they concern data requested periodically for a patient. They will always be available in the 'Other Forms' section (e.g. Laboratory form, Adverse Events form, Follow-up form).

Layout of a form

All forms are identified by their name (e.g. Adverse Event form, Follow-up Form, etc) and code (number or letter combination). On each form, several fields appear. Each field is identified (and gueried) by its field number, listed to the left of the guestion.

The VISTA-RDC screens have an identical set-up for all forms. The VISTA-RDC screens are divided into 3 areas (see example below):

- (1) Action buttons that help you manage the VISTA-RDC form i.e. Save & close, Save, Cancel, Delete.
- (2) Identification of the current user (username), the EORTC study number (study), patient SeqID and form.
- (3) Fields to report the requested data.

Complete Save & Close Save Cancel Delete	
Username dictus - Study TrainingRDC - SeqID 1 (AAA) - Form 6 (Biochemistry Form) version 3 - Class. Date 23-May-2013	
Form 6	
1: Trial period O=Pre-treatment I=Concomitant phase O=2=Adjuvant/Maintenance phase O=3=Follow Up Empty Unknown	
111:TEST 🔘 0=no 🔘 1=yes 🖲 Empty 🔘 Unknown	
2: Cycle number (only to be filled in in the Adjuvant/Maintenance phase)	3
3: Was the blood sample taken 💿 0=no 💿 1=yes 💿 Empty 💿 Unknown	
4: Date of sample 23 May 2013 Unknown (if sample was not taken, please fill in the theoretical date that it should have been taken)	
If blood sample was taken:	
5: Blood sample analysed 💿 1=in your center 💿 2=in another laboratory (all LLN / ULN fields to be filled in) 💿 Empty 💿 Unknown	
6: If in another laboratory, specify (name + city)	

Fields to report the requested data

- Date: should be reported in dd mmm yyyy format e.g. 12 Feb 2014 (day, month and year are a seperate box and a dropdown list appears when you type in the field). If the date is not available, tick the checkbox 'unknown'.
- Question with predefined answers (labeled question: e.g. a no/yes question): radiobuttons allow to select the appropriate answer. By
 default the answer will be empty. If the answer is not available, select 'Unknown'.
- Numeric question (e.g. value of irradiated dose): unit and format (number of integers and decimals) is displayed next to the field.
- Text field: enter text in this field or tick the checkbox 'unknown' if the answer is not known.

Edit a form

Forms with an **incomplete or complete** status **can be modified** after creation ('edit'). Data reported on forms that have been **sent** to the EORTC database can **no longer be modified directly in the system** (only 'view'). In case data needs to be modified on 'sent' forms, please Create an electronic Data Correction. These corrections will be implemented by the EORTC Data Manager.

Edit an unsent form

Once a form is created you can enter the data requested. Answer each question by typing your answer in the field allocated to the question or by selecting the answer.

Activate the cursor in the answer field by clicking on it. You can go through the questions by either using the **TAB** button on your keyboard or by repositioning the cursor with your **mouse**. You can go through radiobuttons answers by using the arrow keys.

Answer fields in the RDC forms have 2 levels of reporting importance.

- Mandatory data
- Other data

Mandatory data

The requested data may not remain empty.

- You can save a form as incomplete when mandatory data is missing
- The form can be set to a **complete** status if all mandatory data have been completed: if you try to save a form as complete when mandatory data is missing, a warning message will appear under the fields which need to be completed before the form can be saved as complete (see example below).

1: Trial period This is a requi	 0=Pre-treatment 1=Concomitant phase 2=Adjuvant/Maintenance phase 3=Follow Up Empty Unknown
2: Cycle numbe	r [##] Unknown
(only to be f	illed in in the Adjuvant/Maintenance phase)
3: Was the bloc	od sample taken 🔘 0=no 🔘 1=yes 💿 Empty 🔘 Unknown
This is a requi	red field.

Other data

These data can not always be completed because:

- The requested information is not yet available
- e.g. "Date of first documented progression of disease" remains empty on the follow-up form until progression was diagnosed.
 The requested information is <u>conditional</u> to the answer of another variable
- e.g. for treatment information: only if there was a modification done, you'll have to provide the reason for this modification. If no modification was done, this box should stay empty.
- The data is <u>not applicable</u> e.g. Some lab values can be entered in 2 units, only one box has to be completed.

Special issues for entering data

General

Unless it is for a clear reason, do not leave applicable fields empty otherwise a query will be issued to ascertain whether information was unknown or simply missing.

Where information is not known, tick the checkbox 'unknown'. For example if a test was not performed or a response or toxicity not assessed or the patient source data are incomplete, do not leave the relevant field(s) empty but tick them as "unknown".

Complete Save & Close Save Cancel Delete Username dictus - Study Demonstration - SeqID 3 (AGED) - Form 12 (END of treatment form) version 3 - Class. Dat	e empty
Form 12	
1: Date of LAST treatment administration [dd/mm/yyyy] V Unknown (CTX, RT or surgery)	
2: Performance status at the END of the last treatment \bigcirc 0 \bigcirc 1 \bigcirc 2 \bigcirc 3 \bigcirc 4 \bigcirc Empty \bigcirc Unknown	
3: Did the patient receive PREOPERATIVE chemotherapy 💿 0=No 🔿 1=Yes 🛇 Empty 🛇 Unknown	
4: If no, reason for no preoperative chemotherapy administration	Unknown

In case the value reported is out of the range defined in the database, a warning **message** will appear. You can correct the value or force the answer by clicking on the button 'Force value'.

8: Observed value	8000 µmol/L [####] 🔲 Unknown	
Value out of rang	e. Do you want to force the value?	Force Value

Text fields

Text fields have to be completed in ENGLISH

Dates

All dates should be reported in dd mmm yyyy format e.g. 12 Feb 2013 (day, month and year are a seperate box and a dropdown list appears when you type in the field). If the date is not available, tick the checkbox 'unknown'.

2: Date hematology sample	12	2	2013	🔲 Unknown
3:WBC 15.00 E+9/L [###.	##]	Feb		

It is not possible to enter partial dates. If the day (and month) is not available, tick the checkbox 'unknown' and put the field number and the available info in the comment fields at the bottom of the form.

Numeric fields

Depending on the laboratory of your institution, some parameters can be expressed in different units: e.g. Hemoglobin can be expressed in "mmol/l" or "g/dl". Choose the field corresponding to the unit used by your laboratory and **leave the other field empty** (no need to flag it UNKNOWN).

- If your laboratory uses other units, make the easiest conversion: e.g. "g/l" to "g/dl".
- If your laboratory provides results in both units, select and document one unit only, but consistently.

Never report values in a different unit than the one(s) provided on the VISTA-RDC form! Warning messages will inform you if the wrong format has been used, informing you about the expected format.



Numbers with decimals

• Only use the number of spaces foreseen in each data field. When excess precision in numbers with decimals is observed, adjust the number according to the following rounding rules (ref: NIST Guide to SI Units):

For rounding xxx,abc where a, b, c, x are figures

- If (c < 5), xxx,abc is rounded into xxx,ab: example: 7,124 7,12
 If (c 5), xxx,abc is rounded into xxx,a(b+1):example: 7,128 7,13
- If some decimals are not filled in, it will automatically filled in with 0 by the system after saving. example: 46,-- 46,00

Collapsible sections

For some studies collapsible optional sections are incorporated into the electronic forms. To access the fields for a collapsed section, click on the panel with arrow.

Concomitant medication form					
1: Has concomitant medication been administered	○ 0=no ○ 1=yes	Please complete a section for each therapy hereunder			
	© Empty ⊚ Unknown				
▶ Medication 1 तीन्					
Hedication 2					
Medication 3					
▶ Medication 4					
Medication 5					
Medication 6					

The fields for that section will appear. As soon as data is completed in these sections or queries are made, the section will always appear as uncollapsed.

Concomitant medication form		
1: Has concomitant medication been administered	0=no 0=res	Please complete a section for each therapy hereunder
	© Empty ⊚ Unknown	
 Medication 1 		
2: Drug name Fluconazole	🔲 Unknown	
3: Route 1=Per os 2=Intravenous 3=Intramuscular 4=Subcutaneous 5=Rectal 6=Transdermal (patch) 7=Other Empty Unknown	w	
4: If other, specify route	🔲 Unkno	wn
5: Dose/Units/Schedule 200mg OD		Unknown
6: Indication for use (a) 1=Adverse event (c) 2=Prophylaxis (c) 3=Other (c) Empty (c) Unknown		
7: Specify indication for use Fungal infection		🔲 Unknown
8: Start date 🕢 01 Mar 2013 🔲 Unkn	own	
9: Ongoing 💿 0=no 💿 1=yes 💿 Empty 💿 Unkno	wn	
10: End date 🕢 day month year 🕅 U	nknown	
Medication 2		
Medication 3		
Medication 4		
Medication 5		
Medication 6		

Comments

At the end of each RDC-form there is a space for writing down **comments.** These fields can be used to mention remarks (example: give reason why a lab value is not available). This could help to reduce the number of queries, but make sure that all information that should be reported in one of the RDC-fields is also mentioned in the appropriate field(s) and not only in these comment fields.

27: Ca++ 10.3 mg/dL [##.##] Unknown					
28: CEA	µg/L [#####.#] <mark>✓ Unknown</mark>				
in case patie	nt receives oral anticoagulant treatment				
29: INR	[##.#] Unknown				
30: If INR not	calculated, PT sec [###.#] Unknown				
Please use th	Please use the following boxes to express any comment you might have on the data of this form				
31: Comment	field 28: CEA value unknown because lab test failed				
32: Comment		 ⊻ เ			
33: Comment					

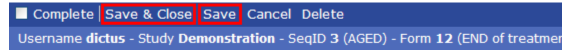
Cancel changes

In case you do not wish to save your changes, click on the 'cancel' button at the top of your screen.

Complete Save & Clo	se Save Cancel Delete
Username dictus - Study	Demonstration - SeqID 3 (AGED) - Form 12 (END of t
Form 12	
1: Date of LAST tr (CTX, RT or sur	ows Internet Explorer
2: Performance sta	Are you sure you want to navigate away from this page?
3: Did the patient	 This will cancel all your changes. Press OK to continue, or Cancel to stay on the current page.
4: If no, reasor	
5: Did the patient	OK Cancel

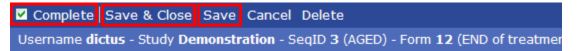
Save a form as incomplete

A form may be saved without being complete and without the mandatory values being reported. To save a form as incompleye, click on the **Save** or on the **Save & Close** button at the top of your screen.



Save a form as complete

A form may be set as complete only if all mandatory questions (required fields) have been correctly answered and if all out-of-range data have been corrected or confirmed and forced. To save a form as complete, tick the field next to 'Complete' and click on the **Save** or on the **Save & Close** button at the top of your screen.



Delete a form

Forms can be deleted by the local Data Manager/Investigator if they were not yet sent to the EORTC (status 'incomplete' or 'complete'). In order to delete a form that has already been sent to the EORTC, the local Data Manager/Investigator should Create an electronic Data Correction. Only the EORTC Data Manager is authorized to delete a form that has already been sent to the EORTC.

Delete an unsent form

A form can be deleted before it has been sent. In case data have already been reported on the form, these data will be deleted as well. To delete an unsent form, click on the 'delete' button at the top of the screen.

Complete Save & Close Save Cancel Delete				
Username dictus - Study Demon	stration - SeqI(3 (AGED) - Form 12 (END of			
Form 12	Y			
1: Date of LAST treatment adm (CTX, RT or surgery)	Message from webpage			
2: Performance status at the E	Are you sure you want to delete this form ?			
3: Did the patient receive PREC	OK Cancel			
4: If no, reason for no preo				

Approve & send a form

Authorization to approve and send forms is given on the signature and delegation of responsibilities log of the study. Forms can be approved and sent from the 'complete' tab.

Users entitled to approve and send forms

Users entitled to approve and send forms will see the message below. Tick the boxes next to the forms you wish to approve, enter your password and click the 'approve & send' button.

Code	Name	Class. date	Last modif.	Approve	
5	Hematology Form	20-Jan-2010	14-Jul-2010		edit - view
70	Adverse Event Form	13-Jan-2010	14-Jul-2010	1	edit - view
6	Biochemistry Form	12-Jan-2010	14-Jul-2010		edit - view
5	Hematology Form	6-Jan-2010	14-Jul-2010		edit - view
6	Biochemistry Form	6-Jan-2010	14-Jul-2010		edit - view
5	Hematology Form	30-Dec-2009	14-Jul-2010	1	edit - view
6	Biochemistry Form	30-Dec-2009	14-Jul-2010		edit - view
8	Concomitant CCI-779 Treatment form	23-Dec-2009	14-Jul-2010	1	edit - view
70	Adverse Event Form	23-Dec-2009	14-Jul-2010		edit - view
6	Biochemistry Form	23-Dec-2009	14-Jul-2010		edit - view
5	Hematology Form	23-Dec-2009	14-Jul-2010		edit - view
5	Hematology Form	Unknown	3-Dec-2013		edit - view

Users not entitled to approve and send forms

Users not entitled to approve and send the forms will see the following message:

Blank (2	2) Incomplete (8) Complete (12)	Sent Qu	ueries (3)	
Code	Name	Class. date	Last modif.	
5	Hematology Form	20-Jan-2010	14-Jul-2010	edit - view
70	Adverse Event Form	13-Jan-2010	14-Jul-2010	edit - view
6	Biochemistry Form	12-Jan-2010	14-Jul-2010	edit - view
5	Hematology Form	6-Jan-2010	14-Jul-2010	edit - view
6	Biochemistry Form	6-Jan-2010	14-Jul-2010	edit - view
5	Hematology Form	30-Dec-2009	14-Jul-2010	edit - view
6	Biochemistry Form	30-Dec-2009	14-Jul-2010	edit - view
8	Concomitant CCI-779 Treatment form	23-Dec-2009	14-Jul-2010	edit - view
70	Adverse Event Form	23-Dec-2009	14-Jul-2010	edit - view
6	Biochemistry Form	23-Dec-2009	14-Jul-2010	edit - view
5	Hematology Form	23-Dec-2009	14-Jul-2010	edit - view
5	Hematology Form	Unknown	3-Dec-2013	edit - view

The Principal Investigator (or an authorized staff member) of this trial should now check, approve and sign the completed forms to send them to the EORTC database.

Save a form as PDF

When opening a form by clicking on 'view' next to the form in the 'Incomplete', 'Complete' or 'Sent' tabs, the button 'Save as pdf' is available.

Click on the button 'Save as pdf' to print the form to PDF. A dialog window 'get pdf' will appear.

Back to List	Print	Save as	pdf
Username va	n den	bossche -	- Study

The pdf version of the form is displayed when clicking on 'here' in the text 'You can get the pdf here'. You can save or print the PDF document.

Get pdf	×
You can get the pdf here	
	Cancel
	Cancel

Print a form

When opening a form by clicking on 'view' next to the form in the 'Incomplete', 'Complete' or 'Sent' tabs, the button 'Print' is available.

By clicking on the button 'Print', the form can be printed.

Back to List Print Save as pdf Username van den bossche - Study De

Answer an electronic query

After sending data, the CRF will be verified by the EORTC Data Manager. In case of missing data or inconsistencies, queries will be generated. These queries will be displayed in VISTA RDC and should be answered in VISTA RDC. You can quickly select the patients with queries to be answered by clicking on the hyperlink 'with queries' on the left part of the screen. The number mentioned in the grey square is the number of patients with queries to be answered.

Study TrainingRDC	• 1	Manage	e Patients		
Study	Your	Pati	ents with	Que	eries
Documentation Metrics	SeqID	Code	Birthdate	Chart	Registration
Institution All	1	AAA	27-Feb-1957		14-Dec-2009
Patients 23	2	aper	27-Mar-1951		13-Jan-2010
with requested forms 6	3	AAB	10-Nov-1956		1-Feb-2010
with incomplete forms 14	4	нн	8-Nov-1953		19-Feb-2010
with complete forms 10 with queries 11	5	MP	29-Sep-1958		2-Mar-2010
	6	AO	15-Nov-1950		19-Mar-2010
Patient	7	GKJ	22-Jan-1982		1-Apr-2010
SeqID	9	POLO	20-Jul-1962		13-Apr-2010
User: dictus	10	BAPH	7-Feb-1950		15-May-2010
	16	GB	21-Jul-1967		20-Jul-2010
Logout	17	mmr	13-Aug-1951		26-Jul-2010
Contacts					

To see the number of pending queries per form, go to the 'Queries' tab. The pending queries can be 'DM' queries or 'DCF' queries.

- · DM queries: raised by the EORTC data manager
- · DCF queries: data corrections raised by the investigator side

Answer queries

To see and answer the DM query/queries on a form, click on 'edit' next to the form. The form will be displayed and the query will be displayed under the field to which the query is related.

nk (2) Incomplete (8) Complete (12) Sent Queries (3)	Queries (3)	Complete (12)	Incomplete (8)	Blank (2)
--	-------------	---------------	----------------	-----------

Queries on Forms and Variables

Code	Name	Class, date	Sent	Pending queries*		Ready	
Code	Name	Class, date	Sent	DM	DCF	to send	
5	Hematology Form	31-Jan-2010	6-Jul-2011	1		1	edit
3	On-Study Form	Unknown	22-Mar-2010	1			edit

*DM: Queries raised by EORTC data manager; DCF: Data corrections

The answer to a query can be a new value or you can confirm the current value.

1: Trial period 0=Pre-treatment
Please check the trial period as the date of sample is after the pre-treatment period. New value: 0=Pre-treatment 1=Concomitant phase 2=Adjuvant/Maintenance phase 3=Follow Up
OConfirm current value
Comment
Ready to send

New value

If you want to enter a new value, tick the radiobutton 'new value' and tick the appropriate radiobutton or complete the new value in the field. Mind using the correct format. After the query has been answered, mark it as 'Ready to send' by ticking the checkbox. Clicking on 'Save' in the header of the form will save the answer(s) and close the form.

1: Trial period 0=Pre-treatment
! Please check the trial period as the date of sample is after the pre-treatment period.
 New value: 0=Pre-treatment 1=Concomitant phase 2=Adjuvant/Maintenance phase 3=Follow Up
Confirm current value
Comment
Ready to send

Confirm current value

In case the answer that was in the field is correct, tick the radiobutton 'confirm current value'. In that case, you must also put a comment to justify why you confirm the value. After the query has been answered, mark it as 'Ready to send' by ticking the checkbox. Clicking on 'Save' in the header of the form will save the answer(s) and close the form.

1: Trial period 0=Pre-treatment
Please check the trial period as the date of sample is after the pre-treatment period. New value: 0=Pre-treatment 1=Concomitant phase 2=Adjuvant/Maintenance phase 3=Follow Up Empty
Confirm current value
Comment The date of sample is wrong (see DCF field 3)
✓ Ready to send

Do not forget to actually send the queries that are marked as 'Ready to send' in order to make them visible in the clinical database (Send electronic query answers / data corrections (studies with electronic queries)).

Create an electronic Data Correction

A **data correction** is a request made by the local study staff as the Investigator, Research Nurse, local Data Manager to ask for a change on data that were already sent to the EORTC study database. Only the EORTC Data Manager is able to modify these forms in the database.

It might happen that you notice that data that were already 'sent' are incorrect and want to correct it. It might also happen that the patient data (code, date of birth) need to be corrected. To do so, you can ask for a data correction.

There are 3 kinds of data corrections:

- On a field level
- On a form level
- On a patient level

Field correction

Select the form in the tab 'Sent' and open it by clicking on 'view'. Next to each field there is a button 'Correct data'.

You can also do a Form correction

Form 70			
1: Trial period	0=Pre-treatment 1=Concomitant phase 2=Adjuvant/Maintenance phase 3=Follow Up	Correct data	
	empty unknown		
2: Cycle numbe (only to be	er filled in in the Adjuvant/Maintenar	Correct data	

When clicking on this button, a dialog field appears and you can select a new value and/or write a clarification in the 'comment' field.

Data Correction for Box No. 1 *
New value O0=Pre-treatment O1=Concomitant phase O2=Adjuvant/Maintenance phase O3=Follow Up Oempty
Comment
>
After saving, don't forget to go to the 'Queries' tab to send it
Cancel Save

Click on 'Save'. The data correction appears below the box. The red background indicates that the data correction was not yet sent. Click on the 'back to list' button at the top of the form to close it. If necessary, the data correction can be edited or deleted in the tab 'Queries'.

Form 70
1: Trial period 0=Pre-treatment
Data Correction New value: 0=Pre-treatment 1=Concomitant phase 2=Adjuvant/Maintenance phase 3=Follow Up Empty
Comment Ready to send Delete

Do not forget to actually send the data corrections in order to make them visible in the clinical database.

Form correction

Select the form in the tab 'Sent' and open it by clicking on 'view'. A comment about the form can be created by clicking on 'Form Comment' in the header of the form.

Back to List	Form Comment Display All Queries Print Save as	s pdf
Username di	$ heta$ - Study ${f Training RDC}$ - SeqID ${f 1}$ (AAA) - Form ${f 70}$ (Advers
Form 70		
1: Trial period	0=Pre-treatment 1=Concomitant phase 2=Adjuvant/Maintenance phase 3=Follow Up	
	empty unknown	

When clicking on this button, a dialog field appears and you can select a new value and/or write a clarification in the 'comment' field.

Form Comment		×
Comment about this forr Please delete this form (form by mistake).		
	Cancel	Save

Click on 'Save'. The data correction appears on the top of the form. The red background indicates that the data correction was not yet sent. Click on the 'back to list' button at the top of the form to close it. If necessary, the data correction can be edited or deleted in the tab 'Queries'.

Save Cancel Display All Queries
Username dictus - Study TrainingRDC - SeqID 1 (AAA) - Form 70 (Adverse Event Form) version
Data Correction Comment Please delete this form (I duplicated the form by mistake).
Ready to send Delete
Form 70
1: Trial period 0=Pre-treatment Correct data

Do not forget to actually send the data corrections in order to make them visible in the clinical database.

Patient data correction

To ask for a correction of the patient data (code, date of birth), click on the tab 'Queries' and click on the button 'Correct patient data' in the section 'Other queries'.

Blank (2) Incomplete (8)) Complete	(12) Sent	Queries (3)		
Queries on Forms and Variables						
Code	Name	Class, date	Sent	Pending queries		
Code	Name	Class, date	Sent	DM	DCF	
5	Hematology Form	31-Jan-2010	6-Jul-2011	1		
3	On-Study Form	Unknown	22-Mar-2010	1		
*DM: Queries raised by EORTC data manager; DCF: Data corrections				S		
Other Queries Save Correct patient data						

A dialog box will appear where you can describe the change to be made. Click on 'Save' to close the dialog box.

Patient Comment		×			
Comment about this patien	t				
The date of birth should be corrected to 28/09/1958.					
20/03/1338.					
	Cancel	Save			

The data correction will be displayed in the section 'Other queries'. The red background indicates that the data correction was not yet sent. If necessary, the data correction can be deleted by clicking on the 'Delete' button. The data correction can be modified by unticking the 'Ready to send' tickbox. Once the modification has been done, mark the data correction as 'Ready to send'.

Blank (2) Incomplete (8)) Complete	Complete (12) Sent		Queries (3)	
Querie	es on Forms and	d Variables				
Code	Name	Class, date	Sent	Pending	queries	
Code	Name	Class, date	Sent	DM	DCF	
5	Hematology Form	31-Jan-2010	6-Jul-2011	1		
3	On-Study Form	Unknown	22-Mar-2010	1		
*DM: Queries raised by EORTC data manager; DCF: Data corrections					S	
Other	Queries	1				
Data Correction						
Comment The date of birth should be corrected to 28/09/1958.						
Ready to send Delete						
Save Correct patient data						

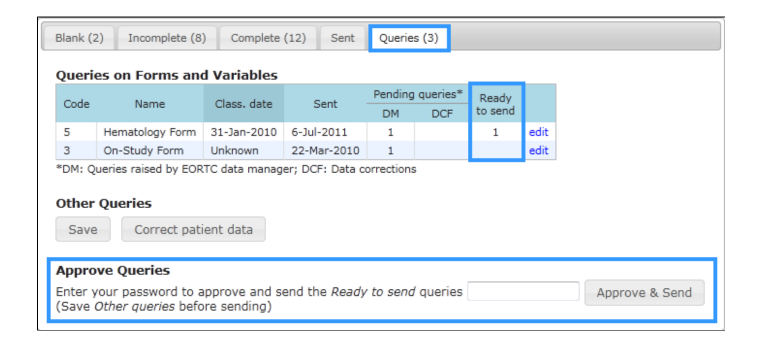
Do not forget to actually send the data corrections in order to make them visible in the clinical database.

Send electronic query answers / data corrections

Persons authorized to approve and send e-CRF, can also send answers to queries / data corrections. The number of queries and data corrections that were marked as 'Ready to send' will be displayed in the column 'Ready to send'. When entering your password and clicking the 'Approve and send' button, all queries and data corrections that were marked as 'Ready to send' will be sent. After sending the answer to a query / the data correction, the query / data correction will disappear from the tab 'Queries' (except for the 'Other Queries'). Query answers / data corrections that have been sent to EORTC can be viewed in the related forms of the tab 'Sent' and their background will be yellow.

Users entitled to send query answers / data corrections

Users entitled to approve and send forms will see the message below.



Users not entitled to send query answers / data corrections

Users not entitled to approve and send the forms will see the following message:

Code Name		Class, date	Sent	Pending queries*		Ready		
Loue	Naitte	class, uate	Sent	DM	DCF	to send		
5	Hematology Form	31-Jan-2010	6-Jul-2011	1		1	edit	
3	On-Study Form	Unknown	22-Mar-2010	1			edit	
Other Queries								
Save Correct patient data								

Display of electronic queries / data corrections

All queries / data corrections can be displayed when opening a form in the 'Sent' tab by clicking on 'view' next to the form.

- (Un)answered queries or data corrections not yet sent will appear highlighted in red
- Answered queries and sent data corrections not yet processed by the EORTC Data Manager will appear in yellow.

Query answers / data corrections that were sent and processed by the EORTC Data Manager do not appear by default. They can be made visible by clicking the button 'Display All Queries' if available at the top of the form. If this button is not available, it means that there are no such "closed" queries on this form. These queries / data corrections can be hidden again by clicking on the button 'Hide Closed Queries' that appears instead of the 'Display All Queries' button once ticked.

Save Cancel Display All Queries
Username $dictus$ - Study $TrainingRDC$ - SeqID $f 1$ (AAA) - Form $f 10$ (Adjuvant Treatment form (TM
Form 10
1: Cycle number 1
Data Correction New value: 2 [##] Comment Ready to send Delete
PATIENT CHARACTERISTICS AT THE START OF THIS CYCLE
2: Weight 58.0 kg Correct data
Please verify the weight of the patient. New value: 60 kg Comment
3: Body surface area 2.00 m2 Correct data
4: Performance status (WHO: 0-4) 2 Correct data

versus

Save Cancel Hide Closed Queries
Username $dictus$ - Study $TrainingRDC$ - SeqID $f 1$ (AAA) - Form $f 10$ (Adjuvant Treatment form (TM
Form 10
1: Cycle number 1
Data Correction New value: 2 [##]
Comment Comment Ready to send Delete
PATIENT CHARACTERISTICS AT THE START OF THIS CYCLE
2: Weight 58.0 kg Correct data
Please verify the weight of the patient. New value: 60 kg Comment
3: Body surface area 2.00 m2 Correct data
: Please verify the body area. Confirm current value
Comment The weight was wrong.

Summary of color codes for queries

Color	Label	Description
Red	Unsent	Query answer / data correction that still needs to be sent
Yellow	Sent	Query answer / data correction that has been sent but not yet processed by the EORTC Data Manager
Green	Closed	Query answer / data correction that has been sent and processed by the EORTC Data Manager