



1. Content

1.1. The Registry will accept prospective registration of interventional clinical trials submitted by Responsible Registrants

| 1.1.1. Does the Registry register trials before the first participant has been recruited? | Yes | No |
|---|----------|----|
| a) Does the Registry query submissions where registration is being sought after the first participant has been recruited? | Yes | No |
| EMA: All trials have to be submitted to the National Competent Authority and loaded into EudraCT before they are authorised, so this query is not applicable. EMA queries late entries by NCAs. | | |
| b) Does the Registry make it clear to Responsible Registrants that prospective registration means that a trial must complete the registration process, <u>and</u> have a trial registration number issued, <u>before</u> the recruitment of the first participant. | Yes | No |
| 1.1.2. Does the Registry register trials that have already recruited the first participant? (Also referred to as retrospective registration). | Yes | No |
| EMA: This occurred in the early days of EudraCT as Member States were implementing legislation. | | |
| 1.1.3. Does the Registry register other types of studies, including observational studies | Yes | No |
| 1.1.4. Does the Registry register all trials submitted by Responsible Registrants. | Yes | No |
| EMA: The Clinical Trials appearing in the EU-CTR are those that meet the publication criteria as required by the EC Regulations. | | |
| a) Does the Registry accept trials submitted by Responsible Registrants? EMA: See above. | Yes | No |
| b) Does the Registry ask the person submitting a trial for registration to verify that they meet the terms and conditions for being a Responsible Registrant before being able to proceed to trial registration. | Yes | No |
| c) Does the Registry use the contact details provided by the Responsible Registrant to verify that those details are correct? | Yes ⊠ | No |
| d) As a minimum, Registries must send an email to the address given and receive a reply from that same address. When possible, the telephone number and/or surface mail address will also be verified in a similar fashion. | Yes | No |
| EMA: This is checked by NCAs who are in contact with the sponsor when authorizing the trial. | · | |
| e) Does the Registry ensure that all Responsible Registrants are associated with an institution or organization? | Yes | No |
| EMA: This happens via NCAs who authorize the trials. | | |





| f) Does the Registry obtain institutional contact details for the Responsible Registrant? g) including name and telephone number of the institution | Yes ⊠ | No □ |
|---|----------|---------|
| 1.1.5. Does the Registry accept studies for registration when the data is submitted as an electronic data file (eg as an xml file) | Yes | No |
| Note: This is not a requirement and is being asked for information only. | | |





1.2. The registry will be open to all prospective registrants (either internationally or within one or more specific countries) (ICMJE requirement)

| a) Which type of study does the Registry accept for registration? | | |
|---|---------|-----------------------|
| | | |
| ☐ Observational studies | | |
| b) If registration is restricted in some way (eg only accepts trials from sponsor, or in a particular health care condition (eg cancer) or interspecify how it is restricted: | om a p | articular) please |
| EMA: Registration is not limited to any particular kind of trial but it of interventional clinical trials and medicinal products. | only ap | plies to |
| c) From which countries does the Registry accept trials for registration: | | |
| EMA: All interventional clinical studies conducted with at least 1 centre conducted completely outside of the EU if they form part of an agreed PIP (st March 2011) are registered in EU-CTR. | | |
| 1.3. The Registry will be able to collect and publicly display the WHO Tria Data Set (TRDS) (ICMJE requirement). | l Regis | tration |
| Does the Registry collect and display, on a publicly accessible web site, all of the items in the WHO Trial Registration Data Set | Yes | No |
| b) Does the Registry have quality control procedures in place to ensure all items in the TRDS contain meaningful data? | Yes | No |
| c) Does the Registry collect the optional TRDS data items? If yes, please specify: | Yes | No |
| EMA: Approval | | |
| 1.4. The Registry will make an effort to keep registered information up-to- | date. | |
| a) Does the Registry permit Responsible Registrants to update information about their trial? | Yes | No |
| Does the Registry have a reminder system to facilitate the submission of updated information by the Responsible Registrants. | Yes | No 🗆 |
| If yes, please state how often the Registry reminds Responsible Registrants to update their data (eg once ever 6 months; once every year): | | |
| EMA: Applicants are required by law to submit substantial amendments and end of trial notices. NCAs are reminded by EMA to complete information — we run data quality checks and these are being increased now that we have data warehouse. | | 77. |
| Note: This is not a requirement and is being asked for information only. c) Does the Registry display the date the trial record was last updated? | Voc | No |
| EMA: Not at the moment but it will be implemented. We are going to add this data as part of a release in the next future. | Yes | No 🗵 |



| Note: This is not a requirement and is being asked for information only. | | |
|--|--|---------|
| d) Does the Registry request updates from Responsible Registrants at least annually until the Registrant has recorded meaningful information about the publication of the trial results (e.g. has listed a citation in a "Publications" field). | Yes | No ⊠ |
| EMA: Applicants are required by law to submit substantial amendments and end of trial notices. NCAs are reminded by EMA to complete information—we run data quality checks and these are being increased now that we have data warehouse. They will be reminded about results once we have enabled these to appear in the Register. | | |
| Note: This is not a requirement and is being asked for information only. | | |
| | | |
| 1.5. The Registry will <u>never</u> remove a trial once it has been registered. | · | |
| a) Does the Registry ever delete a trial record from their database once a registration number has been issued? | Yes | No ⊠ |
| If yes, please explain the circumstances under which a record would be deleted: | | |
| EMA: There is the technical possibility to remove records from the EU-CTR but it applies only under exceptional circumstances due to excessively misleading or fraudulent information, and even in those cases the information may remain public but with warnings attached. The only other case would be in the rare event that a duplicate record entered the system (highly unlikely). | The state of the s | |
| and cyanama (mgm, amma-y). | | |
| b) Does the Registry inform Responsible Registrants at the time of registration that a trial cannot be deleted once it has been registered? | Yes | No |
| EMA: We will put a notice on the website about this. | | |
| c) Does the Registry have clear and transparent processes for dealing with requests to remove a trial record from public view, and have these processes been documented in the SOPs? | Yes | No |
| EMA: We do not remove records – but there is in the new results guidance some information on this. We have a clear process and will document this. | | |
| 2. Quality and Validity 2.1. The Registry will have processes in place to make sure that registered complete and accurate. | d data i | is |
| a) Does Registry staff routinely check all data submitted about a trial for completeness and meaningfulness to ensure that all TRDS fields are populated and comply with the minimum standards? | Yes | No |
| EMA: This process takes place especially at National level when the NCAs perform the review of the CTA application. EMA and NCAs have agreed to introduce a procedure to address this combining routine checks system validation rules and periodic control reports. | | |
| b) If one or more items in the TRDS submitted for registration are incomplete or not meaningful, does Registry staff contact the Responsible Registrant and attempt to obtain complete and | Yes | No |





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|---|----------------------|-------------|
| meaningful data. | | |
| EMA: Same as above | | |
| | | |
| c) Does the Registry's database system apply automated checking | Yes | No |
| procedures (e.g. range checks, logic rules) to data items to facilitate | \boxtimes | |
| validity checking. | E.S | ليبا |
| EMA: Business rules apply when fulfilling the CTA and when loading to | | |
| EudraCT, plus applicants have to submit validation report to NCAs and justify | | |
| | | |
| any deviations. | | |
| a) Does the Registry have processes in place for deciding whether to | | |
| register trials where the Responsible Registrant remains non- | Yes | No |
| compliant with requests to provide complete and meaningful data? | $ \boxtimes $ | |
| compliant with requests to provide complete and meaningful data: | | |
| EMA: Sponsors have to submit a valid application to NCAs in order to get a | İ | |
| trial authorized. | | |
| ordi doctronizodi. | | |
| b) Does the Registry undertake regular internal quality control audits to | Yes | No |
| assess the level of completeness and accuracy of the data collected? | | |
| * | | |
| 2.2. The Registry will have documented Standard Operating Procedures (S | :ODe) | Thoso |
| SOPs will be aligned with the International Standards for Clinical Trial Reg | | |
| a) Registries must have written standards for all procedures and | Yes | |
| processes employed by the registry. These are known as Standard | | No |
| Operating Procedures (SOPs). | \boxtimes | |
| b) Does the Registry use these SOPs to train all staff processing trial | Yes | No |
| registrations to ensure that common standards for ensuring data | | |
| completeness and meaningfulness are adhered to. | \boxtimes | |
| c) Are the internal registry-specific SOPs aligned with the WHO | Yes | No |
| International Clinical Trial Registry Platform International Standards | | |
| for Clinical Trial Registration? | \boxtimes | |
| | : I | |
| EMA: These take the form of commission guidance, on line help and use | | |
| opens. | | |





2.3.1. Does the Registry make sure that the person registering the trial exists Yes No and that they are the appropriate Responsible Registrant? \times EMA: This is a legal process via the clinical trial application to the NCAs. 2.3.2. Does the Registry make sure that the trial exists? Yes No \boxtimes If yes, please briefly describe what the registry does to make sure that the trial exists. This is a legal process as it is controlled by the Clinical Trial Application and authorisation legislation. EMA: The EU-CTR displays information based on official applications provided by the sponsors to the relevant NCAs and IECs when requesting authorization on starting a clinical trial in their territory. Status of the trial is updated by the sponsor and information provided the NCAs that update the EudraCT database. a) Does the Registry obtain written third-party confirmation that a trial Yes No exists? \boxtimes If yes, please specify the method of confirmation: See answer above, the "third party" is the National Competent Authority b) Does the Registry display in the trial record: Yes No a. if the registry has obtained written third party confirmation of \boxtimes the trial's existence, and b. the name of the third party from whom confirmation was received (eg the name of the ethics committee) It is stated which authority has authorized the trial. 2.4. The Registry will have a publicly accessible audit trail so that changes made to the WHO TRDS for an individual trial can be tracked. a) Does the Registry allow Responsible Registrants to update their Yes No registered trial records? \boxtimes EMA: Via legally required substantial amendment and end of trial notification to the NCAs. b) Does the Registry make available a publicly accessible audit trail of Yes No any changes to any TRDS items? X П EMA: We will display the versions as of EU CTR 1.1. Does the have quality control procedures in place to ensure any Yes No updated information continues to fulfil the standards for each TRDS \boxtimes d) Does the Registry use the most up-to-date information as the default Yes No display? \boxtimes Can the TRDS, as originally registered, be accessed at all times? Yes No \boxtimes EMA: All versions are recorded in EudraCT and visible to NCAs - not yet in public but will with Version 1.1.

2.3. The Registry will have processes in place to make sure that people and trials exist





Registries (ISCTR).

| a) | Does the Registry Administrator have a thorough working knowledge of the operational aspects of their registry. | Yes | No |
|----|---|-----|----|
| b) | Is the Registry Administrator committed to ensuring that all Registry staff are familiar with the standards described in the ISCTR? | Yes | No |
| c) | Are all Registry staff familiar with the contents of the ISCTR? | Yes | No |





3. Accessibility

| 3.1. The Registry will make the <u>WHO TRDS</u> for all registered trials accessi public at no charge (ICMJE requirement). | ble to 1 | he |
|--|----------|---|
| a) Does the Registry make the WHO TRDS items for all studies in their register (i.e. the registry database) accessible online at no charge to the end user? | Yes | No |
| 3.2. The Registry will make it possible for the <u>WHO TRDS</u> for all registered searched electronically (<i>ICMJE requirement</i>). | l trials | to be |
| a) Is it possible to search the Registry online using electronic searches of text words and phrases via a simple, single search box? | Yes | No |
| b) Does the online search allow users to search in at least the condition field and the intervention field? | Yes | No □ |
| c) When the results of a trial identified by a search are displayed, are all items in the WHO TRDS visible? | Yes | No □ |
| d) Does the online search have an advanced search option? Note: This is not a requirement and is being asked for information only. | Yes | No |
| 3.3. The Registry will allow Responsible Registrants to submit a trial for reany time of day on any day of the week (24 hours a day, seven days a wee | | tion at |
| a) Is it possible to submit a trial to the registry 24 hours a day, seven (7) days a week? | Yes | No |
| EMA: EudraCT is available 24/7 for completion of CTA information by sponsors and for its upload by NCAs. EU CTR checks EudraCT at least every 30 minutes for updates. Data are first submitted by the sponsors to the NCAs and then by the NCAs loaded in EudraCT during the normal working days, but the system per se is constantly available. | | Province of the second |
| b) If the Registry is planning downtime, does it publish advance notice of downtime at least one (1) week beforehand? | Yes | No 🗆 |
| 3.4. The Registry will allow their register database to be searched at any ti any day of the week (24 hours a day, seven days a week). | me of (| day on |
| a) Is it possible to search the register online 24 hours a day, 7 days a week? | Yes ⊠ | No |





3.5. It is desirable that Registries in the WHO Registry Network also make the <u>WHO TRDS</u> available in the language(s) of the country or countries served by the registry.

| Does the Registry accept and/or display trial information in languages others than English? | Yes | No |
|--|---|---------|
| If yes, please specify the languages used: | | |
| <i>EMA:</i> Information is displayed in the EU-CTR in the language in which the data are loaded in EudraCT database. EU sponsors and citizens have the right to use an EU Official language. It means that for single country clinical trials the information can appear on the official language of that country where the application is submitted and the trial conducted and it might not always have been in English. As of 10 th March 2011 and launch of EudraCT v8.0 all free text fields can be entered first in English and then in any additional languages and each language entry is in a separate added field, labelled with the name of the language, but part of the same registration file for each country. | T THE PROPERTY AND A STATE OF | |
| Only answer the remaining questions in this section if the answer to the above q | uestion | is yes. |
| b) Does the Registry have quality control procedures in place to ensure that all translations are accurate? | Yes | No ⊠ |
| EMA: This is controlled by NCAs at CTA review stage. | | |
| c) Are all TRDS items for all records also available in English? | Yes | No |
| EMA: In general yes and for future see 3.5 above. | | |
| d) Are all trial records translated into English by the Responsible Registrant checked by registry staff against the non-English submission before being accepted for registration? | Yes | No |
| EMA: By NCAs when applicable. | | |
| e) Are all trial records translated by Registry staff checked by at least one other staff member? Not applicable | Yes | No ⊠ |
| f) If there is a discrepancy in a translation, is the translation checked by a third person? Not applicable it would be queried by NCA with sponsor. | Yes | No ⊠ |
| g) Does the Registry make users of the Registry aware of who performed the translation (the Responsible Registrant or Registry staff) of a registered record. Not applicable | Yes | No ⊠ |
| h) If a trial is registered in more than one language then will the Registry submit the "Scientific Title", and a language identifier, to the ICTRP Search Portal for each language used? | Yes ⊠ | No |





4. Unambiguous Identification

| 4.1. The Registry will have in place processes to prevent the registrati trial more than once on their database. | on of a | a single | |
|--|---------|----------|--|
| a) Does the Registry make sure that a trial that has been submitted for registration is not already been included in their register by first searching and checking their own database? Not applicable, as a trial is only registered once per Member State due to clinical trial authorization process. | | No ⊠ | |
| b) Does the Registry have policies and procedures in place to deal with inadvertent duplicate registration of the same trial within their own register? It would not happen see a above. | Yes | No | |
| 4.2 The Registry will facilitate the retrospective linking (or bridging) on the WHO Search Portal of a single trial registered with more than one registry by entering secondary identifiers. This includes the <u>UTN</u> , and the unique identifiers allocated by other registries in the WHO Registry Network. | | | |
| a) Does the Registry require responsible Registrants to make an entry in the Secondary Identifiers field? | Yes | No | |
| EMA: Sponsors are reminded to do so but it is not possible to check at present. We will in future seek to index against other public information. In particular, we will add a default entry of information not available. | | | |
| b) If there are no known secondary identifiers, does the Registry require Responsible Registrants to enter 'Nil known' in the Secondary Identifiers field? EMA: We can if this is required, not at present. | Yes | No | |
| c) Does the Registry require Responsible Registrants to enter a UTN? Note: This is not a requirement and is being asked for information only. | Yes | No ⊠ | |
| 4.3. It is desirable that Primary Registries will search the ICTRP Sear attempt to determine if the trial has already been registered by an Registry in the WHO Registry Network or an ICMJE approved registry. | | | |
| a) Does the Registry attempt to determine whether a submitted trial has been registered in another Primary Registry or an ICMJE approved registry before registration? | Yes | No | |
| EMA: We may do indexing in future against other public registers. Note: This is not a requirement and is being asked for information only. | | | |





5. Technical Capacity

5.1. The Registry will submit the <u>WHO TRDS</u> for <u>all</u> records on their register, in English, to the WHO ICTRP Central Repository.

| to the WHO ICTRP Central Repository. | | |
|--|---|---|
| a) Will the Registry submit the WHO TRDS items for all records on their register, in English, to the WHO ICTRP Central Repository? Note: If a registry accepts study types other than interventional trials (i.e. observational studies) these must be provided as well. | Yes | No |
| EMA: All public items in EU CTR will go to WHO ICTRP. | | |
| b) Will the Registry submit records in the format requested by the WHO ICTRP (e.g. xml file) at least once per month? | Yes | No |
| c) Will the Registry, after the initial data transfer of all records, only submit new or updated records each time (rather than the entire data set every time). | Yes | No |
| EMA: Copy of the data will take place as required by the WHO ICTRP. | | |
| 5.2. The Registry will have access to a database that is used to store an submitted data. | d man | age the |
| a) Does the Registry use database software and hardware that guarantees reliable access to registered data and data safety at all times? | Yes ⊠ | No |
| 5.3. The Registry will have access to adequate information technology s | upport | • |
| a) Does the Registry have access to reliable information technology support? | Yes | No |
| b) Does the Registry have access to all of the following? a. reliable application, database, backup and mail servers b. good internet connectivity speed c. sound operating systems d. appropriate software for servers, desktops and laptops e. database and web development and maintenance personnel f. other skilled information technology personnel to support these systems, as required | Yes 🖂 | No |
| Please briefly describe the Registry's information technology infra separate document may be submitted separately if necessary) | | |
| EMA: The EudraCT database and the EU-CTR are hosted and managed by Medicines Agency, a decentralised Agency of the European Union, located in Agency is responsible for the scientific evaluation of medicines developed by p companies for use in the European Union. Role of the Agency has been first de Regulation (EEC) No 2309/93 replaced by Regulation (EC) 726/2004. The advanced and powerful infrastructure and a large / strong ICT department com term to deliver and maintain large amounts of ICT projects. Its technologies are latest industry standards (multi-tier web applications backed by robust database | Londo harmadescribed EMA I mitted based | on. The ceutical d in the has an in long on the |

on robust middleware servers) and its ICT processes are strictly controlled and properly managed. Back up procedures on-side and off-side and tools are in place for the full data set managed at the Agency. Finally, since both the Agency itself and the EU-CTR are established

by European Union legislation their long term availability is guaranteed.





5.4. The Registry will have adequate security and other provisions against data corruption and loss.

| a) | Does the Registry have documented procedures for ensuring adequate data security and other provisions to prevent data corruption and loss. | Yes | No |
|---------|--|----------|----|
| If yes, | Does the Registry issue alerts in advance of website downtime? please briefly describe how these alerts are circulated and who is sible for circulating them. | Yes ⊠ | No |
| 1 Capon | sible for circulating them. | | |





6. Administration and Governance

| 6.1. The Registry will have at least a national remit, and the support of within the country (or region) to act as the Primary Registry for that countries and not a group of states within a countries. | ıntry oı | rnment r region |
|---|---------------------------------|---------------------------|
| a) Does the Registry have at least a national remit? | Yes | No |
| FIRES It is board on Art 44 Directive 2004/20/FC Art 57/0\ (FD) 1 1 (FO) | | П |
| EMA: It is based on Art 11 Directive 2001/20/EC, Art 57(2) of Regulation (EC) No. 726/2004 and Art 41 of Regulation (EC) No.1901/2006. | | J |
| 1 10. 720/2004 and Art 41 of Negulation (EC) No. 190 1/2000. | | |
| b) Does the Registry have a letter of support, or other appropriate | Yes | No |
| documentation, from the Ministry of Health or other relevant national | | _ |
| or regional agencies? | | |
| Note: The letters of support must be submitted to the ICTRP Secretariat as part of the application | <u></u> | |
| c) From which country (or countries) does the Registry have the reminational (or regional) clinical trial registry? | to act | as the |
| EMA: As described by law from all the EU Member States – so all EU Member | States. | |
| d) Please specify the name of the national agencies that have given their Registry: | suppo | rt to the |
| EMA: All the 27 EU Member States and 3 additional EEA Member States. | | |
| 6.2. The Registry will publicly disclose ownership, governance structur profit status. | e and i | not-for- |
| a) Is the Registry managed by a not-for-profit agency? It is managed by | Yes | No |
| law by EMA which is a public EU body. | | |
| b) Does the Registry publicly disclose its ownership, governance | | |
| structures and not-for-profit status in a prominent place on the registry's website? EMA is a public body | Yes | No |
| c) What is the web address for the page where the ownership, | | |
| governance and not-for-profit information is displayed? | | ĺ |
| www.ema.europa.eu | | |
| | | |
| d) Will the Registry inform the ICTRP immediately if their ownership, | Yes | No |
| governance structures or not-for-profit status change in any way. | Yes | No |
| d) Will the Registry inform the ICTRP immediately if their ownership, governance structures or not-for-profit status change in any way. It is not possible for this to change as it is set out in law. | | No |
| governance structures or not-for-profit status change in any way. It is not possible for this to change as it is set out in law. 6.3. The Registry agrees that, should it cease to function, at least the (original and updated) for all trial records will be transferred to a Primathe WHO Registry Network. | ⊠ <u>WHO</u> ry Regi | TRDS |
| governance structures or not-for-profit status change in any way. It is not possible for this to change as it is set out in law. 6.3. The Registry agrees that, should it cease to function, at least the (original and updated) for all trial records will be transferred to a Prima the WHO Registry Network. a) Will the Registry transfer at least the WHO TRDS (original and | ⊠ who | TRDS |
| governance structures or not-for-profit status change in any way. It is not possible for this to change as it is set out in law. 6.3. The Registry agrees that, should it cease to function, at least the (original and updated) for all trial records will be transferred to a Prima: the WHO Registry Network. a) Will the Registry transfer at least the WHO TRDS (original and updated) for all trial records to another Primary Registry in the WHO | ⊠ <u>WHO</u> ry Regi | TRDS |
| governance structures or not-for-profit status change in any way. It is not possible for this to change as it is set out in law. 6.3. The Registry agrees that, should it cease to function, at least the (original and updated) for all trial records will be transferred to a Primathe WHO Registry Network. a) Will the Registry transfer at least the WHO TRDS (original and updated) for all trial records to another Primary Registry in the WHO Registry Network if it ceases to function? | | TRDS istry in |
| governance structures or not-for-profit status change in any way. It is not possible for this to change as it is set out in law. 6.3. The Registry agrees that, should it cease to function, at least the (original and updated) for all trial records will be transferred to a Primathe WHO Registry Network. a) Will the Registry transfer at least the WHO TRDS (original and updated) for all trial records to another Primary Registry in the WHO Registry Network if it ceases to function? It is not possible for this to happen as it is set out in law. 6.4. The Registry will have a strategy in place ensure the medium sustainability of the registry | ⊠ WHO ry Regi Yes ⊠ | TRDS istry in |
| governance structures or not-for-profit status change in any way. It is not possible for this to change as it is set out in law. 6.3. The Registry agrees that, should it cease to function, at least the (original and updated) for all trial records will be transferred to a Primathe WHO Registry Network. a) Will the Registry transfer at least the WHO TRDS (original and updated) for all trial records to another Primary Registry in the WHO Registry Network if it ceases to function? It is not possible for this to happen as it is set out in law. | ⊠ WHO ry Regi Yes ⊠ | TRDS istry in |
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| governance structures or not-for-profit status change in any way. It is not possible for this to change as it is set out in law. 6.3. The Registry agrees that, should it cease to function, at least the (original and updated) for all trial records will be transferred to a Primathe WHO Registry Network. a) Will the Registry transfer at least the WHO TRDS (original and updated) for all trial records to another Primary Registry in the WHO Registry Network if it ceases to function? It is not possible for this to happen as it is set out in law. 6.4. The Registry will have a strategy in place ensure the medium sustainability of the registry | | TRDS istry in No g term |





7. The Trial Registration Data Set (TRDS)

7.1. The Registry will collect and publicly display all items in the WHO Trial Registration Data Set (TRDS)

| | Item/Label | Does the registry collect and publicly display this data item? | |
|----|--|--|---------|
| 1 | Primary Registry and Trial Identifying Number | Yes | No |
| 2 | Date of Registration in Primary Registry | Yes | No |
| 3 | Secondary Identifying Numbers | Yes ⊠ | No □ |
| 4 | Source(s) of Monetary or Material Support EMA note: field added as of March 2011 | Yes | No |
| 5 | Primary Sponsor | Yes | No |
| 6 | Secondary Sponsor(s) | Yes | No |
| 7 | Contact for public queries EMA note: field added as of March 2011 | Yes | No |
| 8 | Contact for scientific queries EMA note: Same contact as in 7 is used in accordance with legal guidance of EU. | Yes | No |
| 9 | Public title | Yes ⊠ | No |
| 10 | Scientific title | Yes ⊠ | □ & |
| 11 | Countries of Recruitment EMA note: a separate record is displayed for each country in the EU involved in the trial | Yes | No |
| 12 | Health condition(s) or problem(s) studied | Yes ⊠ | No |
| 13 | Interventions | Yes | No |
| 14 | Key Inclusion and Exclusion Criteria | Yes ⊠ | No □ |
| 15 | Study type | Yes ⊠ | No □ |
| 16 | Date of first enrolment EMA note: Currently date of authorisation, i.e. from which point enrolment is permitted, enrolment per se will be added in v8.2 planned in 2012 | Yes | No ⊠ |
| 17 | Target sample size | Yes | No |
| 18 | Recruitment status EMA Note: Information on trial status (ongoing, completed, prematurely ended) is available at the moment. Trials are "ongoing" from the date of authorisation until the end of the trial (i.e. last patient last visit). Actual recruitment status will be available from EudraCT v8.2 planned in 2012 | Yes | No ⊠ |



International Clinical Trials
Registry Platform

| | Item/Label | Does the registry collect and publicly display this data item? |
|----|--------------------------|--|
| 19 | Primary Outcome(s) | Yes No □ |
| 20 | Key Secondary Outcome(s) | Yes No ⊠ □ |

7.2. The Registry may choose to collect and publicly display other data items. It is recommended that registries consider the following optional, additional data items:

| | Item/Label | | Does the registry collect and publicly display this data item? | | |
|----|---|-----|--|--|--|
| 21 | Lay Summary / Synopsis EMA note: Title for lay people is available at the moment, summary is not required, scientific information is favoured. | Yes | No | | |
| 22 | Approvals | Yes | No | | |
| 23 | Results links EMA note: Not available at the moment but it will take place in future – in v9.0 of EudraCT and future versions of EU-CTR. | Yes | No ⊠ | | |





8. Partner Registries

| 8.1. Primary Registries in the WHO Registry Network will have the capacity to partner with other Registries. | | | |
|---|----------|---------|--|
| a) Is the Registry willing and able to form partnerships with other Registries that do not themselves fulfil the criteria for a Primary Registry in the WHO Registry Network? | Yes | No ⊠ | |
| EMA: It is not very clear what this really implies. As EudraCT has a legal framework and mandate it cannot simply start to accept data from other sources. | | | |
| b) Does the Registry currently have any Partner Registries? | Yes | No | |
| If yes, please provide the name(s) of these partners: Not applicable, there are not partnerships in place at the moment. | | | |
| c) If the registry has partners, are they listed on the proposed Primary Registry's web site? | Yes | No | |
| Not applicable If yes, please provide the address of this web page: | | | |
| Note: A completed Registry Profile form is required for all Partner Registries. This profile will be published on the ICTRP's web site. | | | |
| 8.2. Primary Registries in the WHO Registry Network will ensure that pote Registries meet WHO minimum standards requirements. | ntial Pa | artner | |
| a) Does the proposed Primary Registry agree that, before agreeing to accept a Partner Registry and their trial registration records, they will make sure that the Partner Registry meets all the WHO minimum standards listed in the International Standards for Clinical Trial Registries. | Yes | No | |
| b) Has a Registry Profile form been completed and submitted for all of the Registry's Partner Registries? Not applicable | Yes | No | |
| 8.3. Primary Registries will have procedures in place to enable exchange Partner Registries. Not applicable see above. | of data | with | |
| a) Is the Registry able to accept data (that is, as electronic data files) from Partner Registries or other appropriate data providers? Not applicable | Yes | No | |
| b) Does the Registry agree to establish a Memorandum Of Understanding (MOU) or other such agreement with each Partner Registry or other data providers, as per the requirement described in the International Standards for Clinical Trial Registries? | Yes | No | |
| c) Does the Primary Registry agree the area of coverage/responsibility of their Partner Registries or other data providers (such as geographical location, health condition, intervention type, etc) and incorporate this into their SOPs and instructions to Registrants to avoid any confusion or unintentional duplicate registration? | Yes | No | |
| d) Does the Primary Registry record the identification number and date of registration in the Partner Registry within the trial record on the Primary Registry? | Yes | No | |



| e) | Does the Primary Registry identify records that have been sourced from Partner Registries or other data providers so users are aware of the data source? | Yes | No |
|----|--|-----|----|
| f) | Before announcing Partner Registries, Primary Registries must have successfully imported data into the Primary Registry. | Yes | No |





Contact Information

Administrator

The Administrator is the person employed to manage the Registry and will be the primary point of contact between the Registry and the ICTRP Secretariat.

| Title (Dr/Prof/Mr/Mrs/Ms/Miss) | Dr |
|--------------------------------|------------------------------|
| Given Name | Fergus |
| Family Name | Sweeney |
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| Fax | +44 207 418 8595 |
| Email | Fergus.sweeney@ema.europa.eu |

Secondary Contact

The Secondary Contact must not be the same as the Administrator. Secondary contact details are requested for circumstances when the Administrator is unavailable.

| Title (Dr/Prof/Mr/Mrs/Ms/Miss) | Dr |
|--------------------------------|-----------------------------|
| Given Name | Ana |
| Family Name | Rodriguez Sanchez Beato |
| Telephone number | +44 207 523 7160 |
| Fax | +44 207 418 8595 |
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Information Technology Officer

The Information Technology Officer should be the person who will be responsible for submitting the data for inclusion on the ICTRP Central Repository. They should have a good understanding of all of the IT issues relevant to the registry.

| Title (Dr/Prof/Mr/Mrs/Ms/Miss) | Mr |
|--------------------------------|------------------------------|
| Given Name | Olivier |
| Family Name | Simoen |
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| Fax | +44 20 74188670 |
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