



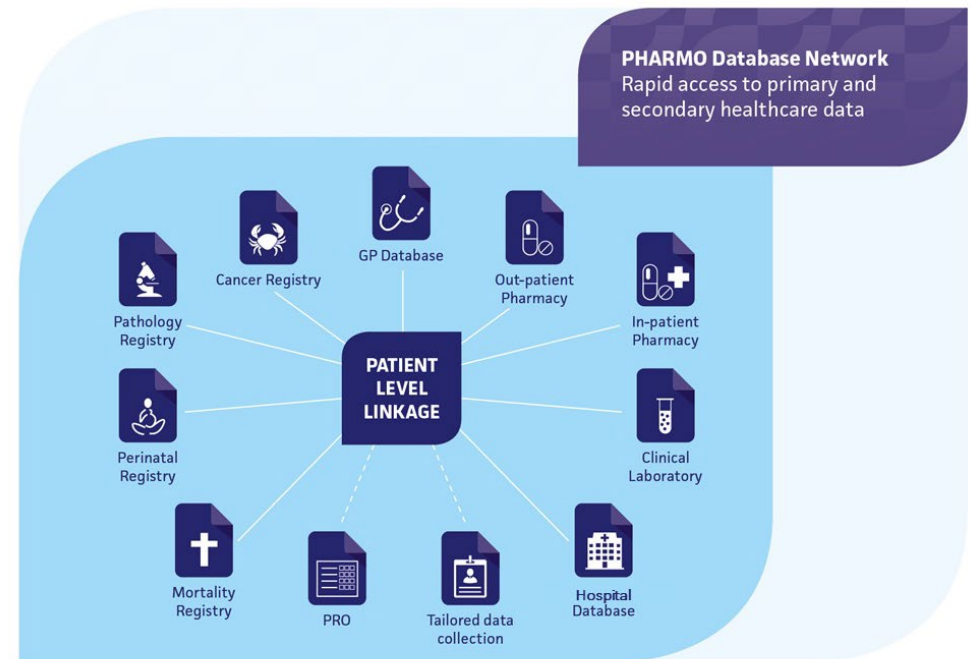
Real World research on medicinal products: key contribution of the ENCePP

Jetty A. Overbeek

**How can ENCePP contribute to
research?
Experience from an EU academic**

Background

- The PHARMO Institute
- PASS and PAES
- PHARMO & ENCePP
 - Research centre since 2011
 - First collaboration in PASS/PAES 2012
 - First coordination of PASS/PAES in 2013



Transparency: EU PAS Register

The screenshot shows the ENCePP website header with the logo and navigation menu. The main content area is titled 'The European Union electronic Register of Post-Authorisation Studies (EU PAS Register)'. It provides instructions on how to register a new study, resume a draft application, or update an existing study. There are buttons for 'Add Study', 'Edit Study', and 'Search'. A sidebar on the left contains links to various resources like News, About Us, and the Code of Conduct. The footer includes copyright information, contact details, and logos for the European Medicines Agency and the European Union.

ENCePP European Network of Centres for Pharmacoepidemiology and Pharmacovigilance

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Home > EU PAS Register

The European Union electronic Register of Post-Authorisation Studies (EU PAS Register)

On this page you can register (or resume a draft application for) a new study, update existing study records or search the EU PAS Register.

To **register a new study** please click on 'Add Study' below:
(If this is a study related to the coronavirus pandemic, please include the text **COVID-19** in the study title)

Add Study

To **resume a draft application** saved previously (but not submitted yet) or to amend and re-submit a previously rejected application, please follow the link:
[Resume Draft/Rejected application](#).

To **update** an existing study record please click on 'Edit Study' below:

Edit Study

To search for registered studies, please click on 'Search' below:

Search

For assistance please contact us at: EU_PAS_Register@ema.europa.eu

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For the UK, as from 1.1.2021, EU Law applies only to the territory of Northern Ireland (NI) to the extent foreseen in the Protocol on Ireland/NI

EUROPEAN MEDICINES AGENCY

Version: 4.4.2.0

Transparency: ENCePP Resources Database



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Home > Resources Database

ENCEPP Resources Database

The Resource Database is an electronic index of available EU research organisations, networks and data sources, including patient registries, in the fields of pharmacoepidemiology and pharmacovigilance, and is a key component of the ENCePP web portal. All information contained in this database is provided and maintained by the listed organisations, data providers or registry holders.

The Resource Database includes information on expertise and research experience across Europe and serves as a hub for both researchers and study sponsors seeking to identify organisations and data sets for conducting specific pharmacoepidemiology and pharmacovigilance studies in Europe.

The database is fully searchable and allows the identification of research centres, data sources and patient registries by country, type and other relevant criteria. Guidance on how to search for patient registries is provided in the document "[ENCEPP Resource Database – Inventory of Patient Registries](#)". Please click on the following button to search the Database:

Search Database

If you and/or your organisation are interested and/or actively involved in performing research in the field of pharmacoepidemiology and pharmacovigilance and would like to join the ENCePP network please click "Add Centre/Network" or contact us at encepp_secretariat@ema.europa.eu

Existing patient registries are invited to enter their details in the Inventory of Data Sources. Guidance on how to upload patient registries is provided in the document "[ENCEPP Resource Database – Inventory of Patient Registries](#)".

Click "Add Data Source" if you are managing a data set or patient registry that can be used in pharmacoepidemiology and pharmacovigilance research.

Click "Edit Resource" if you already have an entry in the Resources Database and would like to update it.

If you wish to resume a draft application saved previously (but not submitted yet) or update a returned application, please follow the link:
[Resume Draft/Returned application](#).

Add Centre/Network

(By adding a centre or network/collaboration, you automatically join the ENCePP network)

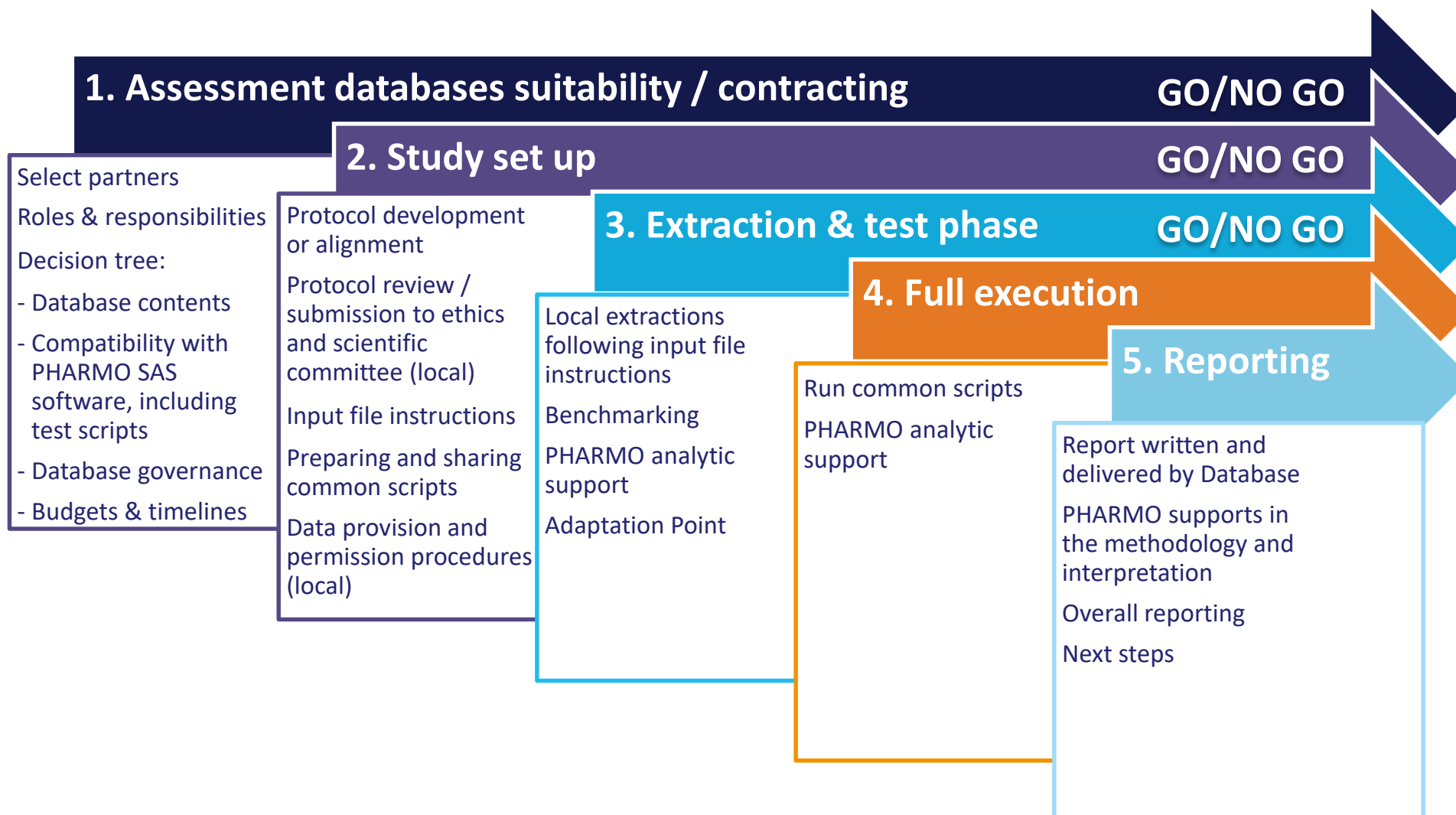
Expanding EU Network



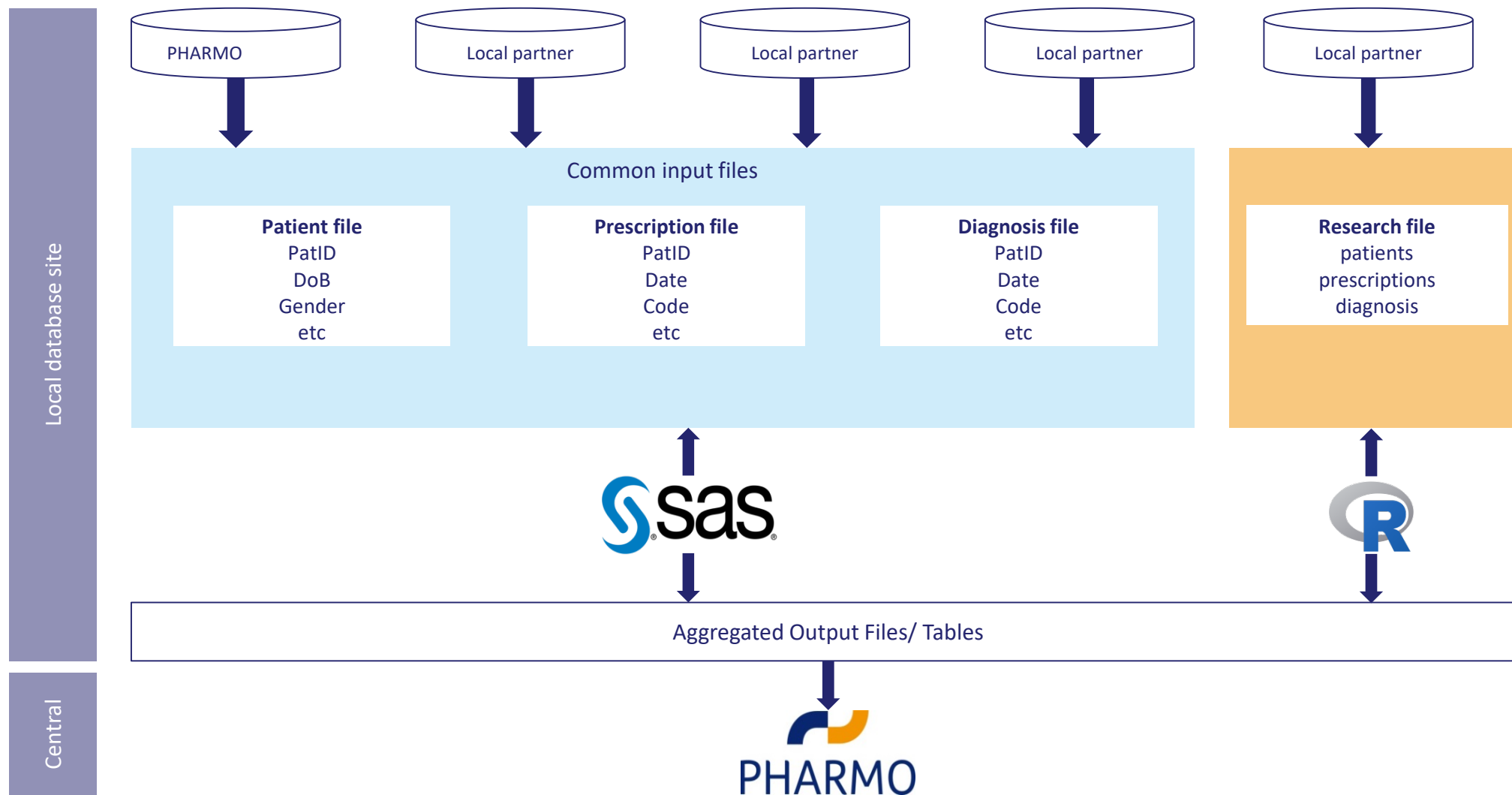
Expanding EU Network



Approach for multi-database studies

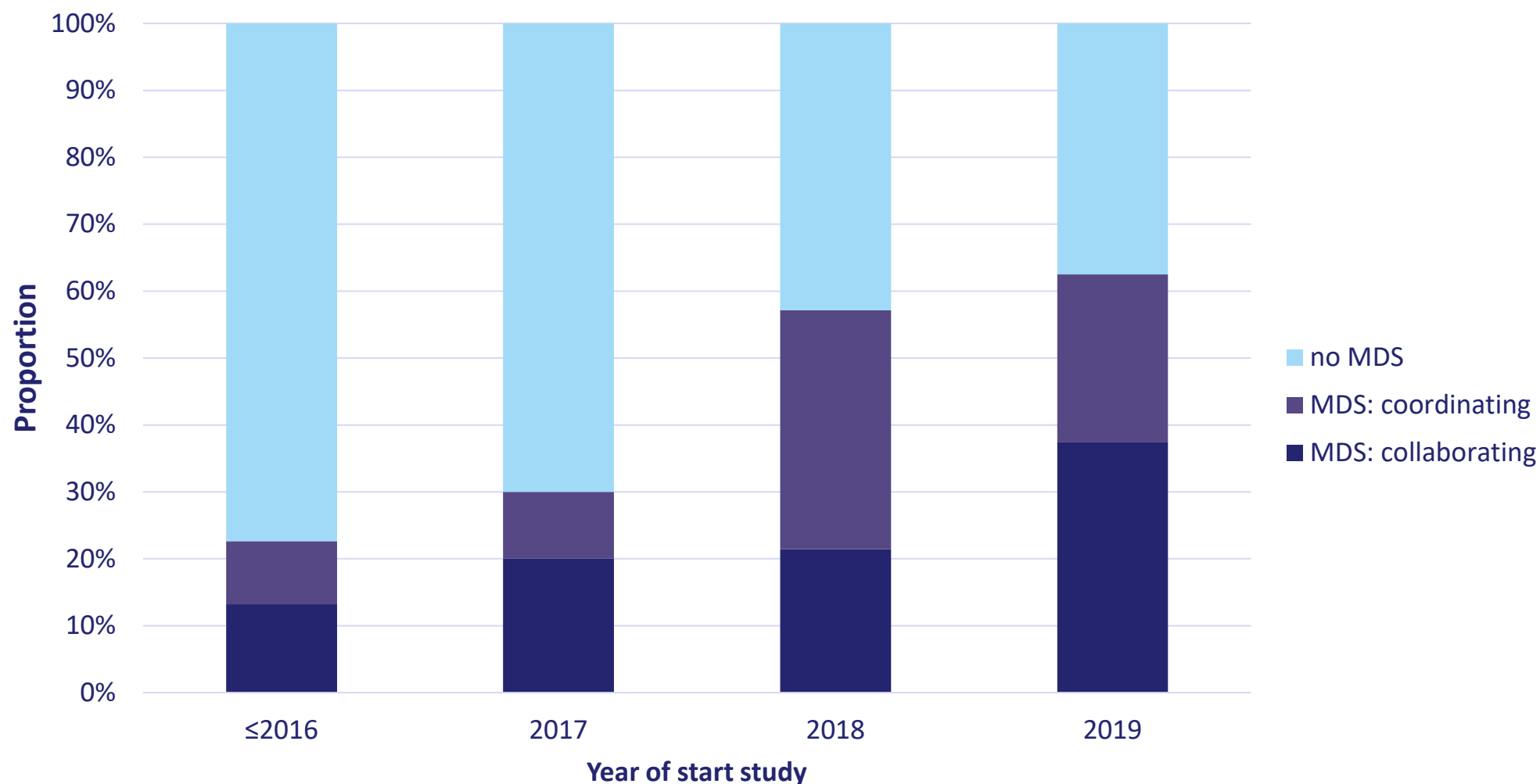


Approach for multi-database studies





Track record multi-database studies (MDS)



Value of ENCePP tools

- EU PAS Register
- Resources Database
- Standards and Guidance
 - Checklist for Protocols
 - Methodological Guide
- Code of Conduct
- ENCePP Study Seal

Recommendations and wishes

- More interaction/involvement
- For the future
 - Accreditation for research centres?
 - Added value of ENCePP seal?

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