

Session 4: Looking towards the future: Brainstorming on funding mechanisms for PAS

Example: Collaborative vaccine studies

ADVANCE



Accelerated Development of VAccine beNefit-risk Collaboration in Europe

- Public-private partnership created in the framework of the Innovative Medicines Initiative (EC+EFPIA)
- Coordinated by Erasmus University + GSK
- Public health agencies, academic institutions, regulatory authorities, vaccine manufacturers

Objective:

To create an infrastructure and a sustainable framework for vaccine benefit-risk monitoring in Europe



ADVANCE



WP1: Best practice Guidance

- Code of conduct: principles for collaboration between study participants
- Quality management: "minimum" quality management principles to be defined
- Governance models: roles & responsibilities, mechanisms of interactions
- Communication recommendations: how to communicate about vaccines safety and results of vaccine studies

WP2: Synergies with other projects

WP3: Data sources

WP4: Methods

WP5: Proof-of-concept studies

WP1/WG3 Governance models



Objective:

- to identify a few "typical" governance models for vaccine studies involving several stakeholders
- to provide recommendations on governance aspects
- to provide recommendations on which model(s) could be applied in different situations

Based on experience of studies that worked well or did not work well Identification of typical scenarios (still on-going)

Scenarios (1)



A public health institute intents to conduct a study to evaluate the effectiveness of a vaccine within a vaccination program.

Hypothesis:

Operational support needed; PHI data owner; vaccine effectiveness to be measured is not brand-specific.

A public health institute intents to raise awareness about the benefits of its vaccination program. The vaccine manufacturers who provide the vaccines are willing to support this initiative.

Hypothesis:

Involvement of several MAHs; not brand-specific.



Scenarios (2)



A vaccine manufacturer intents to conduct a study to assess the burden of disease to a candidate vaccine and use a large public health database it has no direct access to.

Hypothesis: Academia/CRO can have access to public health database; brand-specific.

A vaccine manufacturer intents to conduct a study to generate more evidence about the effectiveness of its vaccine in routine use.

Hypothesis: MAH can generate its VE own data or can use data from surveillance system; brand-specific.



Scenarios (3)



Due to concerns regarding the safety/effectiveness of a vaccine, a regulatory authority requests the vaccine manufacturer to investigate the safety/effectiveness/benefit-risk profile of its vaccine.

- Legal responsibility for the vaccine manufacturers
- Study protocol and report to be endorsed by regulators
- Vaccine manufacturers required to revise protocol/report
- Brand-specific investigation
- Secondary use of data vs. primary data collection
- For some public health institutions, legal/public perception issues to involvement of vaccine manufacturers

Which governance model(s) could allow participation of vaccine manufacturers?

ADVANCE Governance models



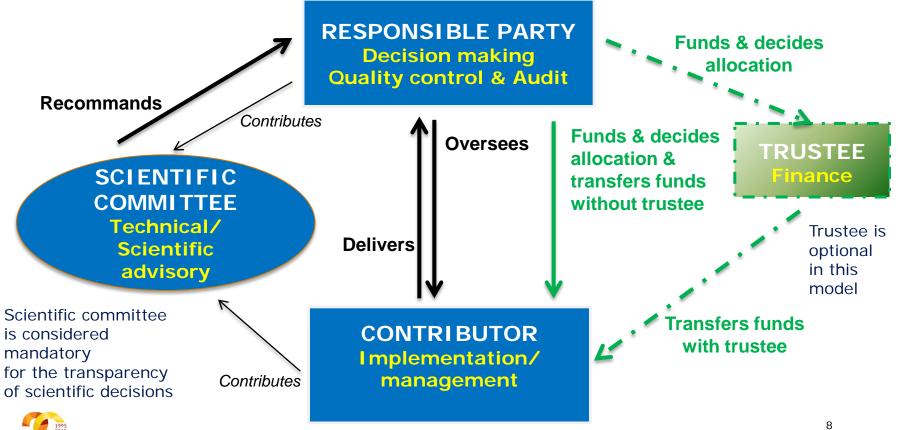
Five basic functions in all studies

- Decision-making
- Implementation/management
- Technical/Scientific advisory function
- Quality control & audit
- Finance

R&R for each function should be clearly identified at the start of the study.

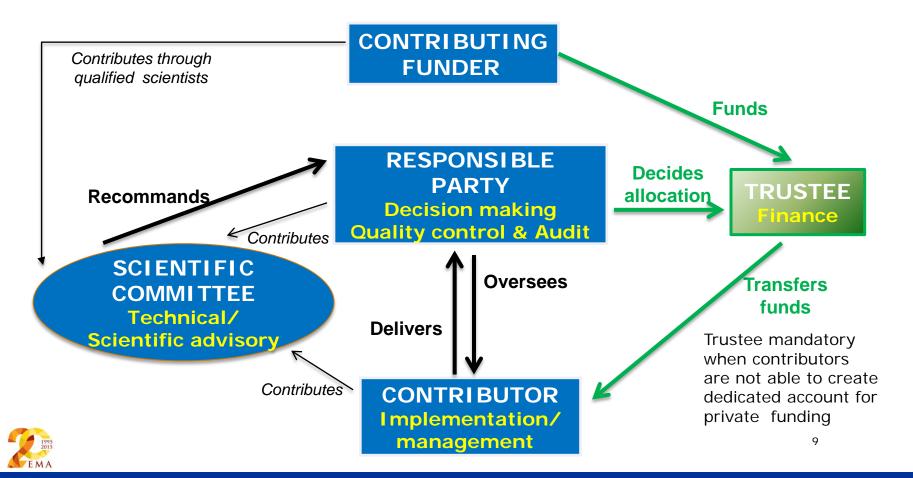






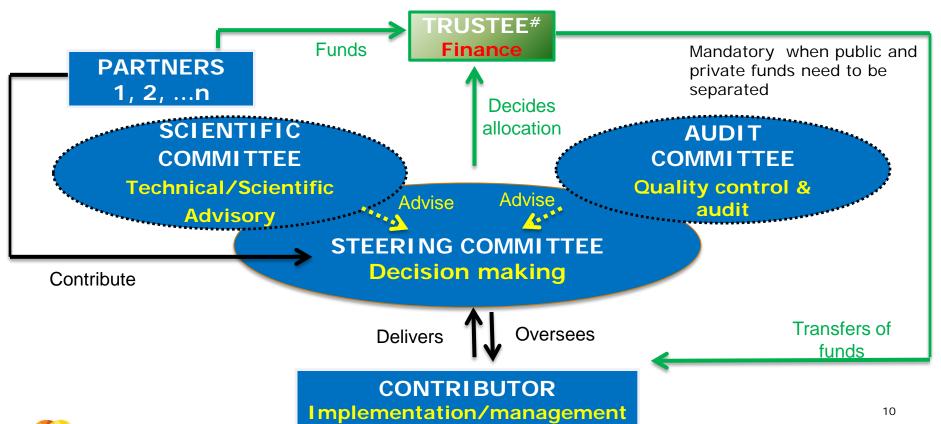
Model 2: Collaboration with contributing funder





Model 3. Partnership with shared funding/tasks







Due to concerns regarding the safety/effectiveness of a vaccine, a regulatory authority requests the vaccine manufacturer to investigate the safety/effectiveness/benefit-risk profile of its vaccine.

Issue: how to accommodate different constraints

- Vaccine manufacturers have legal obligations
- Regulators want to ensure best possible design for research question
- Some public health institutions may not interact with vaccine manufacturers
- The public is concerned by possibility of conflicts of interest





Model 1 (collaboration self-funded by the responsible party): Study conducted by Academia/CRO – VM is the decision maker

Model 2 (collaboration with contributing funder): VM provides funds and contributes to the project – PHI is the decision maker

Model 3 (partnership): Study conducted jointly by PHI and VM through a CRO/Academia

Model	1	2	3
Responsible party	VM	PHI	PHI & VM
Contributing funder	VM	VM	VM
Restricted /partial data access	Academia/CRO VM	PHI	PHI & VM
Results ownership / co authorship	VM	PHI co-ownership/authorship possible	PHI & VM
Comments	Most challenging situation: model 2 preferred by PHI for public trust in safety results and model 3 preferred by VM due to accountability in regulatory context		





Some suggestions from public health authorities present in ADVANCE:

- Regulator involved in steering committee to provide assurance to the public about lack of conflicts of interest
- Regulator liaises directly with public health authority / academic institution
- Choice of contributors for data collection and analysis independent from vaccine manufacturer (/regulator)
- Only centres accredited at national level allowed to participate in regulatory studies,
- etc...





Thank you for your attention

European Medicines Agency

30 Churchill Place • Canary Wharf • London E14 5EU • United Kingdom Telephone +44 (0)20 3660 6000 Facsimile +44 (0)20 3660 5555 Send a question via our website www.ema.europa.eu/contact



