

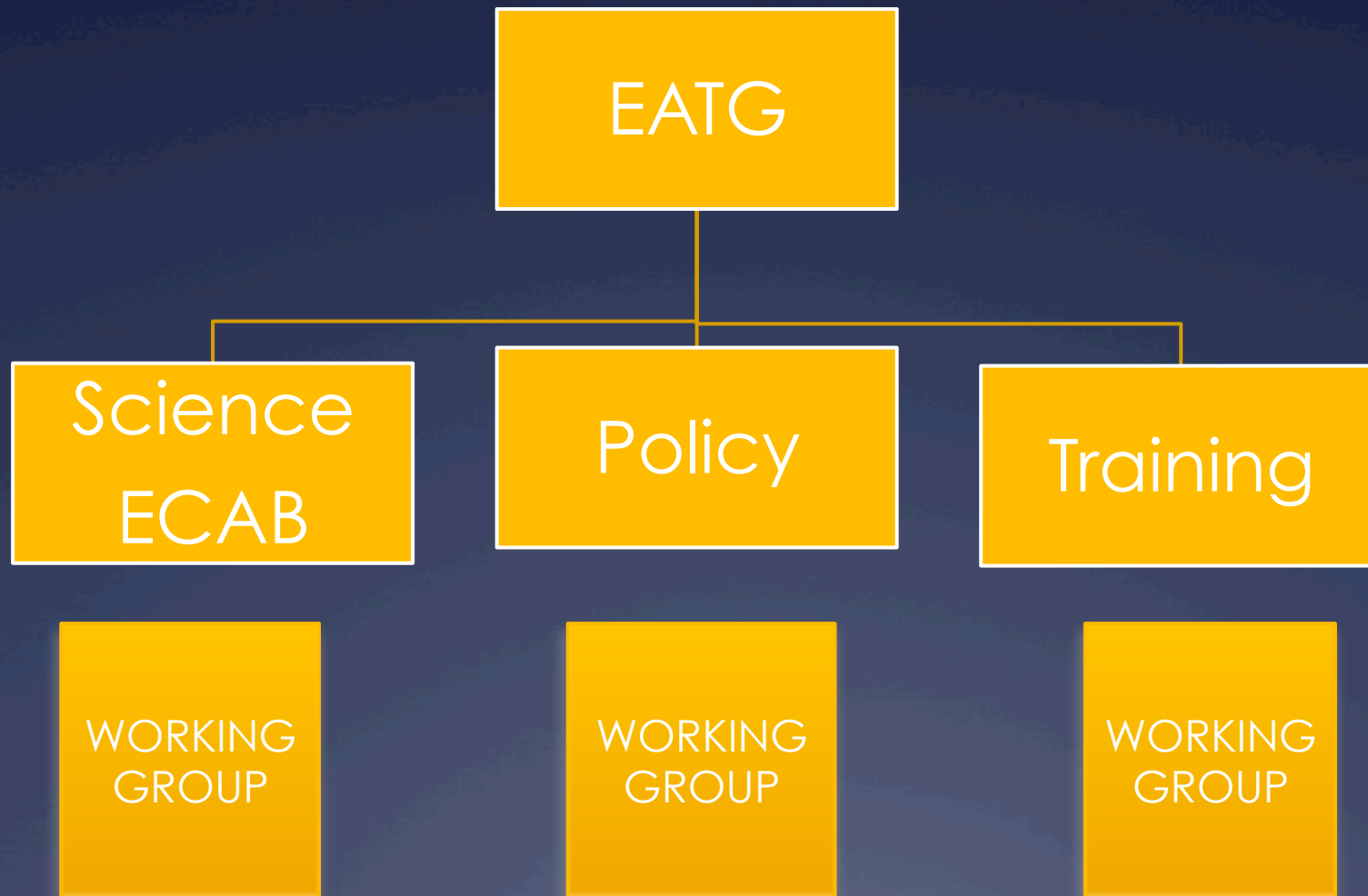
Patient involvement in clinical trials

European AIDS Treatment Group

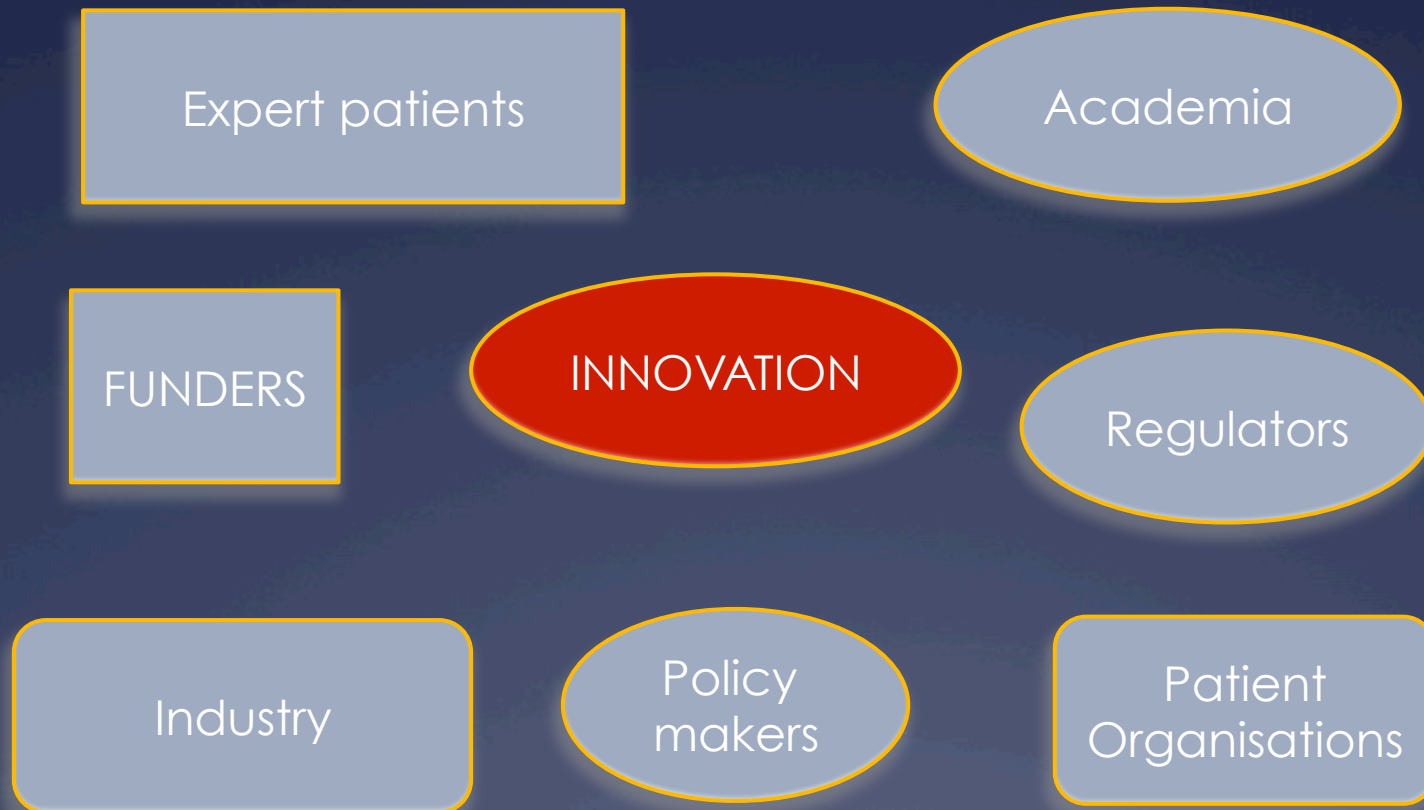
Damian Kelly



STRUCTURE



Who's the partners?



Trial Design

- Protocol review
- Informed consent review (appropriate language)
- Site selection (countries hospitals etc.)
- Demographics (women into trials, colour, co-infections)
- Inclusion / exclusion criteria
- Peer based dissemination – results, options of upcoming trials,
- Proposing sub studies – clinical relevance

ECAB

- European, National and International Guidelines

EACS, WHO, BHIVA

- Regulators (EMA) Patient information

- Conference Organisation – content and agenda setting!

- Data monitoring committee (DSMB)

- Ethics committee (national and regional)

Take Home Message!

When PATIENTS are an integral part of clinical design, its beneficial to the whole process!

Thank you

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