



Topic 3 – Timely completion of paediatric investigation plans (PIPs) Clinical Trial Application Perspective

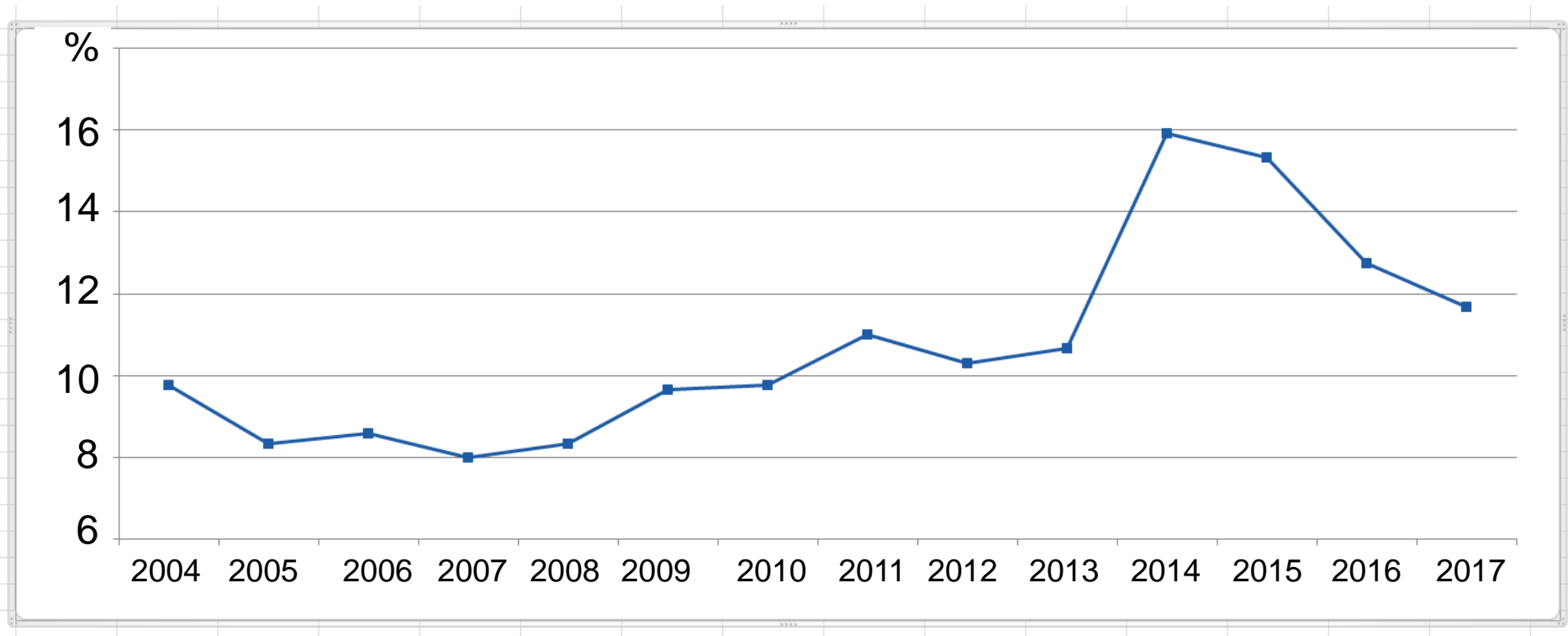
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Co-Chair Clinical Trial Facilitation Group (CTFG, working group under Heads of Medicines Agency, HMA)

EMA/EC multi-stakeholder workshop to further improve the implementation of the Paediatric Regulation, London March 20 2018

Clinical trial percentage involving minor subjects



Clinical Trial applications reported by sponsors to involve minors in at least one MS as a percentage of the total number of trials (not restricted to trials only involving paediatric population, *year reported when data generated in EudraCT – risk for mistake if data not uploaded year of trial application submission*).

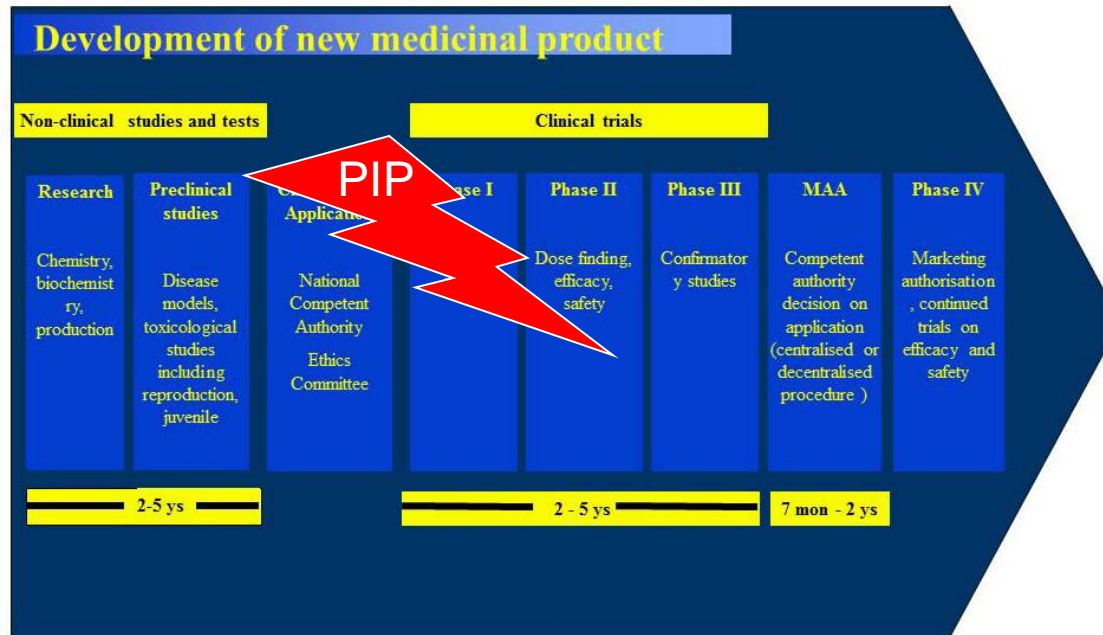
Datawarehouse search 2018-03-19 in EudraCT database

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When is timely application of PIP during development program related to individual clinical trial applications?



When, if complex clinical trial design not structured in traditional separate phases?

How do clinical trial applications (CTAs) interplay with an application for a paediatric investigation plan (PIP)?

	Scope	Decision/opinion /responsibility	Applicant	Possible contradiction PIP-CTA?
PIP				
CTA				

CTFG open for actions to increase contacts between assessors from different structures - PDCO, clinical trial units of NCAs and Ethics Committees

- Future **single decision National Competent Authority and Ethics Committee** on clinical trial applications per Member State when **Clinical Trial Regulation (CTR, EU No 536/2014)** replaces national laws on interventional clinical trials on safety and efficacy of medicinal products
CTR will apply when Portal and Database built by EMA has reached full functionality
- CTFG (Clinical Trial Facilitation Group) open for **exchange on issues relating to pediatric trials** between PDCO delegates and assessors from clinical trial units of National Competent Authorities and colleagues from Ethics Committees
- CTFG plans **future EU training workshop of paediatric clinical assessors from national competent authorities trial units and ethics committees** (date and program to be decided) promoting exchange of ideas on scientific and ethical aspects of paediatric clinical trials and harmonisation within EU

Thanks for your attention – questions welcome!



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