

Manuscript Number:

Title: ADVANCE system testing: benefit-risk analysis of a marketed vaccine using multi-criteria decision analysis and cohort modelling

Article Type: SI: ADVANCE

Keywords: benefit-risk assessment; pertussis vaccines; methodological study; electronic health record databases; Europe

Corresponding Author: Dr. Kaatje Bollaerts,

Corresponding Author's Institution: P95

First Author: Kaatje Bollaerts

Order of Authors: Kaatje Bollaerts; Edouard Ldent; Tom de Smedt; Daniel Weibel; Hanne-Dorthe Emborg; Giorgia Danieli; Talita Duarte-Salles; Consuelo Huerta; Elisa Martín-Merino; Gino Picelli; Lara Tramontan; Miriam Sturkenboom; Vncent Buchau

Abstract: Background

The Accelerated Development of VAccine beNefit-risk Collaboration in Europe (ADVANCE) is a public-private collaboration aiming to develop and test a system for rapid benefit-risk (B/R) monitoring of vaccines using electronic health record (eHR) databases in Europe. Proof-of-concept studies were designed to assess the proposed processes and system for generating the required evidence to perform B/R assessment and near-real time monitoring of vaccines. Most B/R assessment methodologies have been developed for medicinal products. We aimed to test B/R methodologies for vaccines, using the comparison of the B/R profiles of whole-cell (wP) and acellular pertussis (aP) vaccine formulations in children as an example.

Methods

We used multi-criteria decision analysis (MCDA) to structure the B/R assessment combined with cohort modelling to build the B/R effects table. In the cohort model, we simulated the number of events in two hypothetical cohorts of 106 children followed from first pertussis dose till pre-school-entry booster (or six years of age, whichever occurred first), with one cohort receiving wP, and the other aP. The benefits were reductions in pertussis incidence and complications. The risks were increased incidences of febrile convulsions, fever, hypotonic-hyporesponsive episodes, injection-site reactions and persistent crying. The model parameters were informed by estimates (coverage, background incidences, relative risks) from eHR databases from Denmark (SSI), Spain (BIFAP and SIDIAP), Italy (Pedianet) and the UK (RCGP-RSC and THIN). Preferences were elicited from clinical and epidemiological experts.

Results

Using cohort modelling to build the B/R effects table facilitated the comparison of different effect measures (e.g. immediate relative risks vs long-term vaccine effectiveness). Estimates from eHR databases could be used to inform the cohort model, with the cohort modelling results being easily combined with preference weights to obtain B/R scores.

Conclusion

Existing B/R methodology, cohort modelling and estimates from eHR databases can be successfully used for B/R assessment of vaccines.

Dr Gregory A Poland
Editor-in-Chief, Vaccine

22 November 2018

Dear Dr Poland

We are pleased to submit our paper 'ADVANCE system testing: benefit-risk analysis of a marketed vaccine using multi-criteria decision analysis and cohort modelling' to your Journal, Vaccine for the ADVANCE supplement. This paper describes the results from the benefit/risk assessment using the comparison of the B/R profiles of whole-cell (wP) and acellular pertussis (aP) vaccine formulations in children as an example. It is the 9th of the series of 10 papers that will be included in the supplement.

On behalf of all co-authors

Dr Kaatje Bollaerts

I, the undersigned, dr. Kaatje Bollaerts declare that all authors have seen and approved the final version of the manuscript being submitted. We warrant that the article is our original work that has not been previously published and is not under consideration for publication elsewhere

A handwritten signature in blue ink, appearing to read 'Kaatje Bollaerts', with a long horizontal stroke extending to the right.

Kaatje Bollaerts

***Suggested Reviewers**

Name	Institute	email
Shahrul Mt-Isa	Merck	shahrul.mt-isa@merck.com
Lydie Marcelon	Sanofi	lydie.marcelon@sanofi.com
Nicolas Praet	GSK	nicolas.x.praet@gsk.com
Andrew Thomson	European Medicines Agency	andrew.thomson@ema.europa.eu
Rob Hemmings	CHMP Biostatistics working party	rob.hemmings@mhra.gov.uk
Tjeerd van Staa	University of Manchester	tjeerd.vanstaa@manchester.ac.uk
Ed Waddingham	Imperial College, London	e.waddingham@imperial.ac.uk

1 **Abstract**

2 **Background**

3 The Accelerated Development of Vaccine benefit-risk Collaboration in Europe
4 (ADVANCE) is a public-private collaboration aiming to develop and test a system for rapid
5 benefit-risk (B/R) monitoring of vaccines using electronic health record (eHR) databases in
6 Europe. Proof-of-concept studies were designed to assess the proposed processes and system
7 for generating the required evidence to perform B/R assessment and near-real time monitoring
8 of vaccines. Most B/R assessment methodologies have been developed for medicinal
9 products. We aimed to test B/R methodologies for vaccines, using the comparison of the B/R
10 profiles of whole-cell (wP) and acellular pertussis (aP) vaccine formulations in children as an
11 example.

12 **Methods**

13 We used multi-criteria decision analysis (MCDA) to structure the B/R assessment combined
14 with cohort modelling to build the B/R effects table. In the cohort model, we simulated the
15 number of events in two hypothetical cohorts of 10^6 children followed from first pertussis
16 dose till pre-school-entry booster (or six years of age, whichever occurred first), with one
17 cohort receiving wP, and the other aP. The benefits were reductions in pertussis incidence and
18 complications. The risks were increased incidences of febrile convulsions, fever, hypotonic-
19 hyporesponsive episodes, injection-site reactions and persistent crying. The model parameters
20 were informed by estimates (coverage, background incidences, relative risks) from eHR
21 databases from Denmark (SSI), Spain (BIFAP and SIDIAP), Italy (Pedianet) and the UK
22 (RCGP-RSC and THIN). Preferences were elicited from clinical and epidemiological experts.

23 **Results**

24 Using cohort modelling to build the B/R effects table facilitated the comparison of different
25 effect measures (e.g. immediate relative risks vs long-term vaccine effectiveness). Estimates

26 from eHR databases could be used to inform the cohort model, with the cohort modelling
27 results being easily combined with preference weights to obtain B/R scores.

28 **Conclusion**

29 Existing B/R methodology, cohort modelling and estimates from eHR databases can be
30 successfully used for B/R assessment of vaccines.

1 ADVANCE system testing: benefit-risk analysis of a marketed vaccine using multi-criteria
2 decision analysis and cohort modelling

3 Kaatje **Bollaerts**^a, Edouard **Ledent**^b, Tom **de Smedt**^a, Daniel **Weibel**^c, Hanne-Dorthe
4 **Emborg**^d, Giorgia **Danieli**^e, Talita **Duarte-Salles**^f, Consuelo **Huerta**^g, Elisa **Martin**^g, Gino
5 **Picelli**^e, Lara **Tramontan**^e, Miriam **Sturkenboom**^{a,h,i}, Vincent **Bauchau**^b

6 ^a P95 Epidemiology and Pharmacovigilance, Koning Leopold III laan 1 3001 Heverlee,
7 Belgium (Kaatje.Bollaerts@p-95.com; tom.desmedt@p-95.com; miriam.sturkenboom@p-
8 95.com)

9 ^b GSK, Av. Fleming 20, 1300 Wavre, Belgium (edouard.y.ledent@gsk.com;
10 vincent.g.bauchau@gsk.com)

11 ^c Erasmus University Medical Center, Post box 2040, 3000 CA Rotterdam, The Netherlands
12 (d.weibel@erasmusmc.nl)

13 ^d Statens Serum Institut, Artillerivej 5, 2300 Copenhagen, Denmark (HDE@ssi.dk)

14 ^e Epidemiological Information for Clinical Research from an Italian Network of Family
15 Paediatricians (PEDIANET), Padova, Italy (gdanieliconsorzioarsenal@gmail.com;
16 g.picelli@virgilio.it; ltramontan@consorzioarsenal.it)

17 ^f Institut Universitari d'Investigació en Atenció Primària Jordi Gol (IDIAP Jordi Gol),
18 Barcelona, Spain (tduarte@idiapjgol.org)

19 ^g Base de Datos Para la Investigación Farmacoepidemiológica en Atención Primaria (BIFAP),
20 Spanish Agency of Medicines and Medical Devices (AEMPS). Madrid, Spain
21 (chuerta@aemps.es; emartinm@aemps.es)

22 ^h VACCINE.GRID, Spitalstrasse 33, Basel, Switzerland (mcjm.sturkenboom@gmail.com)

23 ⁱ Julius Global Health, University Medical Center Utrecht, Heidelberglaan 100, The
24 Netherlands (m.c.j.sturkenboom@umcutrecht.nl)

- 25 **Corresponding author:** Kaatje Bollaerts, Vlierbeeklaan 18, 3010 Kessel-lo, Belgium
- 26 Tel: + 32 485 789657; email: Kaatje.Bollaerts@p-95.com

1 ADVANCE system testing: benefit-risk analysis of a marketed vaccine using multi-criteria
2 decision analysis and cohort modelling

3 Kaatje **Bollaerts**^a, Edouard **Ledent**^b, Tom **de Smedt**^a, Daniel **Weibel**^c, Hanne-Dorthe
4 **Emborg**^d, Giorgia **Danieli**^e, Talita **Duarte-Salles**^f, Consuelo **Huerta**^g, Elisa **Martín-**
5 **Merino**^g, Gino **Picelli**^e, Lara **Tramontan**^e, Miriam **Sturkenboom**^{a,h,i}, Vincent **Bauchau**^b

6 ^a P95 Epidemiology and Pharmacovigilance, Koning Leopold III laan 1 3001 Heverlee,
7 Belgium (Kaatje.Bollaerts@p-95.com; tom.desmedt@p-95.com; miriam.sturkenboom@p-
8 95.com)

9 ^b GSK, Av. Fleming 20, 1300 Wavre, Belgium (edouard.y.ledent@gsk.com;
10 vincent.g.bauchau@gsk.com)

11 ^c Erasmus University Medical Center, Post box 2040, 3000 CA Rotterdam, The Netherlands
12 (d.weibel@erasmusmc.nl)

13 ^d Statens Serum Institut, Artillerivej 5, 2300 Copenhagen, Denmark (HDE@ssi.dk)

14 ^e Epidemiological Information for Clinical Research from an Italian Network of Family
15 Paediatricians (PEDIANET), Padova, Italy (gdanieliconsorzioarsenal@gmail.com;
16 g.picelli@virgilio.it; ltramontan@consorzioarsenal.it)

17 ^f Institut Universitari d'Investigació en Atenció Primària Jordi Gol (IDIAP Jordi Gol),
18 Barcelona, Spain (tduarte@idiapjgol.org)

19 ^g Base de Datos Para la Investigación Farmacoepidemiológica en Atención Primaria (BIFAP),
20 Spanish Agency of Medicines and Medical Devices (AEMPS). Madrid, Spain
21 (chuerta@aemps.es; emartinm@aemps.es)

22 ^h VACCINE.GRID, Spitalstrasse 33, Basel, Switzerland (mcjm.sturkenboom@gmail.com)

23 ⁱ Julius Global Health, University Medical Center Utrecht, Heidelberglaan 100, The
24 Netherlands (m.c.j.sturkenboom@umcutrecht.nl)

25 **Corresponding author:** Kaatje Bollaerts, Vlierbeeklaan 18, 3010 Kessel-lo, Belgium

26 Tel: + 32 485 789657; email: Kaatje.Bollaerts@p-95.com

27

28 **Abbreviations used**

ADVANCE	Accelerated Development of VAccine beNefit-risk Collaboration in Europe
aP	acellular pertussis
B/R	benefit-risk
eHR	electronic health record
HHE	hypotonic-hyporesponsive episode
ISR	injection-site reaction
MCDA	multi-criteria decision analysis
PROTECT	Pharmacoepidemiological Research on Outcome and Therapeutics
POC	proof-of-concept
RR	relative risk
VE	vaccine effectiveness
wP	whole-cell pertussis

29

30

31 **Abstract**

32 **Background**

33 The Accelerated Development of VAccine beNefit-risk Collaboration in Europe
34 (ADVANCE) is a public-private collaboration aiming to develop and test a system for rapid
35 benefit-risk (B/R) monitoring of vaccines using electronic health record (eHR) databases in
36 Europe. Proof-of-concept studies were designed to assess the proposed processes and system
37 for generating the required evidence to perform B/R assessment and near-real time monitoring
38 of vaccines. Most B/R assessment methodologies have been developed for medicinal
39 products. We aimed to test B/R methodologies for vaccines, using the comparison of the B/R
40 profiles of whole-cell (wP) and acellular pertussis (aP) vaccine formulations in children as an
41 example.

42 **Methods**

43 We used multi-criteria decision analysis (MCDA) to structure the B/R assessment combined
44 with cohort modelling to build the B/R effects table. In the cohort model, we simulated the
45 number of events in two hypothetical cohorts of 10^6 children followed from first pertussis
46 dose till pre-school-entry booster (or six years of age, whichever occurred first), with one
47 cohort receiving wP, and the other aP. The benefits were reductions in pertussis incidence and
48 complications. The risks were increased incidences of febrile convulsions, fever, hypotonic-
49 hyporesponsive episodes, injection-site reactions and persistent crying. The model parameters
50 were informed by estimates (coverage, background incidences, relative risks) from eHR
51 databases from Denmark (SSI), Spain (BIFAP and SIDIAP), Italy (Pedianet) and the UK
52 (RCGP-RSC and THIN). Preferences were elicited from clinical and epidemiological experts.

53 **Results**

54 Using cohort modelling to build the B/R effects table facilitated the comparison of different
55 effect measures (e.g. immediate relative risks vs long-term vaccine effectiveness). Estimates

56 from eHR databases could be used to inform the cohort model, with the cohort modelling
57 results being easily combined with preference weights to obtain B/R scores.

58 **Conclusion**

59 Existing B/R methodology, cohort modelling and estimates from eHR databases can be
60 successfully used for B/R assessment of vaccines.

61

62 **Keywords:** benefit-risk assessment; pertussis vaccines; methodological study; electronic
63 health record databases; Europe

64

65 **1. Introduction**

66 The Accelerated Development of VAccine beNefit-risk Collaboration in Europe project
67 (ADVANCE), launched in 2013 and funded by the Innovative Medicines Initiative (IMI), is a
68 public-private partnership aiming to develop and test a system for rapid benefit-risk (B/R)
69 assessment and near-real time monitoring of vaccines in the post-marketing setting [1] (see
70 Appendix for list of consortium members). A series of proof-of-concept (POC) studies were
71 designed to assess the proposed processes and system. The present study aimed to test a
72 methodology for the B/R assessment of vaccines and assess the use of European electronic
73 health record (eHR) databases for informing the B/R assessment.

74 There are several methodologies for B/R assessments that can support medical decision-
75 making [2]. In particular, the B/R ‘effects table’ is widely used following its recent
76 introduction in European Public Assessment Reports. However, other tools exist, including
77 frameworks, metrics, estimation and modelling techniques, as well as preference elicitation
78 techniques [2, 3]. The ‘Pharmacoepidemiological Research on Outcome and Therapeutics’
79 (PROTECT) consortium completed pioneering work in identifying, organising and appraising
80 B/R assessment tools [4]. Based on their experience drawn from eight case studies, they
81 recommended a systematic approach containing five generic steps; 1) planning, 2) evidence
82 gathering, 3) analysis, 4) exploration and 5) communication, rather than a one-size-fits-all
83 approach [5]. However, up to now, most B/R methodologies have been developed for
84 medicinal products while the B/R assessment of vaccines may require different methods [5].

85 When conducting a B/R assessment in a post-marketing setting, different information sources
86 can be used, including clinical trials, observational studies and systematic literature reviews.
87 Currently, there is a growing interest in using large eHR databases to study vaccine outcomes
88 (e.g. Vaccine Safety Datalink [6], the Post-Licensure Rapid Immunization Safety Monitoring

89 programme [7] and ADVANCE [1]) since these potentially enable real-world vaccine effects
90 to be studied on a large scale in geographical diverse settings.

91 To explore B/R assessment methodology for vaccines and the use of large eHR databases for
92 informing the B/R model, we compared the B/R profiles of whole-cell (wP) and acellular
93 pertussis (aP) vaccine formulations in children prior to their pre-school-entry booster as a test
94 case. This test case was selected to mimic the introduction of a new vaccine, where systematic
95 monitoring of changes in B/R profile over time would be needed. This POC study was
96 undertaken for system testing and not to inform clinical, regulatory or public health decisions
97 on pertussis vaccination.

98 **2. Methods**

99 ***2.1. Benefit-risk analysis***

100 We used multi-criteria decision analysis (MCDA) to structure the B/R assessment following
101 the PROTECT recommendations and ISPOR guidelines [5, 8, 9]. MCDA provides a
102 structured, stepwise approach for the assessment and comparison of different treatment
103 alternatives for benefit and risk outcomes [10]. However, the effect measures for vaccine
104 benefits, i.e., vaccine effectiveness (VE) and risks, i.e., relative risks (RRs) are different, with
105 VE being typically long-term and RRs being typically immediate and short-term. To facilitate
106 their comparison, we used cohort modelling with parameters informed by multi-country eHR
107 database studies on pertussis vaccination coverage, benefits and risks [11-13]. The results of
108 the cohort model were then combined with preference weights solicited from clinical and
109 epidemiological experts to obtain overall B/R scores.

110 ***2.2. Multi-criteria decision analysis***

111 Details on the different MCDA steps and their application are given below.

112 *Step 1: Establishment of the decision context*

113 The test case was the comparison of the B/R profiles of wP and aP vaccine formulations in
114 children prior to their pre-school-entry booster (or six years of age, whichever occurred first)
115 in Europe.

116 *Step 2: Identification of key benefit and risk criteria (value tree)*

117 The initial value tree was discussed and agreed by clinical and epidemiological experts from
118 public health, vaccine manufacturers and academia (**Figure 1**). In the final tree, indirect
119 effects were omitted for simplicity, limb swelling was combined with other injection-site
120 reactions to avoid double-counting; and convulsions were defined as febrile convulsions. The
121 final value tree contained reductions in pertussis and its complications (convulsions,
122 pneumonia and death) as benefit outcomes and febrile convulsions, fever, hypotonic-
123 hyporesponsive episodes (HHE), injection-site reactions (ISR) and persistent crying as risk
124 outcomes.

125 *Step 3: Identification of data sources*

126 The parameters of the B/R model were based on results for vaccination coverage, benefits and
127 risks from the ADVANCE multi-country eHR database studies where possible [11-13]. The
128 following databases were included in this study: SSI (Denmark), BIFAP and SIDIAP (Spain)
129 and RCGP RSC and THIN (UK) , PEDIANET (Italy) [14].

130 *Step 4: Construction of the benefit-risk effects table*

131 We used a cohort simulation model to build the B/R effects table. For the participating
132 countries (i.e., Denmark, Italy, Spain and the UK), we built two hypothetical cohorts of
133 1,000,000 children followed from their first pertussis dose until their pre-school booster (or
134 six years of age). One cohort was vaccinated with wP, the other with aP. Unvaccinated
135 children were not included as the B/R assessment focused on direct effects only (**Figure 1**).
136 To avoid the impact of time-varying confounding or changes in the background incidence

137 rates on the wP-aP comparison, the two hypothetical cohorts were identical with respect to the
138 age-specific background incidence rates, vaccination coverage and age at vaccination. Only
139 the vaccine type-specific parameters (i.e., VE and RR) were varied between the aP and wP
140 hypothetical cohorts.

141 The parameters for the simulation model were informed by the results from the ADVANCE
142 multi-country eHR database studies on pertussis vaccination coverage, benefits and risks with
143 the exception of pertussis VE and pertussis complication incidence rates, which were not
144 available when this B/R analysis was undertaken [11-13, 15-18]. Since the vaccination
145 schedules are different across countries, the dose-specific vaccination coverage and age at
146 vaccination were kept country-specific whereas the background incidence rates and vaccine-
147 type-specific RRs were pooled across countries to increase precision. To take into
148 consideration age-related dependencies, a finely disaggregated age-structure (monthly from
149 first dose to age two and 3-monthly afterwards) was used. Within each cohort, the expected
150 number of events for each outcome was estimated through Monte Carlo simulation based on
151 10^3 simulation draws. Median and 95% uncertainty intervals were obtained to account for
152 uncertainty in model parameters. The simulation models were developed using R version
153 3.4.0 [19]. The model input parameters are summarised in **Table 1**.

154 *Step 4.1: model input parameters: coverage*

155 To reflect recent practice, most recent coverage estimates with at least two years of follow up
156 were obtained from the ADVANCE coverage POC study (e.g. Denmark, Spain and UK: birth
157 cohort 2010; Italy: birth cohort 2007) [11]. The 2- and 3-dose country-specific coverage rates
158 at 24 months old and the age at vaccination were estimated for children who had received at
159 least 1 dose (**Figure 2**).

160 *Step 4.2: model input parameters: benefits*

161 The incidence of pertussis in unvaccinated children was derived from the incidence for those
162 who had received only one dose (since unvaccinated children were excluded from the
163 database study) and an estimate of the 1-dose VE obtained from the literature [12, 16]. To
164 reflect recent epidemiology, data from 2005 onwards were used to estimate age-specific
165 pertussis incidences, which were then pooled across databases using random effects meta-
166 analyses (**Figure 3**) [20].

167 *Step 4.3: model input parameters: risks*

168 Baseline incidences (2005 onwards) were used from primary care databases (BIFAP, RCGP
169 RSC, THIN and PEDIANET) for fever, ISR, persistent crying and somnolence (since these
170 mild outcomes are more likely to be reported in primary care); from the hospital discharge
171 database (SSI) for febrile convulsions (since this is a severe outcome likely to require
172 hospitalisation); and from all databases (primary care and hospital-based) for HHE (since this
173 can be a mild to severe outcome and therefore could be captured in any database). The age-
174 specific incidences were combined across databases (simply summing events and person
175 time) and subsequently smoothed using LOWESS (**Figure 4**) [21]. The database-specific RR
176 of adverse events during the exposure risk windows (by vaccine type and dose) were obtained
177 using a self-controlled case series method, which were subsequently pooled using random-
178 effects meta-analyses (estimates in **Table 1**) [13].

179 *Steps 5-6: Definition of value functions and preference weights*

180 Four clinical and epidemiological experts and three observers attended a preference elicitation
181 workshop, organised in compliance with ISPOR guidelines. After a face-to-face training
182 session on preference elicitation and practicing MCDA swing-weighting using D-Sight
183 software (www.d-sight.com) it was agreed to simplify the preference elicitation as the
184 participants found the swing-weighting difficult to understand, especially when non-linear

185 value functions were selected. It was therefore decided to restrict to linear value functions and
186 express, for each outcome, the number of events that would be equivalent to one pertussis
187 event (with pertussis being considered as the most severe outcome). For each outcome, the
188 lower limit of the linear value function was defined as the minimum lower limit of the 95%
189 uncertainty intervals of the number of events in both the hypothetical wP and aP cohort
190 whereas the upper limit of the linear value function was defined as the maximum upper limit
191 of the 95% uncertainty intervals.

192 *Step 7: Calculation of the benefit-risk scores*

193 The overall B/R scores for the wP and aP formulations (BR_j) were calculated as follows:

$$BR_j = \sum_i^I w_i \left(1 - \frac{N_{ij}^* - \min N_i}{\max N_i - \min N_i} \right) \times 100$$

194
195

196 This represents a linear value function, with $N_{ij}^* = \max(\min N_i, \min(N_{ij}, \max N_i))$, where
197 N_{ij} is the median number of events in the hypothetical cohort j for event type i , where $\min N_i$
198 and $\max N_i$ are the lower and upper limit of the linear value function and where w_i is the
199 preference weight.

200 *Step 8: Performing sensitivity analyses*

201 The impact of data uncertainty (as reflected by the 95% uncertainty intervals of the number of
202 events within the hypothetical populations) was assessed through Monte Carlo simulation
203 with 10^3 simulation runs, assuming normal distributions for the number of events for each
204 outcome. The impact of preference weights was assessed by halving and doubling a single
205 non-standardised preference weight, while keeping the others constant, and then standardising
206 again.

207 **3. Results**

208 ***3.1. Model input parameters: coverage, benefits and risks***

209 Within the selected birth cohorts, almost all children in each of the four countries, who
210 received the first dose, completed the schedule, with the 3-dose coverage rate ranging from
211 95.7% to 97.9% (**Figure 2**). The age at vaccination differed across countries in line with the
212 national recommendations. In Spain and the UK, all three doses were administered within the
213 first 10 months of life whereas in Italy and Denmark, the 3rd dose was administered between
214 10-15 months of age. Pertussis incidence after first dose (birth cohorts 2005 onwards) was
215 highest in children aged 2-3 months in all databases, although substantial database
216 heterogeneity existed, particularly for the two youngest age groups (as indicated by the I^2
217 statistic >75%) (**Figure 3**). The smoothed baseline risks are given in **Figure 4**, clearly
218 showing age trends for all event types. The RR of adverse events during the exposure risk
219 windows are given in **Table 1**.

220 ***3.2. Model results: benefit/risk effects table***

221 The total expected number of events for vaccine-related plus -unrelated outcomes, for both
222 the wP and aP cohorts, are summarised in **Table 2**. These estimates reflect the impact of
223 vaccination on the total disease burden in the population assuming the UK 2-dose and 3-dose
224 vaccination coverage rates and age at vaccination. Similar results were obtained assuming the
225 vaccination coverage rates and age at vaccination in Denmark, Italy and Spain
226 (**Supplementary tables S1, S2 and S3**).

227 ***3.3. Preference weights***

228 Good consensus was reached among the participants for all outcomes during the preference
229 elicitation workshop (**Figure 5a**). The experts attributed higher preference weights to the
230 benefit of preventing pertussis than to the risks of vaccination, with an averaged standardised
231 preference weight of 92.8% for prevented pertussis (**Table 2, Figure 5a**).

232 **3.4. Calculation of the benefit-risk scores**

233 Prevented pertussis was shown to make the largest contribution to the overall B/R scores and
234 was the most highly discriminating factor between aP and wP (**Figure 5b**).

235 **3.5. Impact of data uncertainty**

236 The Monte Carlo distributions of the overall B/R scores by vaccine type showed higher scores
237 for wP than for aP, although there was substantial overlap (**Figure 5c**). Changes in the
238 preference weights for pertussis and febrile convulsions had the largest impact on the overall
239 B/R score for wP whereas changes in the preference weight for fever had the largest impact
240 on the overall B/R score for aP (**Figure 5d**).

241 **4. Discussion**

242 These results demonstrated how existing B/R methodology can be used for post-marketing
243 B/R assessment of vaccines using evidence on vaccination coverage, benefits, and risks that
244 was obtained through dedicated studies in eHR databases. We adopted a structured approach
245 for the B/R assessment as recommended by the PROTECT project. To this end, we used
246 MCDA and combined MCDA with cohort modelling to build the B/R effects table. Although
247 MCDA has been used for vaccines before, to our knowledge, this is the first time it has been
248 combined with simulation modelling techniques to build a B/R effects table [22].

249 We used cohort simulation to build the B/R effects table, expressed as the total number of
250 simulated events for the different benefit and risk outcomes within two hypothetical
251 populations, each one receiving a different vaccine. The cohort simulation approach
252 facilitated the comparison of different effect measures (i.e., VE and RR), while accounting for
253 differences in age at vaccination, number of doses given, age-specific baseline risks and
254 differences in outcome-specific risk windows. We simulated the total number of events (i.e.
255 vaccine-related and -unrelated) to assess the impact of vaccination on the total disease burden.
256 We also complemented MCDA with additional Monte Carlo simulations assessing the impact

257 of data uncertainty on the overall B/R scores, which broadened the use of MCDA to decision-
258 making under uncertainty. Additional sensitivity analyses in which the preference weights
259 were varied enabled to assess the robustness of the B/R scores to changes in preference
260 weights. We used a cohort simulation model to build the B/R effects table since we
261 considered only direct effects, however, dynamic transmission models could have been used
262 if indirect effects were to be considered as well.

263 Evidence that can be used to inform the post-marketing B/R assessment models comes from
264 diverse sources and is of variable quality; here we assessed how evidence generated from
265 eHR databases could be used [23]. Compared with using available published evidence, the
266 approach we used has the advantage that different model parameters can be consistently
267 estimated with high levels of granularity, within the same study population.

268 We have shown how preference weights can be easily combined with the results from cohort
269 simulation models to obtain overall B/R scores. We obtained preference weights from clinical
270 and epidemiological experts as we believed they would have a good understanding of both the
271 benefits and risks of vaccination. We solicited preferences using MCDA swing-weighting,
272 which is one of the most efficient methods of obtaining preference weights as preferences can
273 be solicited during a one-day workshop. However, it requires training and a high level of
274 understanding by the participants. In our experience, the participants found the use of strongly
275 non-linear value functions in the swing-weighting process difficult. Therefore, we simplified
276 the preference elicitation by asking the participants to express the severity of each event by
277 giving the number of outcome events that would be equivalent to one pertussis event.

278 Further discussions on when, from whom and how to elicit preferences regarding vaccination
279 is needed as preference elicitation for vaccines raises many questions. Unlike drugs, vaccines
280 are mostly administered to healthy people, often to children as part of a vaccination
281 programme or mandate. This results in a very low public tolerance for vaccination risks,

282 despite the risks being rare. On the other hand, the benefits of vaccination are often invisible
283 as the incidence of many vaccine-preventable diseases has substantially decreased as a result
284 of vaccination. In addition, some vaccines have the potential to induce herd immunity,
285 whereby unvaccinated individuals are protected indirectly by those vaccinated, implying that
286 the benefits are not necessarily borne by the same individuals who take the risks.

287 Numerous methods for preference elicitation exist (e.g. time trade-offs, discrete choice
288 experiments, conjoint analyses) and these can be used in diverse settings, such as focus
289 groups or surveys. Alternatively, it would be possible to use composite burden of disease
290 measures such as disability-adjusted life years (DALY) to perform B/R assessments [24].
291 With this work, we only explored preference elicitation using MCDA swing-weighting.
292 Exploration of different preference elicitation techniques for vaccines is needed as well as
293 more ethical discussions on comparing disease prevented by vaccination and disease induced
294 by vaccination.

295 In conclusion we have shown that it is possible to use existing B/R methodology and
296 estimates from eHR databases to assess vaccines B/R successfully. We illustrated how cohort
297 modelling can be used to build the B/R effects table expressed as the expected number of
298 events in hypothetical cohorts, which facilitates the comparison of different effect measures.

299

300 **Acknowledgements**

301 The authors would like to thank Margaret Haugh, MediCom Consult, Villeurbanne, France
302 for editorial services and Lina Titievsky, Pfizer, USA, for project lead activities.

303

304

305 **Disclaimer**

306 The results described in this publication are from the proof of concept studies conducted as
307 part of the IMI ADVANCE project with the aim of testing the methodological aspects of the
308 design, conduct and reporting of studies for vaccine benefit-risk monitoring activities. The
309 results presented herein relate solely to the testing of these methodologies and are not
310 intended to inform regulatory or clinical decisions on the benefits and risks of the exposures
311 under investigation. This warning should accompany any use of the results from these studies
312 and they should be used accordingly.

313 The views expressed in this article are the personal views of the authors and should not be
314 understood or quoted as being made on behalf of or reflecting the position of the agencies or
315 organisations with which the authors are affiliated.

316

317

318 **Funding source**

319 The Innovative Medicines Initiative Joint Undertaking funded this project under ADVANCE
320 grant agreement n° 115557, resources of which were composed of a financial contribution
321 from the European Union's Seventh Framework Programme (FP7/2007-2013) and in kind
322 contributions from EFPIA member companies.

323

324

325 **Declaration of potential conflicts of interest**

326 Tom de Smedt, Hanne-Dorthe Emborg, Giorgia Danieli, Talita Duarte-Salles, Consuelo

327 Huerta, Elisa Martin, Gino Picelli, Lara Tramontan declared no potential conflicts of interest

328 Kaatje Bollaerts received consultancy fees from GSK unrelated to the submitted work.

329 Edouard Ledent and Vincent Bauchau declared that they are employees of the GSK group of

330 companies and hold company shares. Daniel Weibel declared that he has received

331 consultancy fees from GSK unrelated to this work. Miriam Sturkenboom declared that she has

332 received grants from Novartis, CDC and Bill & Melinda Gates Foundation, for work

333 unrelated to the submitted work.

334

335

336 **References**

- 337 [1] Sturkenboom M, Bahri P, Chiucchiuini A, Grove Krause T, S. H, Khromava A, et al.
338 Why we need more collaboration in Europe to enhance post-marketing surveillance of
339 vaccines. *Vaccine*. 2018;Paper 1 in ADVANCE supplement.
- 340 [2] Mt-Isa S, Hallgreen CE, Wang N, Callreus T, Genov G, Hirsch I, et al. Balancing
341 benefit and risk of medicines: a systematic review and classification of available
342 methodologies. *Pharmacoepidemiol Drug Saf*. 2014;23:667-78.
- 343 [3] Guo JJ, Pandey S, Doyle J, Bian B, Lis Y, Raisch DW. A review of quantitative risk-
344 benefit methodologies for assessing drug safety and efficacy-report of the ISPOR risk-benefit
345 management working group. *Value Health*. 2010;13:657-66.
- 346 [4] Innovative Medicines Initiative PROTECT project 2010. Available from:
347 <http://www.imi-protect.eu/index.shtml>.
- 348 [5] Hughes D, Waddingham E, Mt-Isa S, Goginsky A, Chan E, Downey GF, et al.
349 Recommendations for benefit-risk assessment methodologies and visual representations.
350 *Pharmacoepidemiol Drug Saf*. 2016;25:251-62.
- 351 [6] McNeil MM, Gee J, Weintraub ES, Belongia EA, Lee GM, Glanz JM, et al. The
352 Vaccine Safety Datalink: successes and challenges monitoring vaccine safety. *Vaccine*.
353 2014;32:5390-8.
- 354 [7] Nguyen M, Ball R, Midthun K, Lieu TA. The Food and Drug Administration's post-
355 licensure rapid immunization safety monitoring program: strengthening the federal vaccine
356 safety enterprise. *Pharmacoepidemiol Drug Saf*. 2012;21 Suppl 1:291-7.
- 357 [8] Marsh K, M IJ, Thokala P, Baltussen R, Boysen M, Kalo Z, et al. Multiple criteria
358 decision analysis for health cared decision making--emerging good practices: report 2 of the
359 ISPOR MCDA Emerging Good Practices Task Force. *Value Health*. 2016;19:125-37.

360 [9] Thokala P, Devlin N, Marsh K, Baltussen R, Boysen M, Kalo Z, et al. Multiple criteria
361 decision analysis for health care decision making--an introduction: report 1 of the ISPOR
362 MCDA Emerging Good Practices Task Force. *Value Health*. 2016;19:1-13.

363 [10] Mussen F, Salek,S., Walker, S. Benefit- risk apraisal of mdicines: a systematic
364 approach to decision- making. Chichester: John Wiley & Sons; 2009.

365 [11] Emborg HD, Berensci K, Braeye T, Bauwens J, Bollaerts K, Correa A, et al.
366 ADVANCE system testing: can coverage of pertussis vaccination be estimated in EU
367 countries using electronic health data: an example. *Vaccine*. 2018;Paper 4 in Supplement.

368 [12] Htar MTT, de Ridder M, Braeye T, Correa A, de Lusignan S, Duarte-Salles T, et al.
369 ADVANCE system testing: vaccine benefit studies by using multi-country electronic health
370 data: An example on pertussis vaccination. *Vaccine*. 2018;Paper 5 in Supplement.

371 [13] Weibel D, Dodd C, Mahaux O, Haguinet F, de Smedt T, Duarte-Salles T, et al.
372 ADVANCE system testing: can safety studies be conducted using electronic health data: an
373 example with pertussis vaccination? *Vaccine*. 2018;Paper 6 in Supplement.

374 [14] Sturkenboom M, Weibel D, van der Aa L, Braeye T, Gheorge M, Becker B, et al.
375 ADVANCE database characterization and fit for purpose assessment for multi-country studies
376 on the coverage, benefits and risks of vaccinations. *Vaccine*. 2018;Paper 3 in Supplement.

377 [15] Farrington CP. Relative incidence estimation from case series for vaccine safety
378 evaluation. *Biometrics*. 1995;51:228-35.

379 [16] Bisgard KM, Rhodes P, Connelly BL, Bi D, Hahn C, Patrick S, et al. Pertussis vaccine
380 effectiveness among children 6 to 59 months of age in the United States, 1998-2001.
381 *Pediatrics*. 2005;116:e285-94.

382 [17] Fulton TR, Phadke VK, Orenstein WA, Hinman AR, Johnson WD, Omer SB.
383 Protective effect of contemporary pertussis vaccines: a systematic review and meta-analysis.
384 *Clin Infect Dis*. 2016;62:1100-10.

385 [18] Zanardi L, Pascual FB, Bisgard K, Murphy T, Wharton M. Pertussis --- United States,
386 1997--2000. *MMWR Morb Mortal Wkly Rep.* 2002;51:73-6.

387 [19] R Development Core Team. *R: a language and environment for statistical computing.*
388 Vienna, Austria2013.

389 [20] DerSimonian R, Laird N. Meta-analysis in clinical trials. *Control Clin Trials.*
390 1986;7:177-88.

391 [21] Cleveland WS. LOWESS: A program for smoothing scatterplots by robust locally
392 weighted regression. *Am Stat.* 1981;35:54-.

393 [22] Marcelon L, Verstraeten T, Dominiak-Felden G, Simondon F. Quantitative benefit-
394 risk assessment by MCDA of the quadrivalent HPV vaccine for preventing anal cancer in
395 males. *Expert Rev Vaccines.* 2016;15:139-48.

396 [23] Hallgreen CE, van den Ham HA, Mt-Isa S, Ashworth S, Hermann R, Hobbiger S, et
397 al. Benefit-risk assessment in a post-market setting: a case study integrating real-life
398 experience into benefit-risk methodology. *Pharmacoepidemiol Drug Saf.* 2014;23:974-83.

399 [24] McDonald SA, Nijsten D, Bollaerts K, Bauwens J, Praet N, van der Sande M, et al.
400 Methodology for computing the burden of disease of adverse events following immunization.
401 *Pharmacoepidemiol Drug Saf.* 2018.

402

403 **Figure captions**

404 **Figure 1** Initial and final pertussis vaccination outcome trees. The outcomes that were not
405 retained for the final outcome tree are shaded in grey. (aP: acellular pertussis vaccines; wP:
406 whole-cell pertussis vaccines; HHE: hypotonic-hypo-responsive episodes).

407 **Figure 2** Age-specific vaccination coverage rates (%) for children who received at least one
408 dose, by dose and country (Denmark, Spain and UK: birth cohort 2010; Italy: birth cohort
409 2007). The horizontal lines at the top indicate coverage for children aged 24 months for each
410 dose. Data from [11].

411 **Figure 3** Database-specific and meta-analysed pertussis incidence (/100.000 person-years
412 (py)) among children who received one dose, by age group, 2005 onwards [12]. Estimates
413 that were considered to be outliers, i.e. absolute value of the studentised residual was >2.5
414 (indicated by *) were excluded from the meta-analysis. Study heterogeneity was investigated
415 by the chi-squared test for heterogeneity, (p-values <0.05 indicate a significant amount of
416 heterogeneity), and quantified using the I^2 statistic with low, moderate and high levels of
417 heterogeneity corresponding to I^2 values of 25%, 50% and 75%, respectively.

418 **Figure 4** Database-specific, pooled and LOWESS smoothed pooled estimates (with 95% CI)
419 for age-specific baseline incidences (/1000 persons-years (py)) by risk outcome, 2005
420 onwards [13, 21].

421 **Figure 5** a) Individual and averaged preference weights. b) Overall B/R score and outcome
422 contributions by vaccine formulation (aP vs wP). c) Impact of data uncertainty: distribution of
423 the overall B/R scores by vaccine type obtained through Monte Carlo simulation. d) Impact of
424 preference weights: changes in the overall B/R scores when doubling (red arrows) or halving
425 (blue arrows) the preference weights one-at-the-time.

426

427

428 **Table 1** Cohort simulation: overview of model parameters.

Parameter	Mean [95% CI]		Distribution	Source(s)
	aP	wP		
Coverage				
Coverage at 24 months		Figure 2	Binomial distribution on number of vaccinated children. Probability of being vaccinated is the coverage at 24 mos.	[11]
Age at vaccination (in months)		Figure 2	Empirical distribution	[11]
Benefits*				
Pertussis				
Age-specific incidence among unvaccinated subjects (/100.000 py) (< 6 years)		Figure 3	Empirical: incidences in children with 1 dose only divided by $(1 - VE_{d1})$, $VE_{d1} = 74\%$	[12, 16]
Vaccine effectiveness – dose 1	0.66 [0.56; 0.71]	0.7 [0.62; 0.72]	Log-normal on relative risk ($RR = 1 - VE$). Meta-analysed 3-dose VE estimate multiplied with VE ratio = 74.1%	[17] [16]
Vaccine effectiveness – dose 2	0.83 [0.71; 0.89]	0.88 [0.84; 0.94]	Log-normal on relative risk ($RR = 1 - VE$). Meta-analysed 3-dose VE estimate multiplied with VE ratio = 93.6%	[17] [16]
Vaccine effectiveness – dose 3	0.89 [0.76; 0.95]	0.94 [0.89; 0.97]	Log-normal on relative risk ($RR = 1 - VE$). Meta-analysed estimate	[17]
Pertussis-related pneumonia (age-specific % of cases with complications)	<6 months: 11.8%; 6-11 months: 8.6%; 1-4 years: 5.4%		Binomial distribution on number of cases with complications. Probability of developing complication is age-specific	[18]
Pertussis-related febrile seizures (age-specific % of cases with complications)	<6 months: 1.4%; 6-11 months: 0.7%; 1-4 years: 1.2%		Binomial distribution on number of cases with complications. Probability of developing complication is age-specific	[18]
Pertussis-related deaths (age-specific % of cases with complications)	<6 months: 0.8%; 6-11 months: 0.1%; 1-4 years: <0.1%		Binomial distribution on number of cases with complications. Probability of developing complication is age-specific	[18]
Risks**				
Febrile convulsions				
Age-specific baseline incidence (/1000 py)		Figure 4	Empirical	[13]
Relative risk (0-3d) – dose 1	0.89 [0.51; 1.57]	1.15 [0.63; 2.11]	Log-normal on relative risk. Meta-analysed estimate.	[13]
Relative risk (0-3d) – dose 2	0.94 [0.79; 1.11]	1.51 [0.71; 3.19]	Log-normal on relative risk. Meta-analysed estimate.	[13]

Parameter	Mean [95% CI]		Distribution	Source(s)
	aP	wP		
Relative risk (0-3d) – dose 3	2.19 [1.69; 2.83]	1.89 [1.55; 2.31]	Log-normal on relative risk	Meta-analysed estimate. [13]
Fever				
Age-specific baseline incidence (/1000 py)		Figure 4		[13]
Relative risk (0-3d) – dose 1	1.18 [1.08; 1.29]	1.92 [1.84; 2.00]	Log-normal on relative risk	Meta-analysed estimate. [13]
Relative risk (0-3d) – dose 2	0.89 [0.81; 0.99]	1.47 [1.42; 1.54]	Log-normal on relative risk	Meta-analysed estimate. [13]
Relative risk (0-3d) – dose 3	1.17 [0.98; 1.39]	1.85 [1.78; 1.92]	Log-normal on relative risk	Meta-analysed estimate. [13]
Hypotonic-hyproresponsive episodes				
Age-specific baseline incidence (/1000 py)		Figure 4		[13]
Relative risk (0-2d) – dose 1	2.72 [1.49; 4.96]	1.70 [1.30; 2.24]	Log-normal on relative risk	Meta-analysed estimate. [13]
Relative risk (0-2d) – dose 2	1.42 [0.73; 2.79]	0.71 [0.36; 1.42]	Log-normal on relative risk	Meta-analysed estimate. [13]
Relative risk (0-2d) – dose 3	1.65 [0.81; 3.39]	1.34 [1.01; 1.78]	Log-normal on relative risk	Meta-analysed estimate. [13]
Injection site reactions				
Age-specific baseline incidence (/1000 py)		Figure 4		[13]
Relative risk (0-2d) – dose 1	1.38 [1.15; 1.65]	2.12 [1.89; 2.38]	Log-normal on relative risk	Meta-analysed estimate. [13]
Relative risk (0-2d) – dose 2	1.78 [1.09; 2.91]	2.42 [2.13; 2.74]	Log-normal on relative risk	Meta-analysed estimate. [13]
Relative risk (0-2d) – dose 3	1.65 [0.81; 3.39]	2.19 [1.95; 2.45]	Log-normal on relative risk	Meta-analysed estimate. [13]
Persistent crying				
Age-specific baseline incidence (/1000 py)		Figure 4		[13]
Relative risk (0-1d) – dose 1	1.56 [0.97; 2.50]	4.60 [3.89; 5.45]	Log-normal on relative risk	Meta-analysed estimate. [13]
Relative risk (0-1d) – dose 2	1.28 [0.92; 1.76]	2.76 [1.90; 4.01]	Log-normal on relative risk	Meta-analysed estimate. [13]
Relative risk (0-1d) – dose 3	1.05 [0.57; 1.92]	2.13 [1.82; 2.47]	Log-normal on relative risk	Meta-analysed estimate. [13]
Somnolence				
Age-specific baseline incidence (/1000 py)		Figure 4		[13]
Relative risk (0-3d) – dose 1	3.01 [1.30; 7.00]	4.47 [1.74; 11.5]	Log-normal on relative risk	Meta-analysed estimate. [13]
Relative risk (0-3d) – dose 2	2.16 [1.60; 2.90]	1.10 [0.70; 1.71]	Log-normal on relative risk	Meta-analysed estimate. [13]
Relative risk (0-3d) – dose 3	1.29 [0.78; 2.14]	1.94 [1.28; 2.94]	Log-normal on relative risk	Meta-analysed estimate. [13]

429 Abbreviations: aP = acellular pertussis vaccine, CI = confidence interval, d = days, mos = months, py = person-year, RR = relative risk, VE = vaccine effectiveness.
430 *Pertussis: Probability of developing pertussis is derived from the age-specific incidence in unvaccinated subjects multiplied with (1-VE), where VE is dose-dependent.
431 Probability of developing pertussis complications is age-specific, with the age-specific probabilities being derived from the literature **Risk events: probability of a risk event
432 outside the risk window is derived from the age-specific incidence. Probability of an event during the risk windows is derived from the age-specific incidence multiplied with
433 the corresponding RR.
434

435 **Table 2:** Benefit/risk effects table, lower and upper limit of linear value functions and standardized averaged preference weights for the
 436 outcomes of interest.

Event	Benefit/risk effects table		Linear value function		Preference weights (%)
	Number of events*		Lower limit	Upper limit	Averaged
	Median [95% uncertainty intervals]				
	aP	wP			
Benefits					
Pertussis	1,292 [686; 2,467]	698 [440; 1,186]	420	2,500	92.8
Pertussis complications					
Convulsions	15 [5; 30]	8 [2; 16]	n.a.	n.a.	n.a.
Death	3 [0; 8]	2 [0; 5]	n.a.	n.a.	n.a.
Pneumonia	101 [51; 188]	56 [34; 93]	n.a.	n.a.	n.a.
Risks					
Febrile convulsions	69,702 [68,899; 70,564]	69,377 [68,730; 70,054]	68,000	71,000	3.8
Fever	539,768 [538,798; 540,702]	541,063 [540,102; 542,017]	540,000	541,500	2.4
Hypotonic-hyporesponsive episodes	2,283 [2,187; 2,382]	2,262 [2,167; 2,355]	2,100	2,400	0.9
Injection site reactions	2,819 [2,709; 2,930]	2,896 [2,782; 3,003]	2,700	3,000	0.1
Persistent crying	25,477 [25,135; 25,859]	26,690 [26,268; 27,139]	25,000	27,500	<0.01
Somnolence	1,783 [1,689; 1,877]	1,799 [1,704; 1,917]	1,600	2,000	<0.01

437 * Cohort simulation model; number of events in a hypothetical cohort of 1,000,000 children followed from first dose till pre-school booster; one cohort received aP, the other wP. The
 438 vaccination coverage and age at vaccination are reflective of the UK. n.a. = data was not available at the time of the preference elicitation and these outcomes were excluded from the overall
 439 B/R score.
 440

441 **Supplementary tables**

442

443 **Table S1.** B/R effects table: number of events by type of event in a hypothetical cohort of
 444 1,000,000 children followed from first dose till pre-school booster; one cohort received aP,
 445 the other wP. The vaccination coverage and age at vaccination are reflective of Denmark.

446

Event	Benefit/risk effects table	
	Number of events*	
	Median [95% uncertainty intervals]	
	aP	wP
Pertussis	1,458 [843; 2,391]	885 [522; 1,237]
Convulsions	16 [7; 30]	10 [3; 18]
Death	4 [1; 8]	2 [0; 6]
Pneumonia	115 [66; 184]	73 [41; 105]
Febrile convulsions	77,168 [69,227; 83,023]	75,270 [68,986; 78,921]
Fever	534,203 [533,066; 540,377]	536,572 [535,495; 541,650]
Hypotonic-hypo-responsive episodes	2,119 [2,014; 2,346]	2,104 [1,999; 2,320]
Injection site reactions	2,560 [2,449; 2,878]	2,611 [2,504; 2,959]
Persistent crying	20,173 [19,829; 25,705]	20,954 [20,569; 26,948]
Somnolence	1,619 [1,527; 1,843]	1,633 [1,536; 1,861]

447

448

449

450 **Table S2.** B/R effects table: number of events by type of event in a hypothetical cohort of
451 1,000,000 children followed from first dose till pre-school booster; one cohort received aP,
452 the other wP. The vaccination coverage and age at vaccination are reflective of Italy.

453

Event	Benefit/risk effects table	
	Number of events*	
	Median [95% uncertainty intervals]	
	aP	wP
Pertussis	1,464 [843: 2,407]	892 [522: 1,240]
Convulsions	16 [7: 30]	10 [3: 18]
Death	4 [1: 9]	3 [0: 6]
Pneumonia	116 [66: 184]	74 [41: 105]
Febrile convulsions	77,581 [69,227: 83,612]	75,603 [68,986: 79,272]
Fever	534,464 [533,333: 540,377]	536,837 [535,735: 541,650]
Hypotonic-hyporesponsive episodes	2,125 [2,020: 2,346]	2,109 [2,004: 2,320]
Injection site reactions	2,568 [2,454: 2,878]	2,617 [2,507: 2,959]
Persistent crying	20,330 [19,979: 25,705]	21,104 [20,714: 26,948]
Somnolence	1,625 [1,534: 1,843]	1,638 [1,542: 1,861]

454

455

456

457 **Table S3.** B/R effects table: number of events by type of event in a hypothetical cohort of

458 1,000,000 children followed from first dose till pre-school booster; one cohort received aP,

459 the other wP. The vaccination coverage and age at vaccination are reflective of Spain.

460

Benefit/risk effects table

Number of events*

Median [95% uncertainty intervals]

Event	aP	wP
Pertussis	1,364 [771; 2,462]	758 [505; 1,212]
Convulsions	16 [6; 31]	9 [3; 17]
Death	4 [1; 9]	2 [0; 6]
Pneumonia	109 [61; 189]	63 [39; 97]
Febrile convulsions	72,255 [69,227; 74,831]	71,334 [68,986; 72,962]
Fever	539,255 [538,193; 540,431]	541,001 [540,038; 541,978]
Hypotonic-hyporesponsive episodes	2,222 [2,114; 2,353]	2,201 [2,097; 2,325]
Injection site reactions	2,725 [2,606; 2,878]	2,792 [2,680; 2,959]
Persistent crying	23,055 [22,677; 25,705]	24,102 [23,671; 26,948]
Somnolence	1,723 [1,625; 1,843]	1,734 [1,635; 1,868]

461

462

463 **Appendix: Members of ADVANCE consortium (October 2018)**

464 **Full partners**

465 AEMPS: Agencia Española de Medicamentos y Productos Sanitarios (www.aemps.es)

466 ARS-Toscana: Agenzia regionale di sanità della Toscana (<https://www.ars.toscana.it/it/>)

467 ASLCR: Azienda Sanitaria Locale della Provincia di Cremona (www.aslcremona.it)

468 AUH: Aarhus Universitetshospital (kea.au.dk/en/home)

469 ECDC: European Centre of Disease Prevention and Control (www.ecdc.europa.eu)

470 EMA: European Medicines Agency (www.ema.europa.eu)

471 EMC: Erasmus Universitair Medisch Centrum Rotterdam (www.erasmusmc.nl)

472 GSK: GlaxoSmithKline Biologicals (www.gsk.com)

473 IDIAP: Jordi Gol Fundació Institut Universitari per a la Recerca a l'Atenció Primària de Salut

474 Jordi Gol i Gurina (<http://www.idiapjordigol.com>)

475 JANSSEN: Janssen Vaccines - Prevention B.V. (<http://www.janssen.com/infectious-diseases-and-vaccines/crucell>)

477 KI: Karolinska Institutet (ki.se/meb)

478 LSHTM: London School of Hygiene & Tropical Medicine (www.lshtm.ac.uk)

479 MHRA: Medicines and Healthcare products Regulatory Agency (www.mhra.gov.uk/)

480 MSD: Merck Sharp & Dohme Corp. (www.merck.com)

481 NOVARTIS: Novartis Pharma AG (www.novartisvaccines.com)

482 OU: The Open University (www.open.ac.uk)

483 P95: P95 (www.p-95.com)

484 PEDIANET: Società Servizi Telematici SRL (www.pedianet.it)

485 PFIZER: Pfizer Limited (www.pfizer.co.uk)

486 RCGP: Royal College of General Practitioners (www.rcgp.org.uk)

487 RIVM: Rijksinstituut voor Volksgezondheid en Milieu (www.rivm.nl)

488 SCIENSANO: Sciensano (<https://www.sciensano.be>)

489 SP: Sanofi Pasteur (www.sanofipasteur.com)

490 SSI: Statens Serum Institut (www.ssi.dk)

491 SURREY: The University of Surrey (www.surrey.ac.uk)

492 SYNAPSE: Synapse Research Management Partners, S.L. (www.synapse-managers.com)

493 TAKEDA: Takeda Pharmaceuticals International GmbH (www.tpi.takeda.com)

494 UNIBAS-UKBB: Universitaet Basel – Children’s Hospital Basel (www.unibas.ch)

495 UTA: Tampereen Yliopisto (www.uta.fi)

496 **Associate partners**

497 AIFA: Italian Medicines Agency (www.agenziafarmaco.it)

498 ANSM: French National Agency for Medicines and Health Products Safety (ansm.sante.fr)

499 BCF: Brighton Collaboration Foundation (brightoncollaboration.org)

500 EOF: Hellenic Medicines Agency, National Organisation for Medicines (www.eof.gr)

501 FISABIO: Foundation for the Promotion of Health and Biomedical Research
502 (www.fisabio.es)

503 HCDCP: Hellenic Centre for Disease Control and Prevention (www.keelpno.gr)

504 ICL: Imperial College London (www.imperial.ac.uk)

505 IMB/HPRA: Irish Medicines Board (www.hpra.ie)

506 IRD: Institut de Recherche et Développement (www.ird.fr)

507 NCE: National Center for Epidemiology (www.oek.hu)

508 NSPH: Hellenic National School of Public Health (www.nsph.gr)

509 PHE: Public Health England (www.gov.uk/government/organisations/public-health-england)

510 THL: National Institute for Health and Welfare (www.thl.fi)

511 UMCU: Universitair Medisch Centrum Utrecht (www.umcu.nl)

512 UOA: University of Athens (www.uoa.gr)

- 513 UNIME: University of Messina (www.unime.it)
- 514 Vaccine.Grid: Vaccine.Grid (<http://www.vaccinegrid.org/>)
- 515 VVKT: State Medicines Control Agency (www.vvkt.it)
- 516 WUM: Polish Medicines Agency - Warszawski Uniwersytet Medyczny
517 (<https://wld.wum.edu.pl/>)
- 518

Figure 1

[Click here to download high resolution image](#)

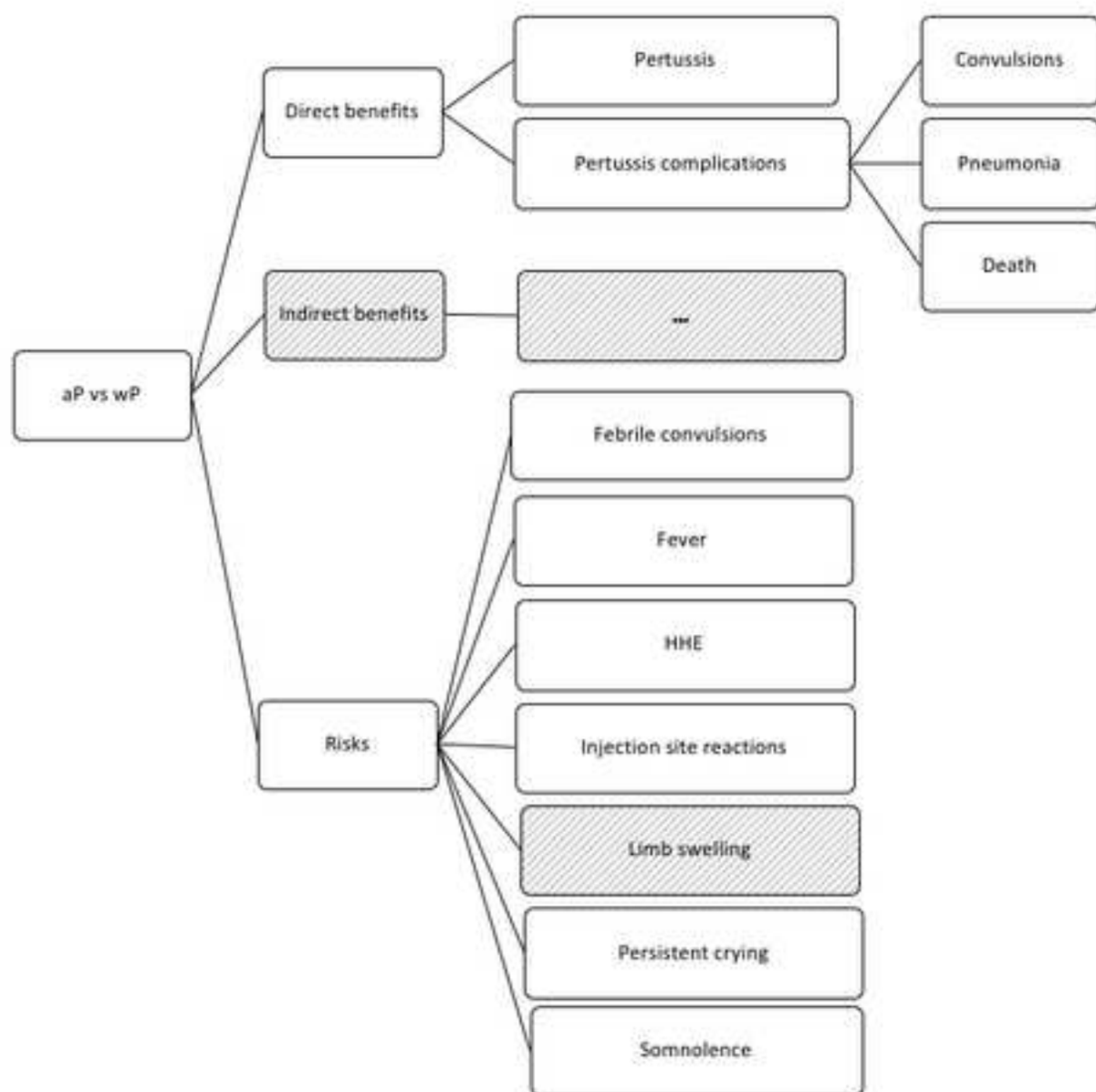


Figure 2

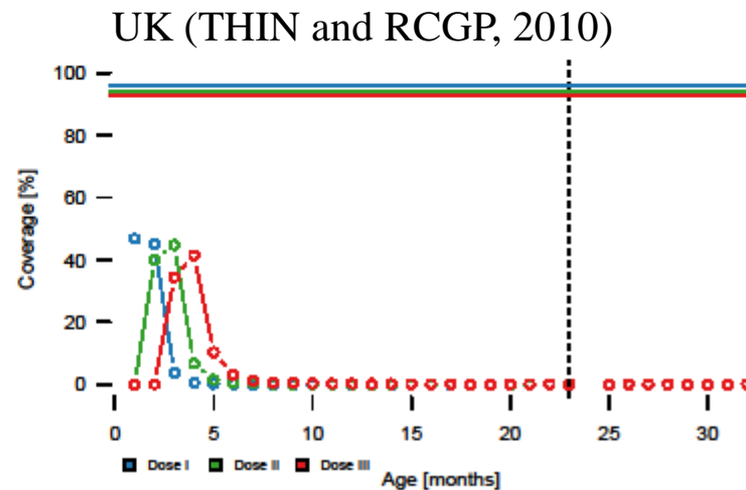
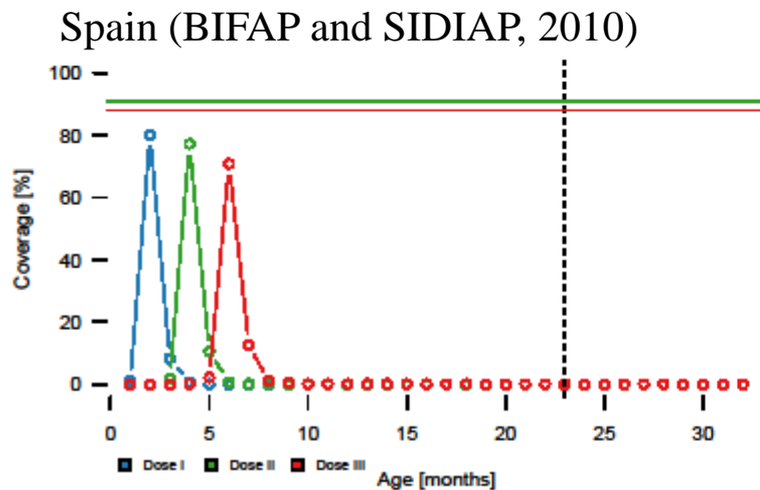
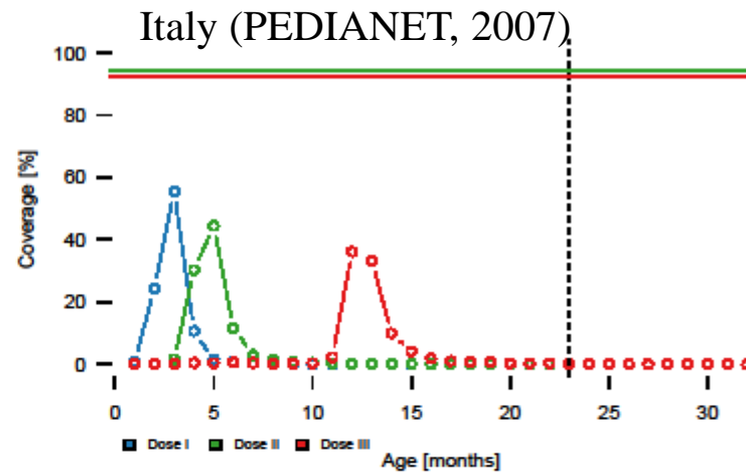
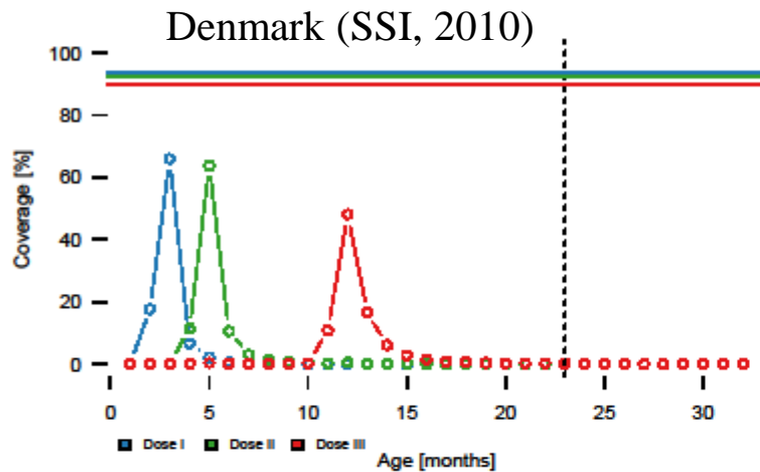


Figure 3
[Click here to download high resolution image](#)

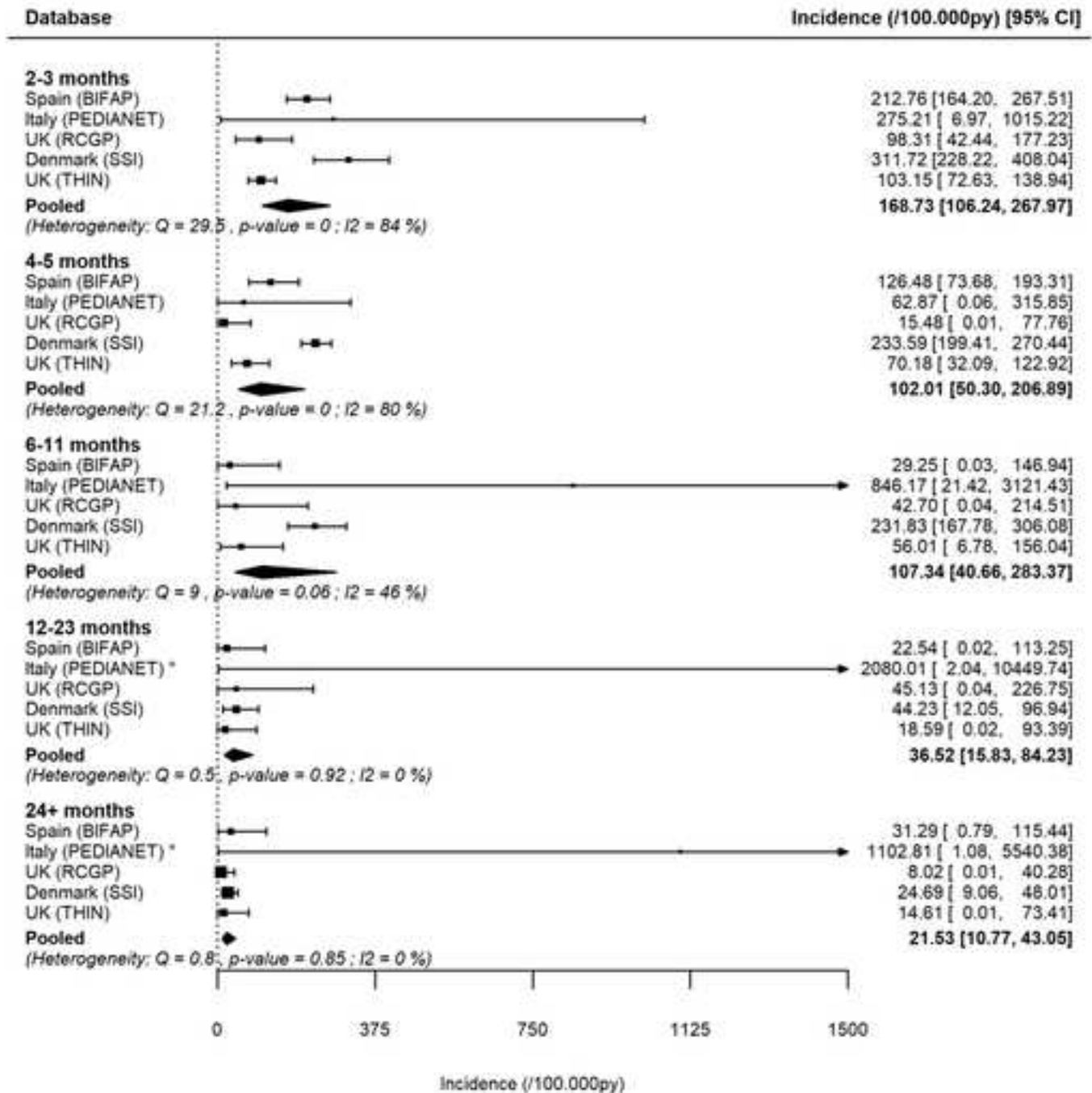


Figure 4

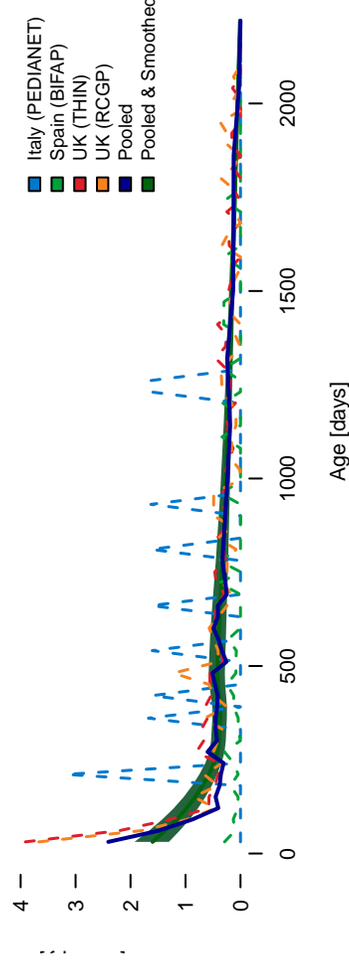
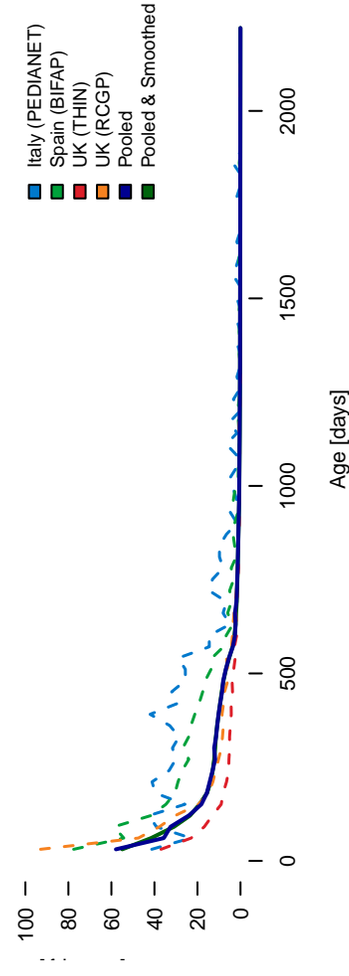
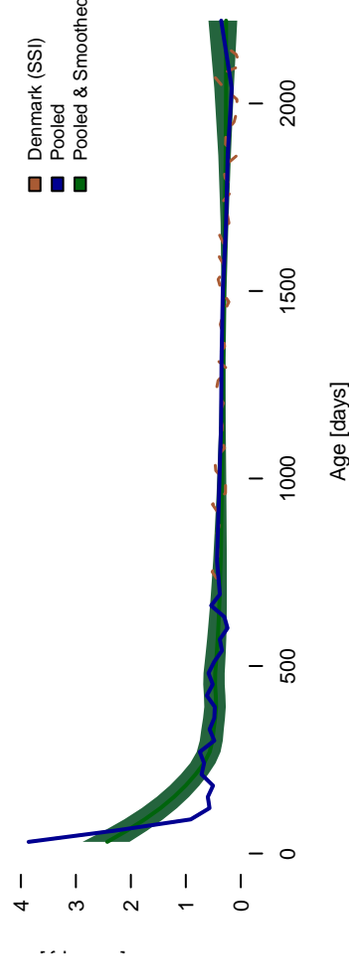
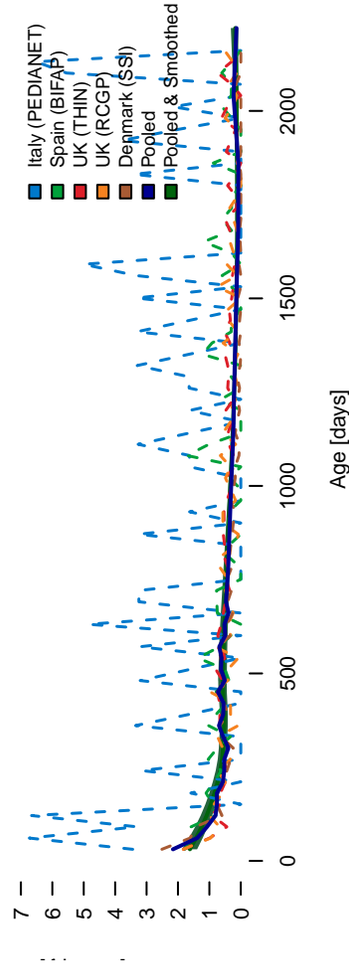
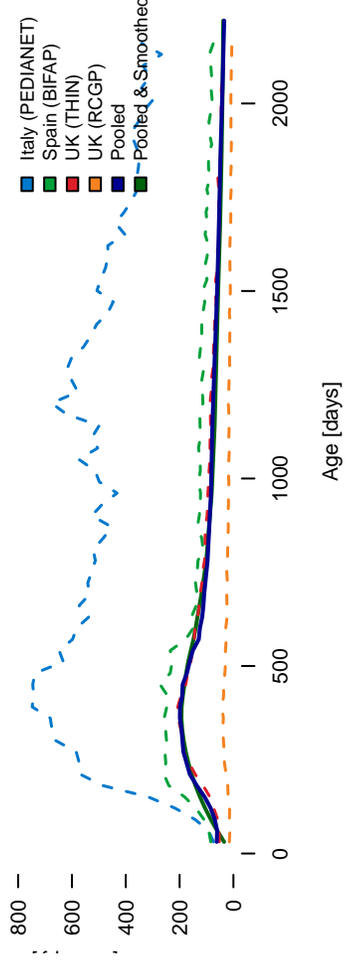
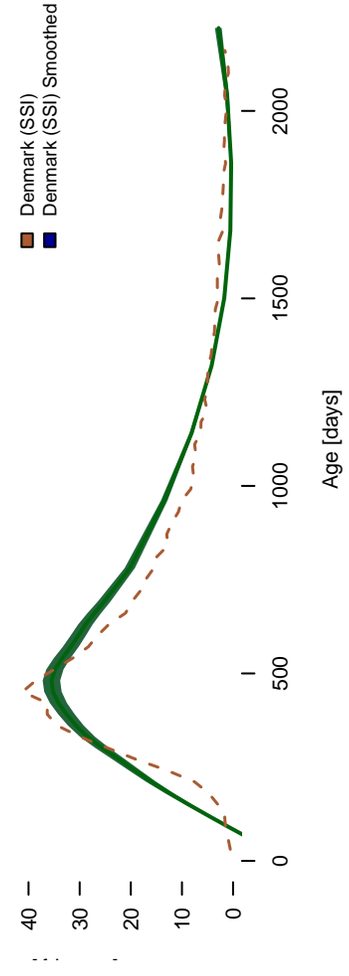


Figure 5
[Click here to download high resolution image](#)

