

Safety of Vaccines Used for Routine Immunization in the United States: An Updated Systematic Review and Meta-analysis

Courtney Gidengil, M.D., M.P.H.,^{a,b} Matthew Bidwell Goetz, M.D.,^c Sydne Newberry, Ph.D.,^d
Margaret Maglione, M.P.P.,^d Owen Hall, M.S.,^d Jody Larkin, M.S.,^d Aneesa Motala, B.A.,^{d, e}
Susanne Hempel, Ph.D.^{d, e}

^aRAND Corporation, 20 Park Plaza, Suite 920, Boston, MA, 02116

^bBoston Children's Hospital, 300 Longwood Avenue, Boston, MA, 02115

^cVA Greater Los Angeles Healthcare System and David Geffen School of Medicine, University of California, Los Angeles, Los Angeles, CA, 90073

^dRAND Corporation, 1776 Main Street, Santa Monica, CA, 90401

^eSouthern California Evidence Review Center, University of Southern California, Keck School of Medicine, 2001 N Soto Street, Los Angeles, CA, 90033

Corresponding author: Courtney Gidengil, MD MPH
RAND Corporation
20 Park Plaza, Suite 920
Boston, MA 02116
Tel: 617-338-2059
Email: gidengil@rand.org

[Click here to view linked References](#)

1 **Safety of Vaccines Used for Routine Immunization in the United States: An Updated**
2 **Systematic Review and Meta-analysis**

3

4 Courtney Gidengil, M.D., M.P.H.,^{a,b} Matthew Bidwell Goetz, M.D.,^c Sydne Newberry, Ph.D.,^d
5 Margaret Maglione, M.P.P.,^d Owen Hall, M.S.,^d Jody Larkin, M.S.,^d Aneesa Motala, B.A.,^{d,e}
6 Susanne Hempel, Ph.D.^{d,e}

7

8 ^aRAND Corporation, 20 Park Plaza, Suite 920, Boston, MA, 02116

9 ^bBoston Children’s Hospital, 300 Longwood Avenue, Boston, MA, 02115

10 ^cVA Greater Los Angeles Healthcare System and David Geffen School of Medicine, University
11 of California, Los Angeles, Los Angeles, CA, 90073

12 ^dRAND Corporation, 1776 Main Street, Santa Monica, CA, 90401

13 ^eSouthern California Evidence Review Center, University of Southern California, Keck School
14 of Medicine, 2001 N Soto Street, Los Angeles, CA, 90033

15

16 Corresponding author: Courtney Gidengil, MD MPH

17 RAND Corporation

18 20 Park Plaza, Suite 920

19 Boston, MA 02116

20 Tel: 617-338-2059

21 Email: gidengil@rand.org

22

23 **Abstract**

24 **Background:** Understanding the safety of vaccines is critical to inform decisions about
25 vaccination. Our objective was to conduct a systematic review of the safety of vaccines
26 recommended for children, adults, and pregnant women in the United States.

27 **Methods:** We searched the literature in November 2020 to update a 2014 Agency for Healthcare
28 Research and Quality review by integrating newly available data. Comparative studies that
29 reported the presence or absence of key adverse events were eligible. Adhering to Evidence-
30 based Practice Center methodology, we assessed the strength of evidence (SoE) for all evidence
31 statements. The systematic review is registered in PROSPERO (CRD42020180089).

32 **Results:** Of 56,603 reviewed citations, 338 studies reported in 518 publications met inclusion
33 criteria. For children, SoE was high for no increased risk of autism following measles, mumps,
34 and rubella (MMR) vaccine. SoE was high for increased risk of febrile seizures with MMR.
35 Rotavirus vaccine was not associated with intussusception at latest time of follow-up (moderate
36 SoE), nor with increased risk of diabetes (high SoE). There was no evidence of increased risk or
37 insufficient evidence for key adverse events for newer vaccines such as 9-valent human
38 papillomavirus and meningococcal B vaccines. For adults, there was no evidence of increased
39 risk (varied SoE) for key adverse events for the new recombinant adjuvanted zoster vaccine and
40 hepatitis B vaccine with novel immunostimulatory adjuvant. We found no evidence of increased
41 risk (varied SoE) for key adverse events among pregnant women following tetanus, diphtheria,
42 and acellular pertussis vaccine, including for preterm labor and stillbirth (moderate SoE).

43 **Conclusions:** Across a large body of research we found few associations of vaccines and
44 serious key adverse events; however, rare events are challenging to study. Any adverse events
45 should be weighed against the protective benefits that vaccines provide.

46

47 **Keywords:** vaccine safety; meta-analysis; systematic review; childhood vaccines; pregnancy

48

49 **Introduction**

50 Vaccines are considered one of the greatest public health achievements and the effectiveness
51 of vaccines in controlling the spread of and even eradicating many infectious diseases is widely
52 acknowledged [1]. Although vaccination rates for children remain high, parents and caregivers
53 still express worries about the safety of childhood vaccines [2-4]. Vaccination rates for adults lag
54 well behind those for children [5]. Only about a third of pregnant women receive both tetanus,
55 diphtheria, and acellular pertussis (Tdap) and influenza vaccines as indicated during their
56 pregnancies [6], due in part to safety concerns [7].

57 Safety concerns about vaccines have persisted in spite of the rigorous, transparent processes
58 that vaccines must undergo, overseen in the United States by the U.S. Food and Drug
59 Administration (FDA) [8]. Once a vaccine is licensed and recommended for use following
60 clinical trials, multiple systems are in place to ensure ongoing assessments of safety through
61 Phase IV studies [9], including post-licensure safety surveillance [10] and the FDA's Post-
62 Licensure Rapid Immunization Monitoring (PRISM) system [11-13]. Multiple databases
63 contribute to surveillance, such as the Vaccine Adverse Event Reporting System (VAERS) [14],
64 Vaccine Safety Datalink [15, 16], and Clinical Immunization Safety Assessment project [17, 18].

65 Reassurance of vaccine safety remains critical for population health in the context of an
66 evolving vaccine landscape and notably the emergence of vaccines against the novel severe acute
67 respiratory syndrome coronavirus (SARS-CoV-2). The purpose of this systematic review was to
68 assess the evidence regarding the safety of vaccines routinely recommended for adults, children,
69 and pregnant women in the United States.

70

71 **Methods**

72 The evidence review assessed and examined adverse events potentially associated with
73 vaccines to determine the safety of vaccines in adults, children, and pregnant women, following
74 the Agency for Healthcare Research and Quality’s Methods Guide for Effectiveness and
75 Comparative Effectiveness Reviews [19] (full details can be found in the online Appendix). The
76 list of included vaccines comprises those licensed by the FDA [20] and included in the CDC’s
77 immunization schedules as of November 2020 (Table 1) [21, 22].

78 This update builds on a 2014 report on the safety of vaccines requested by AHRQ [23],
79 supporting the Office of the Assistant Secretary of Health’s Office of Infectious Disease and
80 HIV/AIDS Policy (OASH/OIDP). The 2014 report built upon a 2011 Institute of Medicine
81 consensus report [24]; the prior 2014 report did not search for or include studies on vaccines that
82 were covered in the IOM report and published prior to 2011. Similarly, in this update only for
83 vaccines for which there were new indications (or for new vaccines) did we perform targeted
84 searches for research published prior to 2014. The review is registered in PROSPERO
85 (CRD42020180089) [25], and the review protocol is posted on the EHC website [26].

86 We searched MEDLINE (including TOXLINE), Embase, CINAHL, Cochrane CENTRAL,
87 Web of Science, and Scopus, through November 2020 (see Appendix for full search strategy).
88 We searched broadly and did not rely on filters for adverse events. Instead, all evaluations of
89 vaccines were obtained and the full text screened for information on adverse events. We
90 reference-mined existing systematic reviews and Advisory Committee on Immunization
91 Practices statements; screened Clinicaltrials.gov; reviewed supplemental material from authors
92 and industry submitted to AHRQ; and consulted with content experts. Experimental and
93 observational studies with a concurrent or historic comparator that reported the presence or
94 absence of adverse events (e.g., self-controlled studies such as those conducted by the Vaccine

95 Safety Datalink [15]) met inclusion criteria. The update allowed for control groups receiving
96 either no vaccine or standard of care (i.e., the previously available vaccine) as comparators. With
97 the assistance of a technical expert panel—comprised of vaccine experts with particular clinical
98 expertise in key populations (children, adults, older adults, and pregnant women), vaccine safety
99 methodologists, and consumers—we determined a set of key adverse events *a priori* to allow an
100 unbiased synthesis across studies.

101 Two trained reviewers (with Master’s degrees and experience in systematic reviews)
102 independently screened the citations and full text publications. Data were abstracted by an
103 experienced subject matter expert with clinical and research expertise in vaccines (C.G.). For
104 each key adverse event, we computed the relative risk (RR) and 95% confidence intervals (CI) of
105 the adverse event among those who received the vaccine of interest compared to controls across
106 all studies. We combined estimates across studies in random effects meta-analyses using
107 Hartung-Knapp correction of standard errors where appropriate. For cases with zero events
108 across studies, we added a constant to the empty cell to enable computation. We determined the
109 most appropriate meta-analysis model (see Appendix), given that for many adverse events only a
110 small number of studies were available, studies reported on rare events, and several studies
111 reported zero events [27-29]. Where studies did not report sufficient detail and could not be
112 combined into the meta-analysis, we reported the risk estimates provided by the authors.

113 All studies that reported rates of adverse events that could be computed were combined in
114 meta-analyses. When studies could not be combined statistically, we narratively synthesized the
115 findings to inform the strength of evidence (SoE) assessment and ensure that all available
116 evidence was considered. For the synthesis we determined whether there was evidence of an
117 increased risk of adverse events relative to a control group. In addition to the relative effect, we

118 also documented the actual incidences, sample sizes, and resulting rates of adverse events in the
119 vaccinated and control groups for each individual study where available. We reviewed all
120 instances where the vaccinated group had reported more instances of adverse events in detail. In
121 the absence of evidence of an increased risk across studies, we also reviewed the risk reported in
122 individual studies and documented the observed rates. For estimates that were imprecise—given
123 the small number of reported events and the small number of samples from which conclusions
124 for the true risk could be estimated—the narrative synthesis also reports observed rates to
125 transparently document the available evidence.

126 We used McHarm [30] for critical appraisal of individual studies, rating studies that reported
127 timing and severity and used standard, precise definitions of adverse events higher than studies
128 that did not. The body of evidence was assessed based on AHRQ Evidence-based Practice
129 Center grading [31]. We used four criteria to grade the SoE: (1) study limitations; (2)
130 consistency; (3) precision; and (4) reporting bias. We differentiated *high*, *moderate*, *low*, and
131 *insufficient* evidence to communicate the confidence for the findings across studies. *High*
132 confidence indicates that the evidence reflects the true effect; further research is very unlikely to
133 change our confidence in the estimate of effect. *Moderate* confidence indicates that the evidence
134 reflects the true effect; further research may change our confidence in the estimate of effect and
135 may change the estimate. *Low* confidence indicates that the evidence reflects the true effect;
136 further research is likely to change our confidence in the estimate of effect and is likely to
137 change the estimate. *Insufficient* indicates that evidence either is unavailable or does not permit a
138 conclusion.

139 Findings are reported below for the selected key adverse events (adverse events identified in
140 the prior report that were not selected as key adverse events for this update are included in the

141 Appendix). We report effect estimates (RR and 95% CI) that could be computed for findings of
142 moderate or high SoE across studies; we also report findings that were of low SoE, but not the
143 effect estimates.

144

145 **Results**

146 Of 56,603 reviewed citations, 189 new studies met inclusion criteria in this update for a total
147 of 338 studies reported in 518 publications across the prior report and update (Figure 1) [32-
148 549]. Study designs included RCTs, cohort studies, pre-post designs, case-control designs, non-
149 randomized controlled clinical trials, and self-controlled studies (either self-controlled risk
150 interval or self-controlled case series analyses). Many studies followed patients for six months or
151 longer, and some for up to 15 years to record emerging adverse events.

152 The methodological rigor and reporting of the adverse events over 15 assessed domains
153 varied widely across studies (Appendix Table 1; Appendix Figure 1). Most studies reported the
154 timing and frequency of the adverse events assessment, but few reported the qualifications of the
155 outcome assessors.

156 Full study characteristics can be found in Appendix Tables 2, 3, and 4.

157 ***Safety of vaccines included in the routine immunization schedule in children***

158 A summary of the strength of evidence for the findings can be found in Table 2 (all effect
159 estimates and assessments of the quality of the evidence are in Appendix Table 5, followed by
160 synthesis of the SoE across the prior report and update in Appendix Table 5a).

161 **9-valent human papillomavirus vaccine.** All but one study compared 9-valent human
162 papillomavirus vaccine to 2- or 4-valent vaccines. We also reviewed studies that combined

163 children and adults. There was no evidence of increased risk of autoimmune disease, birth
164 defects, death, reproductive system events, seizures, or spontaneous abortion (all low SoE).

165 **13-valent pneumococcal vaccine.** Risk estimates were based on comparisons of 13-valent
166 pneumococcal vaccine to 7-valent pneumococcal vaccine, except for death. There was no
167 evidence of increased risk of death (RR 2.02; CI 0.07, 59.88; risk estimate based on 1 study;
168 moderate SoE assessed across all 5 available studies). The risk estimate was imprecise as the
169 sample size was small with only one event (1/193 vs 0/195). There was also no evidence of
170 increased risk of asthma, cardiovascular events, intussusception, meningitis, reproductive system
171 events, or seizures (all low SoE). There was an increased risk of febrile seizures (low SoE).
172 There was insufficient evidence for 23-valent pneumococcal polysaccharide vaccine for the
173 outcomes of interest.

174 **Diphtheria, tetanus, and pertussis (DTaP) vaccine.** There was no evidence of increased
175 risk of type 1 diabetes mellitus (moderate SoE, effect estimate N/A). There was no evidence of
176 increased risk of asthma or death (low SoE).

177 **Tetanus, diphtheria & acellular pertussis vaccine.** There was no evidence of increased risk
178 of cardiovascular events or death (low SoE).

179 ***Haemophilus influenzae* type b vaccine.** There was no evidence of increased risk of serious
180 adverse events in the short term (moderate SoE; effect estimates N/A).

181 **Hepatitis A vaccine.** There was an increased risk of idiopathic thrombocytopenic purpura
182 (moderate SoE, effect estimate N/A).

183 **Hepatitis B vaccine.** There was no evidence of increased risk of multiple sclerosis (moderate
184 SoE, effect estimate N/A).

185 **Inactivated poliovirus vaccine.** There was insufficient evidence for conclusions about
186 increased risk of key adverse events.

187 **Quadrivalent influenza vaccines (IIV).** Quadrivalent IIV was compared to trivalent IIV in
188 all but one of the studies that contributed to risk estimates (this study only contributed to the risk
189 estimate for death). There was no evidence of increased risk of death (RR 1.08; CI 0.02, 53.95;
190 estimate based on 1 study; moderate SoE across 6 studies). The risk estimate was imprecise
191 because it was based on one small study with no deaths (0/99 vs 0/107). There was no evidence
192 of increased risk of anaphylaxis or systemic allergic reaction, asthma, autoimmune disease,
193 cardiovascular events, febrile seizures, or seizures (low SoE).

194 Quadrivalent live attenuated influenza vaccine (LAIV) was compared to placebo or no
195 vaccine in some studies, or another influenza vaccine (trivalent LAIV or IIV) in other studies.
196 There was no evidence of increased risk of death (when compared to trivalent LAIV) or seizures
197 (when compared to placebo or no vaccine) (low SoE).

198 **Measles, mumps, and rubella vaccine.** There was no evidence of an association with autism
199 (RR 0.60; CI 0.09, 4.12; 2 studies; high SoE across studies from 2014 report and update). There
200 was an increased risk of anaphylaxis (in children with allergies; high SoE), febrile seizures (high
201 SoE), and idiopathic thrombocytopenic purpura (moderate SoE) (effect estimates N/A).

202 **Serogroup A, C, W, and Y meningococcal vaccines.** Some studies of serogroup A, C, W,
203 and Y meningococcal vaccines used another meningococcal vaccine as an active comparator,
204 while others used a non-active comparator (placebo or a base treatment received by both
205 intervention and control groups). All estimates below are based on studies of children with a
206 non-active comparator, but studies where an active comparator was used as well as studies of
207 both children and adults also contribute to the SoE. There was no evidence of increased risk of

208 cardiovascular events (RR 0.34; CI 0.02, 5.46; estimate based on 1 study; moderate SoE across 3
209 studies), febrile seizures (RR 0.51; CI 0.18, 1.44; based on 1 study; moderate SoE across 4
210 studies), intussusception (RR 0.46; CI 0.10, 2.03; 1 study; moderate SoE), idiopathic
211 thrombocytopenic purpura (RR 0.17; CI 0.01, 5.09; based on 1 study; moderate SoE across 3
212 studies), Kawasaki disease (RR 1.37; CI 0.15, 12.22; based on 1 study; moderate SoE across 2
213 studies), or seizures (RR 1.51; CI 0.05, 44.86; based on 1 study; moderate SoE across 7 studies).

214 There was no evidence of increased risk of diabetes (RR 1.32; CI 0.00, 21861366; based on 2
215 studies; moderate SoE across 6 studies) but the risk estimate was imprecise due to small samples
216 and few or no cases in the vaccinated and unvaccinated groups (1/396 vs 0/397; 0/392 vs 0/296).

217 There was also no evidence of increased risk of acute disseminated encephalomyelitis,
218 asthma, autoimmune disease, death, encephalitis/encephalopathy, meningitis, multiple sclerosis,
219 reproductive system events, or transverse myelitis (low SoE). There was increased risk of
220 anaphylaxis in children with allergies (moderate SoE; effect estimate N/A), but there was no
221 evidence of increased risk among all children (low SoE).

222 **Serogroup B meningococcal vaccine.** There was no evidence of increased risk of
223 anaphylaxis or systemic allergic reaction (RR 0.56; CI 0.00, 34735108; 2 studies; moderate
224 SoE), but the risk estimate was imprecise due to no cases in the vaccinated and unvaccinated
225 groups (0/198 vs 0/121; 0/992 vs 0/501). There was no evidence of increased risk of reproductive
226 system events (RR 0.89; CI 0.01, 65.20; 3 studies; moderate SoE); again, the risk estimate was
227 imprecise, in this case due to small samples and few or no cases in the vaccinated and
228 unvaccinated groups (1/198 vs 0/121; 1/174 vs 0/99; 0/374 vs 1/378). There was no evidence of
229 increased risk of asthma, death, or seizures (low SoE).

230 **Rotavirus vaccine.** We found no evidence of increased risk of intussusception across all
231 studies that could be combined for an estimate (RR 0.65; CI 0.41, 1.05; based on 19 studies;
232 moderate SoE across 38 studies), though some observational studies indicated increased risk,
233 particularly around the first dose. There was no evidence of increased risk of asthma (RR 1.33;
234 CI 0.65, 2.72; 5 studies; moderate SoE), autoimmune disease (RR 0.65; CI 0.16, 2.67; 2 studies;
235 moderate SoE), death (RR 1.05; CI 0.82, 1.35; based on 14 studies; moderate SoE across 15
236 studies), diabetes (RR 0.74; CI 0.45, 1.22; based on 3 studies; high SoE across 4 studies), febrile
237 seizures (RR 0.82; CI 0.33, 2.05; based on 7 studies; moderate SoE across 9 studies), or seizures
238 (RR 1.02; CI 0.25, 4.16; based on 5 studies; moderate SoE across 8 studies).

239 There was no evidence of increased risk of encephalitis/encephalopathy (RR 0.67; CI 0.00,
240 85995; based on 2 studies; moderate SoE across 4 studies), but the risk estimate was imprecise
241 due to few or no cases in the vaccinated and unvaccinated groups (1/1647 vs 2/1641; 1/34904 vs
242 1/34862). There was no evidence of increased risk of idiopathic thrombocytopenic purpura (RR
243 0.64; CI 0.00, 1778394; 2 studies; moderate SoE); again, the risk estimate was imprecise due
244 there being few or no cases in the vaccinated and unvaccinated groups (1/34904 vs 0/34862;
245 0/4359 vs 2/4328). There was no evidence of increased risk of stroke (RR 1.32; CI 0.00,
246 1459247; 2 studies; moderate SoE). The risk estimate is imprecise given the small number of
247 studies and studies reporting few or no cases in the vaccinated and unvaccinated groups (1/34904
248 vs 0/34862; 1/1666 vs 1/1667).

249 There was no evidence of increased risk of anaphylaxis or systemic allergic reaction,
250 autoimmune thyroiditis (Hashimoto's disease), Kawasaki disease, meningitis, or reproductive
251 system events (all low SoE).

252 **Varicella vaccine.** There was evidence of increased risk of anaphylaxis (high SoE) and
253 idiopathic thrombocytopenic purpura (among children aged 11-17 years; moderate SoE) (effect
254 estimates N/A).

255 ***Safety of vaccines included in the routine immunization schedule in adults***

256 A summary of the strength of evidence for the findings can be found in Table 3 (all effect
257 estimates and assessments of the quality of the evidence are in Appendix Table 6, followed by
258 synthesis of the SoE in Appendix Table 6a).

259 **13-valent pneumococcal conjugate vaccine.** Some studies of 13-valent pneumococcal
260 conjugate vaccine used another pneumococcal vaccine as an active comparator, while others
261 used a non-active comparator (placebo or a base treatment received by both intervention and
262 control groups). All risk estimates below are based on studies with a non-active comparator, but
263 studies where an active comparator was used also contribute to the SoE. There was no evidence
264 of increased risk of cardiovascular events (RR 0.97; CI 0.58, 1.64; based on 4 studies; moderate
265 SoE across 6 studies), myocardial infarction (RR 1.76; CI 0.42, 7.39; based on 4 studies;
266 moderate SoE across 6 studies), or reproductive system events (RR 0.59; CI 0.01, 42.46; based
267 on 3 studies; moderate SoE across 5 studies).

268 There was no evidence of increased risk of herpes zoster (RR 1.49; CI 0.00, 24855526; 2
269 studies; moderate SoE). The risk estimate was imprecise as only two studies reported on the
270 outcome with few or no cases occurring in the vaccinated and unvaccinated groups (0/576 vs
271 0/575; 1/42237 vs 0/42255). There was also no evidence of increased risk of or stroke (RR 1.12;
272 CI 0.00, 451; 2 studies; moderate SoE); the risk estimate was imprecise due to no events in one
273 study (0/551 vs 0/560), and a large sample size with a small number of events in the other
274 (9/42237 vs 8/42255).

275 We found no evidence of increased risk of acute disseminated encephalomyelitis,
276 anaphylaxis or systemic allergic reactions, asthma, autoimmune disease, death,
277 encephalitis/encephalopathy, idiopathic thrombocytopenic purpura, meningitis, or seizures (low
278 SoE).

279 **23-valent pneumococcal polysaccharide vaccine.** We found no evidence of increased risk
280 of death (RR 1.45; CI 0.00, 3455; based on 2 studies; moderate SoE across 4 studies). The risk
281 estimate was imprecise due to a very small study with no events (0/19 vs 0/21) and a larger study
282 with few events (6/725 vs 4/724). We also found high SoE for no evidence of increased risk of
283 cardiovascular events (RR 0.46; CI 0.27, 0.76; based on 4 studies; high SoE across 8 studies) or
284 cerebrovascular events (effect estimate N/A) in people aged 65 years and older.

285 **Hepatitis B vaccine.** For HEPLISAV-B (which was compared to previously available
286 hepatitis B vaccines), there was no evidence of increased risk for asthma, autoimmune disease,
287 cardiovascular events, death, herpes zoster, reproductive system events, and stroke (low SoE).
288 For all hepatitis B vaccine, there was no evidence of increased risk of diabetes (RR 0.61; CI
289 0.55, 0.67; based on 1 study comparing hepatitis B vaccines to no vaccine; moderate SoE across
290 2 studies). For hepatitis B vaccines (not including HEPLISAV-B) there was no increased risk of
291 multiple sclerosis, but there was increased risk of anaphylaxis in patients allergic to yeast (both
292 moderate SoE; effect estimates N/A).

293 **Influenza vaccines (IIV).** Influenza vaccines were compared to an active comparator (either
294 trivalent influenza vaccine or another influenza vaccine). For quadrivalent IIV, we identified no
295 evidence of increased risk of asthma, cardiovascular events, death, myocardial infarction,
296 reproductive system events, seizures, or stroke (low SoE). For adjuvanted IIV (either trivalent or
297 quadrivalent), there was no evidence of stroke (RR 1.18; CI 0.00, 33607; based on 2 studies;

298 moderate SoE across 3 studies). The risk estimate was imprecise as only two studies reported on
299 the outcome with few or no cases occurring in the vaccinated and comparator groups (0/888 vs
300 1/888, 3/3545 vs 2/3537). There was no evidence of increased risk of asthma, autoimmune
301 disease, cardiovascular events, death, encephalitis/encephalopathy, Guillain-Barré syndrome,
302 idiopathic thrombocytopenic purpura, myocardial infarction, or seizures (low SoE). For
303 quadrivalent recombinant influenza vaccine, there was no evidence of increased risk of
304 cardiovascular events, death, encephalitis/encephalopathy, myocardial infarction, reproductive
305 system events, or stroke (low SoE). There was insufficient evidence for conclusions about
306 increased risk of key adverse events for quadrivalent live attenuated influenza vaccine.

307 **Measles, mumps, rubella vaccine.** There was no evidence of increased risk of type 1
308 diabetes mellitus (moderate SoE, effect estimate N/A).

309 **Serogroups A, C, W and Y meningococcal vaccines.** Some studies of serogroup A, C, W,
310 and Y meningococcal vaccines used another meningococcal vaccine as an active comparator,
311 while others used a non-active comparator (placebo or a base treatment received by both
312 intervention and control groups); all risk estimates below are based on studies with a non-active
313 comparator, but studies where an active comparator was used also contribute to the SoE. There
314 was no evidence of increased risk of death (RR 0.99; CI 0.00, 60563320; based on 2 studies;
315 moderate SoE across 4 studies). The risk estimate was imprecise due to two small studies with
316 no events (00/99 vs 0/100; 0/85 vs 0/84). There was no evidence of increased risk of
317 cardiovascular events, myocardial infarction, or stroke (all low SoE).

318 **Tetanus, diphtheria, and acellular pertussis and tetanus and diphtheria vaccines.** There
319 was evidence of increased risk of anaphylaxis (high SoE, effect estimate N/A).

320 **Recombinant zoster vaccine.** We found moderate SoE of no evidence of increased risk of
321 cardiovascular events (RR 0.89; CI 0.66, 1.21; 3 studies), death (RR 0.93; CI 0.78, 1.11; 4
322 studies), myocardial infarction (RR 0.89; CI 0.38, 2.05; 3 studies), or reproductive system events
323 (RR 1.04; CI 0.03, 37.17; 2 studies).

324 For all other adverse events for which there was moderate SoE, the confidence intervals were
325 wide because the risk estimate was based on two studies with few or no events occurring in the
326 vaccinated and non-vaccinated groups: amyotrophic lateral sclerosis (RR 2.60; CI 0.00, 571537;
327 2/6950 vs 0/6950, 2/7695 vs 1/7710), anaphylaxis or systemic allergic reaction (RR 1.32; CI
328 0.00, 1463200; 1/6950 vs 1/6950, 1/7695 vs 0/7710), asthma (RR 0.90; CI 0.00, 493; 2/6950 vs
329 4/6950, 6/7695 vs 5/7710), diabetes (RR 1.00; CI 0.00, 606; 5/6950 vs 6/6950, 3/7695 vs
330 2/7710), encephalitis/encephalopathy (RR 0.50; CI 0.00, 2867570; 0/6950 vs 1/6950, 0/7695 vs
331 1/7710), Guillain-Barré syndrome (RR 0.67; CI 0.00, 86459; 1/6950 vs 2/6950, 1/7695 vs
332 1/7710), idiopathic thrombocytopenic purpura (RR 2.65; CI 0.00, 530690; 1/6950 vs 0/6950,
333 3/7695 vs 1/7710), meningitis (RR 0.50; CI 0.00, 2867570; 0/6950 vs 1/6950, 0/7695 vs 1/7710),
334 seizures (RR 1.34; CI 0.00, 13492; 2/6950 vs 0/6950, 3/7695 vs 3/7710), or stroke (RR 1.44; CI
335 0.03, 71.52; 7/6950 vs 6/6950, 19/7695 vs 12/7710).

336 We found no evidence of increased risk of herpes zoster (RR 0.09; CI 0.02, 0.30; 5 studies;
337 high SoE). There was no evidence of increased risk of acute disseminated encephalomyelitis,
338 angioedema, ataxia, autoimmune disease, or autoimmune thyroiditis (low SoE).

339 **9-valent human papillomavirus vaccine, hepatitis A vaccine, combination hepatitis A
340 and hepatitis B vaccine, serogroup B meningococcal vaccine, and varicella vaccine.**

341 Evidence was insufficient to draw conclusions about increased risk of key adverse events based
342 on studies of adults only.

343 *Safety of vaccines included in the routine immunization schedule in pregnant women (both*
344 *for the woman and her fetus)*

345 A summary of the strength of evidence for the findings is in Table 4 (all effect estimates and
346 assessments of the quality of the evidence are in Appendix Table 7, followed by synthesis of the
347 SoE in Appendix Table 7a).

348 We found insufficient evidence to draw conclusions about increased risk of key adverse
349 events for hepatitis B vaccine, quadrivalent inactivated influenza vaccines, or quadrivalent
350 recombinant influenza vaccine in pregnant women.

351 All studies of Tdap compared to either placebo or base treatment also received by the control
352 groups, except for one study that compared Tdap and Td. There was no evidence of increased
353 risk for maternal cardiovascular events (RR 0.86; CI 0.41, 1.84; 6 studies), maternal death (RR
354 1.52; CI 0.07, 32.25; 4 studies), maternal diabetes (RR 0.98; CI 0.88, 1.10; 4 studies),
355 eclampsia/pre-eclampsia (RR 0.96; CI 0.92, 1.01; 6 studies), preterm labor (RR 0.62; CI 0.46,
356 0.82; 10 studies), maternal reproductive system events (RR 0.52; CI 0.05, 5.91; 3 studies),
357 stillbirth (RR 0.44; CI 0.11, 1.80; 6 studies), cardiovascular events in infants (RR 0.77; CI 0.50,
358 1.20; 4 studies), death in infants (RR 0.15; CI 0.00, 8.88; 3 studies), encephalitis/encephalopathy
359 in infants (RR 1.23; CI 0.60, 2.54; 4 studies), or seizures in infants (RR 1.02; CI 0.76, 1.35; 3
360 studies) (all moderate SoE). There was also no evidence of increased risk of maternal
361 encephalitis/encephalopathy, autism in infants, birth defects in infants, or febrile seizures in
362 infants (low SoE).

363

364 **Discussion**

365 We assessed the evidence for the safety of vaccines currently used for routine immunization
366 in the United States among children, adults, and pregnant women. We conducted extensive
367 literature searches, screened 56,603 citations, and abstracted 338 studies reported in 518
368 publications.

369 Overall, our evidence review found vaccines to be safe across populations with serious
370 adverse events being rare, consistent with other recent systematic reviews of vaccine safety
371 [550]. For adults, there was no evidence of increased risk (varied SoE) or insufficient evidence
372 for the new recombinant adjuvanted zoster vaccine and hepatitis B vaccine with novel
373 immunostimulatory adjuvant. We found either no evidence of increased risk or insufficient
374 evidence among pregnant women following Tdap, including for preterm labor and stillbirth
375 (moderate SoE).

376 For children, across all studies SoE was high for no increased risk of autism following
377 measles, mumps, and rubella (MMR) vaccine. SoE was high for increased risk of febrile seizures
378 with MMR. There was no evidence of increased risk (varied SoE) or insufficient evidence for
379 key adverse events for the newer vaccines such as 9-valent human papillomavirus and
380 meningococcal B vaccines. We found high SoE for no increased risk of diabetes following
381 rotavirus vaccine, and moderate SoE for no increased risk of other adverse events, such as
382 autoimmune disease and idiopathic thrombocytopenic purpura. We also found no evidence of
383 increased risk of intussusception following rotavirus vaccine at the latest time of follow-up
384 across studies that could be pooled, consistent with a recent meta-analysis [551]. However, there
385 were mixed findings across other studies, which included pre-post studies, cohort studies, and
386 self-controlled case series, particularly related to the risk following the first dose. While
387 intussusception is a known possible side effect of rotavirus vaccination (listed in the package

388 inserts for both vaccines and also in the Vaccine Injury Table as a condition covered under the
389 National Vaccine Injury Compensation Program) [552] the finding that there is no increased risk
390 with the longest-term follow-up from clinical trials is noteworthy.

391 Our study had some limitations. While our literature search procedures were extensive, some
392 unpublished data may not have been identified, although we mitigated this by searching trial
393 registries. The importance of trial registries has increased dramatically since reporting of results
394 has become mandatory. Clinicaltrials.gov is set up to capture results that can be used in
395 systematic reviews and meta-analyses, including data on severe adverse events, serious adverse
396 events, and mortality. In general, the harms data in Clinicaltrials.gov have been found to be more
397 complete than in the corresponding publications,[553, 554] although we note that the database
398 tends to better capture the presence of reported adverse events than the absence of such events.

399 However, trials often have insufficient sample sizes to identify rare adverse events and may
400 not follow participants long enough to identify long-term sequelae; even in studies with generous
401 follow-up times, timing of events is not always optimally reported. Indeed, many of the harms
402 we assessed as key adverse events (e.g., acute disseminated encephalomyelitis, Guillain-Barré
403 syndrome, transverse myelitis, anaphylaxis) are quite rare and the number of studies that
404 reported on the events for a vaccine was often small. As a result, despite our extensive searches
405 for data that could be combined across studies, our confidence intervals are often wide and the
406 SoE often low or insufficient. Given the limitations of controlled trials, we included post-
407 marketing surveillance and self-controlled analyses (if they met inclusion criteria) when grading
408 the SoE. For example, in the United States the CDC's Vaccine Safety Datalink uses data obtained
409 from eight large health care organizations, enabling studies that may be particularly useful for
410 identifying safety signals and/or investigating concerns for rare serious adverse events. Such

411 innovative methodologic approaches have improved the analysis of rare adverse events,
412 particularly in the post-marketing phase.

413 We also may have missed studies due to the challenging nature of assessing harms (as
414 contrasted with assessing effectiveness); however, we screened the full text of all identified
415 vaccine intervention studies, and our search terms did not include safety terms in order not to
416 miss relevant data. Wherever possible, we used data that could be combined in meta-analyses to
417 estimate the relative risk based on all available research studies. When we could not combine
418 data in pooled estimates, we integrated findings (including from the prior 2014 report) in a
419 narrative synthesis to inform the SoE.

420 This review excluded studies of vaccines not currently in use in the United States and cannot
421 make evidence statements for other vaccine schedules. We also excluded non-English language
422 studies. Although we considered only vaccines approved for use in the United States, it is
423 possible relevant epidemiological studies have been published in non-English journals.

424 Careful consideration should be given to research gaps, including where the evidence was
425 insufficient to assess the potential associations between some vaccines and particular adverse
426 events and/or where confidence intervals around risk estimates were extremely wide. However,
427 when deciding whether studies are warranted, important factors to consider include the severity
428 and frequency of the adverse event being studied and the challenges of conducting sufficiently
429 powered studies when investigating rare events. Given the rare nature of some of the serious
430 adverse events of interest (e.g., anaphylaxis, immune thrombocytopenia, Guillain-Barré
431 syndrome), ongoing studies of large populations and post-marketing surveillance of vaccines
432 after FDA licensure as noted earlier are needed to identify uncommon adverse events. Future

433 vaccine research will also need to take into account the expanding landscape of new vaccines
434 and vaccine technologies, in particular the new COVID-19 vaccines [555].

435

436 **Conclusion**

437 Across a large body of research, we found few instances in which vaccines are rarely
438 associated with serious adverse events; however, potential risks for rare adverse events should be
439 weighed carefully against the protective benefits that those vaccines provide.

440

441 **Acknowledgments**

442 The authors gratefully acknowledge the following individuals for their contributions to this
443 project: Kim Wittenberg, Stephanie Chang, Christine Chang, Thomas Acciani, David Kim, Kara
444 Elam, Tammy Beckham, Mark Helfand, and Ethan Balk for helpful comments; Jeremy Miles for
445 statistical consultation; Peggy Chen, Alicia Ruelaz Maher, Kelsey O'Hollaren, Nabeel Qureshi,
446 Keller Scholl, Olamigoke Akinniranye, Tim M. Kim, Oluwafemi Jimoh, Lea Xenakis, Weilong
447 (David) Kong, Zichun Xu, and Sangita Baxi for research assistance; and Judy Bearer for
448 administrative assistance.

449 In addition, the authors would like to thank the Technical Expert Panel while acknowledging
450 that the study questions, design, methodologic approaches, and/or conclusions do not necessarily
451 represent the views of individual technical and content experts. The list of technical experts who
452 provided input for the AHRQ report are as follows: Meghan Baker, Sarah Coles, Frank
453 Destefano, Janet McElhaney, Rebecca Reindel, Thomas Yoshikawa, and Ousseny Zerbo. The
454 authors also sought additional peer review for the AHRQ report and would like to thank the
455 following individuals: Haitao Chu, Tamera Coyne-Beasley, Janet Cragan, Matthew Daley, Cody
456 Meissner, and Barbara Mulach.

457 ***Conflict of interest:*** The authors have no potential conflicts of interest to report, including
458 financial interests, activities, relationships, and affiliations.

459 ***Funding source:*** This study was supported by a contract from the Agency for Healthcare
460 Research and Quality (AHRQ), Rockville, MD (Contract No. 290-2015-00010I). The funder had
461 no role in the design, interpretation of results, nor the decision to submit this manuscript. The
462 views expressed in this article are those of the authors and do not necessarily represent the views
463 of the US government; AHRQ; or the US Department of Health and Human Services.

464 ***Author Contributions:*** All authors attest that they meet the ICMJE criteria for authorship.

465

466

467 **References**

- 468 [1] Centers for Disease Control Prevention. Ten great public health achievements--United States,
469 1900-1999. *MMWR Morb Mortal Wkly Rep.* 1999;48:241-3.
- 470 [2] Gidengil C, Chen C, Parker AM, Nowak S, Matthews L. Beliefs around childhood vaccines
471 in the United States: A systematic review. *Vaccine.* 2019;37:6793-802.
- 472 [3] Kempe A, Saville AW, Albertin C, Zimet G, Breck A, Helmkamp L, et al. Parental Hesitancy
473 About Routine Childhood and Influenza Vaccinations: A National Survey. *Pediatrics.* 2020;146.
- 474 [4] Szilagyi PG, Albertin CS, Gurfinkel D, Saville AW, Vangala S, Rice JD, et al. Prevalence
475 and characteristics of HPV vaccine hesitancy among parents of adolescents across the US.
476 *Vaccine.* 2020;38:6027-37.
- 477 [5] Centers for Disease Control and Prevention. Vaccination Coverage Among Adults in the
478 United States, National Health Interview Survey, 2017. 2018.
- 479 [6] Kahn KE, Black CL, Ding H, Williams WW, Lu PJ, Fiebelkorn AP, et al. Influenza and Tdap
480 Vaccination Coverage Among Pregnant Women - United States, April 2018. *MMWR Morb*
481 *Mortal Wkly Rep.* 2018;67:1055-9.
- 482 [7] Lindley MC, Kahn KE, Bardenheier BH, D'Angelo DV, Dawood FS, Fink RV, et al. Vital
483 Signs: Burden and Prevention of Influenza and Pertussis Among Pregnant Women and Infants -
484 United States. *MMWR Morb Mortal Wkly Rep.* 2019;68:885-92.
- 485 [8] Marshall V, Baylor NW. Food and Drug Administration regulation and evaluation of
486 vaccines. *Pediatrics.* 2011;127 Suppl 1:S23-30.
- 487 [9] Salmon DA, Pavia A, Gellin B. Editors' introduction: Vaccine safety throughout the product
488 life cycle. *Pediatrics.* 2011;127 Suppl 1:S1-4.
- 489 [10] U.S Food and Drug Administration. Postmarketing Requirements and Commitments:
490 Introduction. 2016.
- 491 [11] Nguyen M, Ball R, Midthun K, Lieu TA. The Food and Drug Administration's Post-
492 Licensure Rapid Immunization Safety Monitoring program: strengthening the federal vaccine
493 safety enterprise. *Pharmacoepidemiol Drug Saf.* 2012;21 Suppl 1:291-7.
- 494 [12] Mini-Sentinel. 2011.
- 495 [13] U.S Food and Drug Administration. Vaccines. 2020.
- 496 [14] Sentinel. Vaccines, Blood, & Biologics Assessments.
- 497 [15] Centers for Disease Control and Prevention. Vaccine Safety Datalink (VSD). 2019.
- 498 [16] DeStefano F, Vaccine Safety Datalink Research G. The Vaccine Safety Datalink project.
499 *Pharmacoepidemiol Drug Saf.* 2001;10:403-6.
- 500 [17] Centers for Disease Control and Prevention. Clinical Immunization Safety Assessment
501 (CISA) Project. 2020.
- 502 [18] Williams SE, Klein NP, Halsey N, Dekker CL, Baxter RP, Marchant CD, et al. Overview of
503 the Clinical Consult Case Review of adverse events following immunization: Clinical
504 Immunization Safety Assessment (CISA) network 2004-2009. *Vaccine.* 2011;29:6920-7.
- 505 [19] Agency for Healthcare Research and Quality. Methods Guide for Effectiveness and
506 Comparative Effectiveness Reviews. Rockville (MD)2008.
- 507 [20] U.S Food and Drug Administration. Vaccines Licensed for Use in the United States.
- 508 [21] Centers for Disease Control and Prevention. Immunization Schedules: Table 1.
509 Recommended Adult Immunization Schedule for ages 19 years or older, United States, 2021.

510 [22] Centers for Disease Control and Prevention. Immunization Schedules: Table 1.
511 Recommended Child and Adolescent Immunization Schedule for ages 18 years or younger,
512 United States, 2021.

513 [23] Maglione MA, Gidengil C, Das L, Raaen L, Smith A, Chari R, et al. Safety of Vaccines
514 Used for Routine Immunization in the United States. *Evid Rep Technol Assess (Full Rep)*.
515 2014;1-740.

516 [24] Institute of Medicine I. Adverse effects of vaccines: Evidence and causality. Washington,
517 DC: The National Academy Press. 2011.

518 [25] Motala A, Hempel S, Gidengil C, Goetz M, Maglione M, Hall O, et al. Safety of Vaccines
519 Used for Routine Immunization in the United States: An update. PROSPERO 2020
520 CRD42020180089 2020.

521 [26] Agency for Healthcare Research and Quality. Safety of Vaccines Used for Routine
522 Immunization in the United States: Research Protocol. April 6, 2020.

523 [27] Higgins JPT, Green S, (editors). *Cochrane Handbook for Systematic Reviews of*
524 *Interventions* Version 5.1.0 [updated March 2011]. The Cochrane Collaboration, 2011. Available
525 from www.cochrane-handbook.org.

526 [28] IntHout J, Ioannidis JP, Borm GF. The Hartung-Knapp-Sidik-Jonkman method for random
527 effects meta-analysis is straightforward and considerably outperforms the standard DerSimonian-
528 Laird method. *BMC Med Res Methodol*. 2014;14:25.

529 [29] Jackson D, Law M, Rucker G, Schwarzer G. The Hartung-Knapp modification for random-
530 effects meta-analysis: A useful refinement but are there any residual concerns? *Stat Med*.
531 2017;36:3923-34.

532 [30] Santaguida P, Raina P. The Development of the McHarm Quality Assessment Scale for
533 adverse events: Delphi Consensus on important criteria for evaluating harms. McMaster
534 University. 2012.

535 [31] *Methods Guide for Effectiveness and Comparative Effectiveness Reviews*. AHRQ
536 Publication No. 10(14)-EHC063-EF. 2011/03/25 ed. Rockville, MD: Agency for Healthcare
537 Research and Quality; January 2014.

538 [32] Abdelnour A, Silas PE, Lamas MRV, Aragon CFG, Chiu N-C, Chiu C-H, et al. Safety of a
539 quadrivalent meningococcal serogroups A, C, W and Y conjugate vaccine (MenACWY-CRM)
540 administered with routine infant vaccinations: results of an open-label, randomized, phase 3b
541 controlled study in healthy infants. *Vaccine*. 2014;32:965-72.

542 [33] Acosta J, Benages C, Díaz MA, Xiberta M, Muñoz F. Preventing pertussis in the early
543 infant: Development and results of a prenatal vaccination program. *Acta Medica International*.
544 2016;3:78-81.

545 [34] Alberer M, Burchard G, Jelinek T, Reisinger E, Beran J, Hlavata LC, et al. Safety and
546 immunogenicity of typhoid fever and yellow fever vaccines when administered concomitantly
547 with quadrivalent meningococcal ACWY glycoconjugate vaccine in healthy adults. *Journal of*
548 *travel medicine*. 2015;22:48-56.

549 [35] Alberer M, Burchard G, Jelinek T, Reisinger EC, Meyer S, Forleo-Neto E, et al.
550 Immunogenicity and safety of concomitant administration of a combined hepatitis A/B vaccine
551 and a quadrivalent meningococcal conjugate vaccine in healthy adults. *Journal of travel*
552 *medicine*. 2015;22:105-14.

553 [36] Amdekar YK, Lalwani SK, Bavdekar A, Balasubramanian S, Chhatwal J, Bhat SR, et al.
554 Immunogenicity and safety of a 13-valent pneumococcal conjugate vaccine in healthy infants

555 and toddlers given with routine vaccines in India. *The Pediatric infectious disease journal.*
556 2013;32:509-16.

557 [37] Anez G, Hedrick J, Simon MW, Christensen S, Jeanfreau R, Yau E, et al. Immunogenicity
558 and safety of a booster dose of a quadrivalent meningococcal tetanus toxoid-conjugate vaccine
559 (MenACYW-TT) in adolescents and adults: a Phase III randomized study. *Hum Vaccin*
560 *Immunother.* 2020:1-7.

561 [38] Baccarini CI, Simon MW, Brandon D, Christensen S, Jordanov E, Dhingra MS. Safety and
562 Immunogenicity of a Quadrivalent Meningococcal Conjugate Vaccine in Healthy
563 Meningococcal-Naïve Children 2-9 Years of Age: A Phase III, Randomized Study. *Pediatric*
564 *Infectious Disease Journal.* 2020:955-60.

565 [39] Baker MA, Baer B, Kulldorff M, Zichittella L, Reindel R, DeLuccia S, et al. Kawasaki
566 disease and 13-valent pneumococcal conjugate vaccination among young children: A self-
567 controlled risk interval and cohort study with null results. *PLoS medicine.* 2019;16:e1002844.

568 [40] Baker MA, Jankosky C, Yih WK, Gruber S, Li L, Cocoros NM, et al. The risk of febrile
569 seizures following influenza and 13-valent pneumococcal conjugate vaccines. *Vaccine.*
570 2020;38:2166-71.

571 [41] Bart S, Cannon K, Herrington D, Mills R, Forleo-Neto E, Lindert K, et al. Immunogenicity
572 and safety of a cell culture-based quadrivalent influenza vaccine in adults: A Phase III, double-
573 blind, multicenter, randomized, non-inferiority study. *Human vaccines & immunotherapeutics.*
574 2016;12:2278-88.

575 [42] Baxter R, Eaton A, Hansen J, Aukes L, Caspard H, Ambrose CS. Safety of quadrivalent live
576 attenuated influenza vaccine in subjects aged 2-49years. *Vaccine.* 2017;35:1254-8.

577 [43] Baxter R, Lewis E, Goddard K, Fireman B, Bakshi N, DeStefano F, et al. Acute
578 Demyelinating Events Following Vaccines: A Case-Centered Analysis. *Clin Infect Dis.*
579 2016;63:1456-62.

580 [44] Baxter R, Lewis N, Bohrer P, Harrington T, Aukes L, Klein NP. Sudden-Onset
581 Sensorineural Hearing Loss after Immunization: A Case-Centered Analysis. *Otolaryngol Head*
582 *Neck Surg.* 2016;155:81-6.

583 [45] Becerra-Culqui TA, Getahun D, Chiu V, Sy LS, Tseng HF. Prenatal Tetanus, Diphtheria,
584 Acellular Pertussis Vaccination and Autism Spectrum Disorder. *Pediatrics.* 2018;142.

585 [46] Becerra-Culqui TA, Getahun D, Chiu V, Sy LS, Tseng HF. The Association of Prenatal
586 Tetanus, Diphtheria, and Acellular Pertussis (Tdap) Vaccination With Attention-
587 Deficit/Hyperactivity Disorder. *American journal of epidemiology.* 2020;189:1163-72.

588 [47] Beran J, Peeters M, Dewe W, Raupachova J, Hobzova L, Devaster JM. Immunogenicity and
589 safety of quadrivalent versus trivalent inactivated influenza vaccine: a randomized, controlled
590 trial in adults. *BMC Infectious Diseases.* 2013;13:224.

591 [48] Berenson AB, Hirth JM, Rahman M, Laz TH, Rupp RE, Sarpong KO. Maternal and infant
592 outcomes among women vaccinated against pertussis during pregnancy. *Human vaccines &*
593 *immunotherapeutics.* 2016;12:1965-71.

594 [49] Black S, Klein NP, Shah J, Bedell L, Karsten A, Dull PM. Immunogenicity and tolerability
595 of a quadrivalent meningococcal glycoconjugate vaccine in children 2-10 years of age. *Vaccine.*
596 2010;28:657-63.

597 [50] Block SL, Falloon J, Hirschfield JA, Krilov LR, Dubovsky F, Yi T, et al. Immunogenicity
598 and safety of a quadrivalent live attenuated influenza vaccine in children. *Pediatric Infectious*
599 *Disease Journal.* 2012;31:745-51.

600 [51] Block SL, Klein NP, Sarpong K, Russell S, Fling J, Petrecz M, et al. Lot-to-lot Consistency,
601 Safety, Tolerability and Immunogenicity of an Investigational Hexavalent Vaccine in US Infants.
602 *The Pediatric infectious disease journal*. 2017;36:202-8.

603 [52] Block SL, Yi T, Sheldon E, Dubovsky F, Falloon J. A randomized, double-blind
604 noninferiority study of quadrivalent live attenuated influenza vaccine in adults. *Vaccine*.
605 2011;29:9391-7.

606 [53] Bonten MJM, Huijts SM, Bolkenbaas M, Webber C, Patterson S, Gault S, et al.
607 Polysaccharide conjugate vaccine against pneumococcal pneumonia in adults. *The New England*
608 *journal of medicine*. 2015;372:1114-25.

609 [54] Briggs-Steinberg C, Aboudi D, Hodson G, Shah S. Clinical Tolerance of In-Neonatal
610 Intensive Care Unit Administration of Rotavirus Vaccine. *Am J Perinatol*. 2019.

611 [55] Burke RM, Tate JE, Dahl RM, Aliabadi N, Parashar UD. Does Rotavirus Vaccination
612 Affect Longer-Term Intussusception Risk in US Infants? *J Pediatric Infect Dis Soc*. 2020;9:257-
613 60.

614 [56] Carlin JB, Macartney KK, Lee KJ, Quinn HE, Buttery J, Lopert R, et al. Intussusception
615 risk and disease prevention associated with rotavirus vaccines in Australia's National
616 Immunization Program. *Clinical infectious diseases : an official publication of the Infectious*
617 *Diseases Society of America*. 2013;57:1427-34.

618 [57] Caspard H, Steffey A, Mallory RM, Ambrose CS. Evaluation of the safety of live attenuated
619 influenza vaccine (LAIV) in children and adolescents with asthma and high-risk conditions: a
620 population-based prospective cohort study conducted in England with the Clinical Practice
621 Research Datalink. *BMJ Open*. 2018;8:e023118.

622 [58] Chang L-J, Meng Y, Janoszyk H, Landolfi V, Talbot HK, Group QHDS. Safety and
623 immunogenicity of high-dose quadrivalent influenza vaccine in adults ≥ 65 years of age: A
624 phase 3 randomized clinical trial. *Vaccine*. 2019;37:5825-34.

625 [59] Chang LJ, Hedrick J, Christensen S, Pan J, Jordanov E, Dhingra MS. A Phase II,
626 randomized, immunogenicity and safety study of a quadrivalent meningococcal conjugate
627 vaccine, MenACYW-TT, in healthy adolescents in the United States. *Vaccine*. 2020a.

628 [60] Chang LJ, Hedrick J, Christensen S, Pan J, Jordanov E, Dhingra MS. A Phase II,
629 randomized, immunogenicity and safety study of a quadrivalent meningococcal conjugate
630 vaccine, MenACYW-TT, in healthy adolescents in the United States. *Vaccine*. 2020b.

631 [61] Chang YC, Chou YJ, Liu JY, Yeh TF, Huang N. Additive benefits of pneumococcal and
632 influenza vaccines among elderly persons aged 75 years or older in Taiwan - A representative
633 population-based comparative study. *J Infect*. 2012;65:231-8.

634 [62] Chen LF, Chen HP, Huang YS, Huang KY, Chou P, Lee CC. Pneumococcal pneumonia and
635 the risk of stroke: a population-based follow-up study. *PLoS One*. 2012;7:e51452.

636 [63] Chlibek R, Bayas JM, Collins H, de la Pinta MLR, Ledent E, Mols JF, et al. Safety and
637 immunogenicity of an AS01-adjuvanted varicella-zoster virus subunit candidate vaccine against
638 herpes zoster in adults ≥ 50 years of age. *The Journal of infectious diseases*. 2013;208:1953-61.

639 [64] Contopoulos-Ioannidis DG, Halpern MS, Maldonado Y. Trends in Hospitalizations for
640 Intussusception in California in Relationship to the Introduction of New Rotavirus Vaccines,
641 1985-2010. *The Pediatric infectious disease journal*. 2015;34:712-7.

642 [65] Cowling BJ, Perera RAPM, Valkenburg SA, Leung NHL, Iuliano AD, Tam YH, et al.
643 Comparative Immunogenicity of Several Enhanced Influenza Vaccine Options for Older Adults:
644 A Randomized, Controlled Trial. *Clinical infectious diseases : an official publication of the*
645 *Infectious Diseases Society of America*. 2019.

646 [66] Cunningham AL, Lal H, Kovac M, Chlibek R, Hwang S-J, Diez-Domingo J, et al. Efficacy
647 of the Herpes Zoster Subunit Vaccine in Adults 70 Years of Age or Older. *The New England*
648 *journal of medicine*. 2016;375:1019-32.

649 [67] Cutland CL, Nolan T, Halperin SA, Kurugol Z, Ahmed K, Perrett KP, et al.
650 Immunogenicity and safety of one or two doses of the quadrivalent meningococcal vaccine
651 MenACWY-TT given alone or with the 13-valent pneumococcal conjugate vaccine in toddlers:
652 A phase III, open-label, randomised study. *Vaccine*. 2018;36:1908-16.

653 [68] Dagan R, Patterson S, Juergens C, Greenberg D, Givon-Lavi N, Porat N, et al. Comparative
654 immunogenicity and efficacy of 13-valent and 7-valent pneumococcal conjugate vaccines in
655 reducing nasopharyngeal colonization: a randomized double-blind trial. *Clinical infectious*
656 *diseases : an official publication of the Infectious Diseases Society of America*. 2013;57:952-62.

657 [69] Daley MF, Yih WK, Glanz JM, Hambidge SJ, Narwaney KJ, Yin R, et al. Safety of
658 diphtheria, tetanus, acellular pertussis and inactivated poliovirus (DTaP-IPV) vaccine. *Vaccine*.
659 2014;32:3019-24.

660 [70] Deichmann KA, Ferrera G, Tran C, Thomas S, Eymin C, Baudin M. Immunogenicity and
661 safety of a combined measles, mumps, rubella and varicella live vaccine (ProQuad)
662 administered concomitantly with a booster dose of a hexavalent vaccine in 12-23-month-old
663 infants. *Vaccine*. 2015;33:2379-86.

664 [71] DeMeo SD, Raman SR, Hornik CP, Wilson CC, Clark R, Smith PB. Adverse Events After
665 Routine Immunization of Extremely Low-Birth-Weight Infants. *JAMA Pediatr*. 2015;169:740-5.

666 [72] DeSilva M, Vazquez-Benitez G, Nordin JD, Lipkind HS, Klein NP, Cheetham TC, et al.
667 Maternal Tdap vaccination and risk of infant morbidity. *Vaccine*. 2017;35:3655-60.

668 [73] Dhingra MS, Kundu R, Gupta M, Kanungo S, Ganguly N, Singh MP, et al. Evaluation of
669 safety and immunogenicity of a live attenuated tetravalent (G1-G4) Bovine-Human Reassortant
670 Rotavirus vaccine (BRV-TV) in healthy Indian adults and infants. *Vaccine*. 2014;32 Suppl
671 1:A117-23.

672 [74] Dhingra MS, Peterson J, Hedrick J, Pan J, Neveu D, Jordanov E. Immunogenicity, safety
673 and inter-lot consistency of a meningococcal conjugate vaccine (MenACYW-TT) in adolescents
674 and adults: A Phase III randomized study. *Vaccine*. 2020;38:5194-201.

675 [75] Domachowske JB, Pankow-Culot H, Bautista M, Feng Y, Claeys C, Peeters M, et al. A
676 Randomized Trial of Candidate Inactivated Quadrivalent Influenza Vaccine versus Trivalent
677 Influenza Vaccines in Children Aged 3-17 Years. *Journal of Infectious Diseases*. 2013;207:1878-
678 87.

679 [76] Donahue JG, Kieke BA, Lewis EM, Weintraub ES, Hanson KE, McClure DL, et al. Near
680 real-time surveillance to assess the safety of the 9-valent human papillomavirus vaccine.
681 *Pediatrics*. 2019;144.

682 [77] Duffy J, Weintraub E, Hambidge SJ, Jackson LA, Kharbanda EO, Klein NP, et al. Febrile
683 Seizure Risk After Vaccination in Children 6 to 23 Months. *Pediatrics*. 2016;138:1-10.

684 [78] Dunkle LM, Izikson R, Patriarca P, Goldenthal KL, Muse D, Callahan J, et al. Efficacy of
685 Recombinant Influenza Vaccine in Adults 50 Years of Age or Older. *The New England journal*
686 *of medicine*. 2017;376:2427-36.

687 [79] Dunkle LM, Izikson R, Patriarca PA, Goldenthal KL, Muse D, Cox MMJ. Randomized
688 Comparison of Immunogenicity and Safety of Quadrivalent Recombinant Versus Inactivated
689 Influenza Vaccine in Healthy Adults 18-49 Years of Age. *The Journal of infectious diseases*.
690 2017;216:1219-26.

691 [80] Dynavax Technologies C. Safety and Immunogenicity Study of the Hepatitis B Vaccine,
692 HEPLISAV™, Compared to Engerix-B® Vaccine. 2015.

693 [81] Dynavax Technologies Corporation. Safety and Efficacy of HEPLISAV™ Hepatitis B
694 Virus Vaccine Compared With Engerix-B® Vaccine. 2006.

695 [82] Eriksson M, Käyhty H, Saha H, Lahdenkari M, Koskinen P, Mäkisalo H, et al. A
696 randomized, controlled trial comparing the immunogenicity and safety of a 23-valent
697 pneumococcal polysaccharide vaccination to a repeated dose 13-valent pneumococcal conjugate
698 vaccination in kidney transplant recipients. *Transplant Infectious Disease*. 2020;22.

699 [83] Escolano S, Hill C, Tubert-Bitter P. Intussusception risk after RotaTeq vaccination:
700 evaluation from worldwide spontaneous reporting data using a self-controlled case series
701 approach. *Vaccine*. 2015;33:1017-20.

702 [84] Essink B, Fierro C, Rosen J, Figueroa AL, Zhang B, Verhoeven C, et al. Immunogenicity
703 and safety of MF59-adjuvanted quadrivalent influenza vaccine versus standard and alternate B
704 strain MF59-adjuvanted trivalent influenza vaccines in older adults. *Vaccine*. 2020;38:242-50.

705 [85] Esteves-Jaramillo A, Koehler T, Jeanfreau R, Neveu D, Jordanov E, Singh Dhingra M.
706 Immunogenicity and safety of a quadrivalent meningococcal tetanus toxoid-conjugate vaccine
707 (MenACYW-TT) in ≥ 56 -year-olds: A Phase III randomized study. *Vaccine*. 2020;38:4405-11.

708 [86] Eucr Outside EU/EEA. Safety and Immunogenicity of Meningococcal ACWY Conjugate
709 Versus Polysaccharide Vaccine in Children 2 to 10 Years of Age. 2014.

710 [87] Fernandes EG, Leshem E, Patel M, Flannery B, Pellini ACG, Veras MA, et al. Hospital-
711 based surveillance of intussusception among infants. *Jornal de pediatria*. 2016;92:181-7.

712 [88] Fischer L, Gerstel PF, Poncet A, Siegrist C-A, Laffitte E, Gabay C, et al. Pneumococcal
713 polysaccharide vaccination in adults undergoing immunosuppressive treatment for inflammatory
714 diseases--a longitudinal study. *Arthritis Res Ther*. 2015;17:151.

715 [89] Fotso Kamdem A, Vidal C, Pazart L, Leroux F, Pugin A, Savet C, et al. A case-control
716 study of risk factors for intussusception among infants in eastern France after the introduction of
717 the rotavirus vaccine. *Vaccine*. 2019;37:4587-93.

718 [90] Frenck RW, Jr., Gurtman A, Rubino J, Smith W, van Cleeff M, Jayawardene D, et al.
719 Randomized, controlled trial of a 13-valent pneumococcal conjugate vaccine administered
720 concomitantly with an influenza vaccine in healthy adults. *Clin Vaccine Immunol*.
721 2012;19:1296-303.

722 [91] Frey SE, Reyes MRA-DL, Reynales H, Bernal NN, Nicolay U, Narasimhan V, et al.
723 Comparison of the safety and immunogenicity of an MF59-adjuvanted with a non-adjuvanted
724 seasonal influenza vaccine in elderly subjects. *Vaccine*. 2014;32:5027-34.

725 [92] Garland SM, Cheung T-H, McNeill S, Petersen LK, Romaguera J, Vazquez-Narvaez J, et al.
726 Safety and immunogenicity of a 9-valent HPV vaccine in females 12-26 years of age who
727 previously received the quadrivalent HPV vaccine. *Vaccine*. 2015;33:6855-64.

728 [93] Gasparini R, Johnston W, Conversano M, Garscadden A, Alexanderian D, Giglioli N, et al.
729 Immunogenicity and safety of combined tetanus, reduced diphtheria, acellular pertussis vaccine
730 when co-administered with quadrivalent meningococcal conjugate and human papillomavirus
731 vaccines in healthy adolescents. *J Vaccines Vaccin*. 2014;5:231.

732 [94] Geier DA, Geier MR. A longitudinal cohort study of childhood MMR vaccination and
733 seizure disorder among American children. *Brain and Development*. 2020.

734 [95] Gilca V, Sauvageau C, Panicker G, De Serres G, Ouakki M, Unger ER. Immunogenicity
735 and safety of a mixed vaccination schedule with one dose of nonavalent and one dose of bivalent

736 HPV vaccine versus two doses of nonavalent vaccine - A randomized clinical trial. *Vaccine*.
737 2018;36:7017-24.

738 [96] Glanz JM, Clarke CL, Xu S, Daley MF, Shoup JA, Schroeder EB, et al. Association
739 between Rotavirus Vaccination and Type 1 Diabetes in Children. *JAMA Pediatrics*. 2020:E1-E8.

740 [97] Glover C, Crawford N, Leeb A, Wood N, Macartney K. Active SMS-based surveillance of
741 adverse events following immunisation with influenza and pertussis-containing vaccines in
742 Australian pregnant women using AusVaxSafety. *Vaccine*. 2020a;38:4892-900.

743 [98] Greenberg DP, Robertson CA, Landolfi VA, Bhaumik A, Senders SD, Decker MD. Safety
744 and immunogenicity of an inactivated quadrivalent influenza vaccine in children 6 months
745 through 8 years of age. *The Pediatric infectious disease journal*. 2014;33:630-6.

746 [99] Greenberg DP, Robertson CA, Noss MJ, Blatter MM, Biedenbender R, Decker MD. Safety
747 and immunogenicity of a quadrivalent inactivated influenza vaccine compared to licensed
748 trivalent inactivated influenza vaccines in adults. *Vaccine*. 2013;31:770-6.

749 [100] Greenberg DP, Robertson CA, Talbot HK, Decker MD. Safety and immunogenicity of a
750 quadrivalent influenza vaccine in adults 65 y of age and older. *Human vaccines &*
751 *immunotherapeutics*. 2017;13:2058-64.

752 [101] Greenberg RN, Gurtman A, Frenck RW, Strout C, Jansen KU, Trammel J, et al. Sequential
753 administration of 13-valent pneumococcal conjugate vaccine and 23-valent pneumococcal
754 polysaccharide vaccine in pneumococcal vaccine-naive adults 60-64 years of age. *Vaccine*.
755 2014;32:2364-74.

756 [102] Griffin JB, Yu L, Watson D, Turner N, Walls T, Howe AS, et al. Pertussis Immunisation in
757 Pregnancy Safety (PIPS) Study: A retrospective cohort study of safety outcomes in pregnant
758 women vaccinated with Tdap vaccine. *Vaccine*. 2018;36:5173-9.

759 [103] Groome MJ, Tate JE, Arnold M, Chitnis M, Cox S, de Vos C, et al. Evaluation of
760 intussusception after oral monovalent rotavirus vaccination in South Africa. *Clinical infectious*
761 *diseases : an official publication of the Infectious Diseases Society of America*. 2019.

762 [104] Hall C, Abramovitz LM, Bukowinski AT, Ricker AA, Khodr ZG, Gumbs GR, et al. Safety
763 of tetanus, diphtheria, and acellular pertussis vaccination among pregnant active duty U.S.
764 military women. *Vaccine*. 2020;38:1982-8.

765 [105] Halperin SA, Donovan C, Marshall GS, Pool V, Decker MD, Johnson DR, et al.
766 Randomized Controlled Trial of the Safety and Immunogenicity of Revaccination With Tetanus-
767 Diphtheria-Acellular Pertussis Vaccine (Tdap) in Adults 10 Years After a Previous Dose. *Journal*
768 *of the Pediatric Infectious Diseases Society*. 2019;8:105-14.

769 [106] Halperin SA, Langley JM, Ye L, MacKinnon-Cameron D, Elsherif M, Allen VM, et al. A
770 Randomized Controlled Trial of the Safety and Immunogenicity of Tetanus, Diphtheria, and
771 Acellular Pertussis Vaccine Immunization During Pregnancy and Subsequent Infant Immune
772 Response. *Clinical infectious diseases : an official publication of the Infectious Diseases Society*
773 *of America*. 2018;67:1063-71.

774 [107] Hansen J, Timbol J, Lewis N, Pool V, Decker MD, Greenberg DP, et al. Safety of DTaP-
775 IPV/Hib vaccine administered routinely to infants and toddlers. *Vaccine*. 2016;34:4172-9.

776 [108] Hansen J, Zhang L, Eaton A, Baxter R, Robertson CA, Decker MD, et al. Post-licensure
777 safety surveillance study of routine use of quadrivalent meningococcal diphtheria toxoid
778 conjugate vaccine (MenACWY-D) in infants and children. *Vaccine*. 2018;36:2133-8.

779 [109] Hansen J, Zhang L, Klein NP, Robertson CA, Decker MD, Greenberg DP, et al. Post-
780 licensure safety surveillance study of routine use of quadrivalent meningococcal diphtheria
781 toxoid conjugate vaccine. *Vaccine*. 2017;35:6879-84.

782 [110] Hartvickson R, Cruz M, Ervin J, Brandon D, Forleo-Neto E, Dagnew AF, et al. Non-
783 inferiority of mammalian cell-derived quadrivalent subunit influenza virus vaccines compared to
784 trivalent subunit influenza virus vaccines in healthy children: a phase III randomized,
785 multicenter, double-blind clinical trial. *International journal of infectious diseases : IJID : official*
786 *publication of the International Society for Infectious Diseases.* 2015;41:65-72.

787 [111] Hattori F, Kawamura Y, Kawada JI, Kojima S, Natsume J, Ito K, et al. Survey of rotavirus-
788 associated severe complications in Aichi Prefecture. *Pediatr Int.* 2018;60:259-63.

789 [112] Hawken S, Ducharme R, Rosella LC, Benchimol EI, Langley JM, Wilson K, et al.
790 Assessing the risk of intussusception and rotavirus vaccine safety in Canada. *Human vaccines &*
791 *immunotherapeutics.* 2017;13:703-10.

792 [113] Heyward WL, Kyle M, Blumenau J, Davis M, Reisinger K, Kabongo ML, et al.
793 Immunogenicity and safety of an investigational hepatitis B vaccine with a Toll-like receptor 9
794 agonist adjuvant (HBsAg-1018) compared to a licensed hepatitis B vaccine in healthy adults 40-
795 70 years of age. *Vaccine.* 2013;31:5300-5.

796 [114] Hoffman V, Abu-Elyazeed R, Enger C, Esposito DB, Doherty MC, Quinlan SC, et al.
797 Safety study of live, oral human rotavirus vaccine: A cohort study in United States health
798 insurance plans. *Human vaccines & immunotherapeutics.* 2018;14:1782-90.

799 [115] Huang J, Ou HY, Lin J, Karnchanasorn R, Feng W, Samoa R, et al. The impact of hepatitis
800 B vaccination status on the risk of diabetes, implicating diabetes risk reduction by successful
801 vaccination. *PLoS ONE.* 2015;10.

802 [116] Huang W-T, Juan Y-C, Liu C-H, Yang Y-Y, Chan KA. Intussusception and Kawasaki
803 disease after rotavirus vaccination in Taiwanese infants. *Vaccine.* 2020;38:6299-303.

804 [117] Huh WK, Joura EA, Giuliano AR, Iversen O-E, de Andrade RP, Ault KA, et al. Final
805 efficacy, immunogenicity, and safety analyses of a nine-valent human papillomavirus vaccine in
806 women aged 16-26 years: a randomised, double-blind trial. *Lancet (London, England).*
807 2017;390:2143-59.

808 [118] Hung IFN, Leung AYM, Chu DWS, Leung D, Cheung T, Chan C-K, et al. Prevention of
809 acute myocardial infarction and stroke among elderly persons by dual pneumococcal and
810 influenza vaccination: a prospective cohort study. *Clinical Infectious Diseases.* 2010;51:1007-16.

811 [119] Hviid A, Hansen JV, Frisch M, Melbye M. Measles, Mumps, Rubella Vaccination and
812 Autism: A Nationwide Cohort Study. *Annals of internal medicine.* 2019;170:513-20.

813 [120] Iwata S, Nakata S, Ukae S, Koizumi Y, Morita Y, Kuroki H, et al. Efficacy and safety of
814 pentavalent rotavirus vaccine in Japan: a randomized, double-blind, placebo-controlled,
815 multicenter trial. *Human vaccines & immunotherapeutics.* 2013;9:1626-33.

816 [121] Jackson LA, Gurtman A, van Cleeff M, Jansen KU, Jayawardene D, Devlin C, et al.
817 Immunogenicity and safety of a 13-valent pneumococcal conjugate vaccine compared to a 23-
818 valent pneumococcal polysaccharide vaccine in pneumococcal vaccine-naïve adults. *Vaccine.*
819 2013;31:3577-84.

820 [122] Jain A, Marshall J, Buikema A, Bancroft T, Kelly JP, Newschaffer CJ. Autism occurrence
821 by MMR vaccine status among US children with older siblings with and without autism. *JAMA.*
822 2015;313:1534-40.

823 [123] Juergens C, de Villiers PJT, Moodley K, Jayawardene D, Jansen KU, Scott DA, et al.
824 Safety and immunogenicity of 13-valent pneumococcal conjugate vaccine formulations with and
825 without aluminum phosphate and comparison of the formulation of choice with 23-valent
826 pneumococcal polysaccharide vaccine in elderly adults: a randomized open-label trial. *Human*
827 *vaccines & immunotherapeutics.* 2014;10:1343-53.

828 [124] Kantso B, Halkjaer SI, Thomsen OO, Belard E, Gottschalck IB, Jorgensen CS, et al.
829 Immunosuppressive drugs impairs antibody response of the polysaccharide and conjugated
830 pneumococcal vaccines in patients with Crohn's disease. *Vaccine*. 2015;33:5464-9.

831 [125] Kharbanda EO, Vazquez-Benitez G, Lipkind HS, Klein NP, Cheetham TC, Naleway AL,
832 et al. Maternal Tdap vaccination: Coverage and acute safety outcomes in the vaccine safety
833 datalink, 2007-2013. *Vaccine*. 2016;34:968-73.

834 [126] Kieninger D, Sheldon E, Lin W-Y, Yu C-J, Bayas JM, Gabor JJ, et al. Immunogenicity,
835 reactogenicity and safety of an inactivated quadrivalent influenza vaccine candidate versus
836 inactivated trivalent influenza vaccine: a phase III, randomized trial in adults aged ≥ 18 years.
837 *BMC infectious diseases*. 2013;13:343.

838 [127] Kim DS, Shin SH, Lee HJ, Hong YJ, Lee SY, Choi KM, et al. Immunogenicity and safety
839 of 13-valent pneumococcal conjugate vaccine given to Korean children receiving routine
840 pediatric vaccines. *Pediatric Infectious Disease Journal*. 2013;32:266-73.

841 [128] Kirstein J, Pina M, Pan J, Jordanov E, Dhingra MS. Immunogenicity and safety of a
842 quadrivalent meningococcal tetanus toxoid-conjugate vaccine (MenACYW-TT) in adults 56
843 years of age and older: a Phase II randomized study. *Human Vaccines and Immunotherapeutics*.
844 2020;16:1299-305.

845 [129] Klein NP, Fireman B, Yih WK, Lewis E, Kulldorff M, Ray P, et al. Measles-mumps-
846 rubella-varicella combination vaccine and the risk of febrile seizures. *Pediatrics*. 2010;126:e1-8.

847 [130] Klein NP, Lewis E, Fireman B, Hambidge SJ, Naleway A, Nelson JC, et al. Safety of
848 measles-containing vaccines in 1-year-old children. *Pediatrics*. 2015;135:e321-9.

849 [131] Lal H, Cunningham AL, Godeaux O, Chlibek R, Diez-Domingo J, Hwang S-J, et al.
850 Efficacy of an adjuvanted herpes zoster subunit vaccine in older adults. *The New England
851 journal of medicine*. 2015;372:2087-96.

852 [132] Langer-Gould A, Qian L, Tartof SY, Brara SM, Jacobsen SJ, Beaber BE, et al. Vaccines
853 and the risk of multiple sclerosis and other central nervous system demyelinating diseases.
854 *JAMA neurology*. 2014;71:1506-13.

855 [133] Langley JM, Carmona Martinez A, Chatterjee A, Halperin SA, McNeil S, Reisinger KS, et
856 al. Immunogenicity and safety of an inactivated quadrivalent influenza vaccine candidate: a
857 phase III randomized controlled trial in children. *The Journal of infectious diseases*.
858 2013;208:544-53.

859 [134] Langley JM, Wang L, Aggarwal N, Bueso A, Chandrasekaran V, Cousin L, et al.
860 Immunogenicity and Reactogenicity of an Inactivated Quadrivalent Influenza Vaccine
861 Administered Intramuscularly to Children 6 to 35 Months of Age in 2012-2013: A Randomized,
862 Double-Blind, Controlled, Multicenter, Multicountry, Clinical Trial. *Journal of the Pediatric
863 Infectious Diseases Society*. 2015;4:242-51.

864 [135] Layton JB, Butler AM, Li D, Boggess KA, Weber DJ, McGrath LJ, et al. Prenatal Tdap
865 immunization and risk of maternal and newborn adverse events. *Vaccine*. 2017;35:4072-8.

866 [136] Layton JB, Butler AM, Panozzo CA, Brookhart MA. Rotavirus vaccination and short-term
867 risk of adverse events in US infants. *Paediatr Perinat Epidemiol*. 2018;32:448-57.

868 [137] Lee HJ, Choe YJ, Hong Y-J, Kim K-H, Park SE, Kim Y-K, et al. Immunogenicity and
869 safety of a multicomponent meningococcal serogroup B vaccine in healthy adolescents in Korea-
870 -A randomised trial. *Vaccine*. 2016;34:1180-6.

871 [138] Lee HJ, Chung M-H, Kim WJ, Hong YJ, Choi KM, Lee J, et al. Immunogenicity and
872 safety of a novel quadrivalent meningococcal conjugate vaccine (MenACWY-CRM) in healthy

873 Korean adolescents and adults. *International journal of infectious diseases : IJID : official*
874 *publication of the International Society for Infectious Diseases.* 2014;28:204-10.

875 [139] Leslie DL, Kobre RA, Richmand BJ, Aktan Guloksuz S, Leckman JF. Temporal
876 Association of Certain Neuropsychiatric Disorders Following Vaccination of Children and
877 Adolescents: A Pilot Case-Control Study. *Front Psychiatry.* 2017;8:3.

878 [140] Li C-Y, Chen L-C, Lin H-Y, Lee M-S, Hung S-K, Lai C-L, et al. Impact of 23-valent
879 pneumococcal polysaccharide vaccination on the frequency of pneumonia-related hospitalization
880 and survival in elderly patients with prostate cancer: A seven-year nationwide matched cohort
881 study. *Cancer.* 2020.

882 [141] Li R, Stewart B, McNeil MM, Duffy J, Nelson J, Kawai AT, et al. Post licensure
883 surveillance of influenza vaccines in the Vaccine Safety Datalink in the 2013-2014 and 2014-
884 2015 seasons. *Pharmacoepidemiology and drug safety.* 2016;25:928-34.

885 [142] Li R-c, Huang T, Li Y, Luo D, Tao J, Fu B, et al. Human rotavirus vaccine (RIX4414)
886 efficacy in the first two years of life: a randomized, placebo-controlled trial in China. *Human*
887 *vaccines & immunotherapeutics.* 2014;10:11-8.

888 [143] Lombardi F, Belmonti S, Fabbiani M, Morandi M, Rossetti B, Tordini G, et al.
889 Immunogenicity and Safety of the 13-Valent Pneumococcal Conjugate Vaccine versus the 23-
890 Valent Polysaccharide Vaccine in Unvaccinated HIV-Infected Adults: A Pilot, Prospective
891 Controlled Study. *PLoS one.* 2016;11:e0156523.

892 [144] Mallory RM, Nyborg A, Kalyani RN, Yuan Y, Block SL, Dubovsky F. A study to evaluate
893 the immunogenicity and shedding of live attenuated influenza vaccine strains in children 24-
894 48 months of age. *Vaccine.* 2020;38:1001-8.

895 [145] Mallory RM, Yu J, Kameo S, Tanaka M, Rito K, Itoh Y, et al. The safety and efficacy of
896 quadrivalent live attenuated influenza vaccine in Japanese children aged 2-18 years: Results of
897 two phase 3 studies. *Influenza and other respiratory viruses.* 2018a;12:438-45.

898 [146] Maréchal C, Lal H, Poder A, Ferguson M, Enweonye I, Heineman TC, et al.
899 Immunogenicity and safety of the adjuvanted recombinant zoster vaccine co-administered with
900 the 23-valent pneumococcal polysaccharide vaccine in adults ≥ 50 years of age: a randomized
901 trial. *Vaccine.* 2018;36:4278- 86.

902 [147] Marshall GS, Adams GL, Leonardi ML, Petrecz M, Flores SA, Ngai AL, et al.
903 Immunogenicity, Safety, and Tolerability of a Hexavalent Vaccine in Infants. *Pediatrics.*
904 2015;136:e323-32.

905 [148] McClure DL, Jacobsen SJ, Klein NP, Naleway AL, Kharbanda EO, Glanz JM, et al.
906 Similar relative risks of seizures following measles containing vaccination in children born
907 preterm compared to full-term without previous seizures or seizure-related disorders. *Vaccine.*
908 2019;37:76-9.

909 [149] McGeoch LJ, Finn A, Marlow RD. Impact of rotavirus vaccination on intussusception
910 hospital admissions in England. *Vaccine.* 2020;38:5618-26.

911 [150] Mo Z, Mo Y, Li M, Tao J, Yang X, Kong J, et al. Efficacy and safety of a pentavalent live
912 human-bovine reassortant rotavirus vaccine (RV5) in healthy Chinese infants: A randomized,
913 double-blind, placebo-controlled trial. *Vaccine.* 2017;35:5897-904.

914 [151] Morgan JL, Baggari SR, McIntire DD, Sheffield JS. Pregnancy outcomes after antepartum
915 tetanus, diphtheria, and acellular pertussis vaccination. *Obstetrics and gynecology.*
916 2015;125:1433-8.

917 [152] Munnoch SA, Cashman P, Peel R, Attia J, Hure A, Durrheim DN. Participant-Centered
918 Online Active Surveillance for Adverse Events Following Vaccination in a Large Clinical Trial:
919 Feasibility and Usability Study. *Journal of medical Internet research*. 2019;21:e14791.

920 [153] Munoz FM, Bond NH, Maccato M, Pinell P, Hammill HA, Swamy GK, et al. Safety and
921 immunogenicity of tetanus diphtheria and acellular pertussis (Tdap) immunization during
922 pregnancy in mothers and infants: a randomized clinical trial. *JAMA*. 2014;311:1760-9.

923 [154] Naleway AL, Mittendorf KF, Irving SA, Henninger ML, Crane B, Smith N, et al. Primary
924 Ovarian Insufficiency and Adolescent Vaccination. *Pediatrics*. 2018;142.

925 [155] Nelson JC, Yu O, Dominguez-Islas CP, Cook AJ, Peterson D, Greene SK, et al. Adapting
926 group sequential methods to observational postlicensure vaccine safety surveillance: results of a
927 pentavalent combination DTaP-IPV-Hib vaccine safety study. *American Journal of
928 Epidemiology*. 2013;177:131-41.

929 [156] Oberle D, Hoffelner M, Pavel J, Mentzer D, Barth I, Drechsel-Bauerle U, et al.
930 Retrospective multicenter matched case-control study on the risk factors for intussusception in
931 infants less than 1 year of age with a special focus on rotavirus vaccines - the German
932 Intussusception Study. *Human vaccines & immunotherapeutics*. 2020:1-14.

933 [157] Ochoa-Gondar O, Vila-Corcoles A, Rodriguez-Blanco T, de Diego-Cabanes C, Hospital-
934 Guardiola I, Jarrod-Pamies M, et al. Evaluating the clinical effectiveness of pneumococcal
935 vaccination in preventing myocardial infarction: The CAPAMIS study, three-year follow-up.
936 *Vaccine*. 2014;32:252-7.

937 [158] Ofori-Anyinam O, Leroux-Roels G, Drame M, Aerssens A, Maes C, Amanullah A, et al.
938 Immunogenicity and safety of an inactivated quadrivalent influenza vaccine co-administered
939 with a 23-valent pneumococcal polysaccharide vaccine versus separate administration, in adults
940 ≥ 50 years of age: Results from a phase III, randomized, non-inferiority trial. *Vaccine*.
941 2017;35:6321-8.

942 [159] Ostergaard L, Lucksinger GH, Absalon J, Beeslaar J, Eiden J, Jansen KU, et al. A phase 3,
943 randomized, active-controlled study to assess the safety and tolerability of meningococcal
944 serogroup B vaccine bivalent rLP2086 in healthy adolescents and young adults. *Vaccine*.
945 2016;34:1465-71.

946 [160] Perez JZV, Aranda JMR, de la O Cavazos M, Osuna MDZ, Davila JP, Elizondo MRB, et
947 al. Randomized clinical trial of the safety and immunogenicity of the Tdap vaccine in pregnant
948 Mexican women. *Human Vaccines & Immunotherapeutics*. 2017;13:128-35.

949 [161] Perez-Vilar S, Wernecke M, Arya D, Lo A-C, Lufkin B, Hu M, et al. Surveillance for
950 Guillain-Barre syndrome after influenza vaccination among U.S. Medicare beneficiaries during
951 the 2017-2018 season. *Vaccine*. 2019;37:3856-65.

952 [162] Perrett KP, Halperin SA, Nolan T, Martinez Pancorbo C, Tapiero B, Martinon-Torres F, et
953 al. Immunogenicity, transplacental transfer of pertussis antibodies and safety following pertussis
954 immunization during pregnancy: Evidence from a randomized, placebo-controlled trial. *Vaccine*.
955 2019.

956 [163] Petousis-Harris H, Jiang YN, Yu L, Watson D, Walls T, Turner N, et al. A Retrospective
957 Cohort Study of Safety Outcomes in New Zealand Infants Exposed to Tdap Vaccine in Utero.
958 *Vaccines*. 2019;7:15.

959 [164] Petrecz M, Ramsey KP, Stek JE, Martin JC, Klopfer SO, Kuter B, et al. Concomitant use
960 of VAQTA® with PedvaxHIB® and Infanrix® in 12 to 17 month old children. *Human Vaccines
961 and Immunotherapeutics*. 2016;12:503-11.

962 [165] Pfizer. Study Comparing a 13-valent Pneumococcal Conjugate Vaccine With 23-valent
963 Pneumococcal Polysaccharide Vaccine in Adults.
964 <https://ClinicalTrials.gov/show/NCT00427895>; 2007.

965 [166] Puig-Barbera J, Diez-Domingo J, Varea AB, Chavarri GS, Rodrigo JAL, Hoyos SP, et al.
966 Effectiveness of MF59-adjuvanted subunit influenza vaccine in preventing hospitalisations for
967 cardiovascular disease, cerebrovascular disease and pneumonia in the elderly. *Vaccine*.
968 2007;25:7313-21.

969 [167] Richmond PC, Marshall HS, Nissen MD, Jiang Q, Jansen KU, Garces-Sanchez M, et al.
970 Safety, immunogenicity, and tolerability of meningococcal serogroup B bivalent recombinant
971 lipoprotein 2086 vaccine in healthy adolescents: A randomised, single-blind, placebo-controlled,
972 phase 2 trial. *The Lancet Infectious Diseases*. 2012;12:597-607.

973 [168] Rivera L, Schwarz TF, Kim KH, Kim YK, Behre U, Cha SH, et al. Immunogenicity and
974 safety of the quadrivalent meningococcal vaccine MenACWY-TT co-administered with a
975 combined diphtheria-tetanus-acellular pertussis vaccine versus their separate administration in
976 adolescents and young adults: A phase III, randomized study. *Vaccine*. 2018;36:4750-8.

977 [169] Rodriguez Weber MA, Claeys C, Aranza Doniz C, Feng Y, Innis BL, Jain VK, et al.
978 Immunogenicity and safety of inactivated quadrivalent and trivalent influenza vaccines in
979 children 18-47 months of age. *The Pediatric infectious disease journal*. 2014;33:1262-9.

980 [170] Rogers MAM, Basu T, Kim C. Lower Incidence Rate of Type 1 Diabetes after Receipt of
981 the Rotavirus Vaccine in the United States, 2001–2017. *Scientific Reports*. 2019;9.

982 [171] Salas A, Pardo-Seco J, Cebey-Lopez M, Martinon-Martinez JM, Gomez-Rial J, Curras-
983 Tuala MJ, et al. Impact of rotavirus vaccination on childhood hospitalizations for seizures:
984 Heterologous or unforeseen direct vaccine effects? *Vaccine*. 2019;37:3362-8.

985 [172] Sancovski M, Mesaros N, Feng Y, Ceregido MA, Luyts D, De Barros E. Safety of reduced
986 antigen content diphtheria-tetanus-acellular pertussis vaccine when administered during
987 pregnancy as part of the maternal immunization program in Brazil: a single center, observational,
988 retrospective, cohort study. *Human vaccines & immunotherapeutics*. 2019;15:2873-81.

989 [173] Sanofi Pasteur aSC, Sanofi. Safety and Immunogenicity of Fluzone® Quadrivalent,
990 Flublok® Quadrivalent, and Fluzone® High-Dose, Influenza Vaccines, 2018-2019
991 Formulations. 2018.

992 [174] Santolaya ME, O’Ryan ML, Valenzuela MT, Prado V, Vergara R, Munoz A, et al.
993 Immunogenicity and tolerability of a multicomponent meningococcal serogroup B (4CMenB)
994 vaccine in healthy adolescents in Chile: a phase 2b/3 randomised, observer-blind, placebo-
995 controlled study. *Lancet*. 2012;379:617-24.

996 [175] Schwarz TF, Aggarwal N, Moeckesch B, Schenkenberger I, Claeys C, Douha M, et al.
997 Immunogenicity and Safety of an Adjuvanted Herpes Zoster Subunit Vaccine Coadministered
998 With Seasonal Influenza Vaccine in Adults Aged 50 Years or Older. *The Journal of infectious
999 diseases*. 2017;216:1352-61.

1000 [176] Schwarz TF, Flamaing J, Rumke HC, Penzes J, Juergens C, Wenz A, et al. A randomized,
1001 double-blind trial to evaluate immunogenicity and safety of 13-valent pneumococcal conjugate
1002 vaccine given concomitantly with trivalent influenza vaccine in adults aged ≥ 65 years. *Vaccine*.
1003 2011;29:5195-202.

1004 [177] Senders S, Bhuyan P, Jiang Q, Absalon J, Eiden JJ, Jones TR, et al. Immunogenicity,
1005 Tolerability and Safety in Adolescents of Bivalent rLP2086, a Meningococcal Serogroup B
1006 Vaccine, Coadministered with Quadrivalent Human Papilloma Virus Vaccine. *The Pediatric
1007 infectious disease journal*. 2016;35:548-54.

1008 [178] Seo YB, Choi WS, Lee J, Song JY, Cheong HJ, Kim WJ. Comparison of immunogenicity
1009 and safety of an influenza vaccine administered concomitantly with a 13-valent pneumococcal
1010 conjugate vaccine or 23-valent polysaccharide pneumococcal vaccine in the elderly. *Clinical and*
1011 *Experimental Vaccine Research*. 2017;6:38-44.

1012 [179] Shakib JH, Korgenski K, Sheng X, Varner MW, Pavia AT, Byington CL. Tetanus,
1013 diphtheria, acellular pertussis vaccine during pregnancy: pregnancy and infant health outcomes.
1014 *The Journal of pediatrics*. 2013;163:1422-4.

1015 [180] Shimada K, Morinaga H, Kiyonagi T, Miyazaki T, Nishitani-Yokoyama M, Okai I, et al.
1016 Safety and Efficacy of Simultaneous Inoculations of Pneumococcal and Influenza Vaccines in
1017 Patients with Coronary Artery Disease. *J Atheroscler Thromb*. 2020.

1018 [181] Shiramoto M, Hanada R, Juergens C, Shoji Y, Yoshida M, Ballan B, et al. Immunogenicity
1019 and safety of the 13-valent pneumococcal conjugate vaccine compared to the 23-valent
1020 pneumococcal polysaccharide vaccine in elderly Japanese adults. *Human vaccines &*
1021 *immunotherapeutics*. 2015;11:2198-206.

1022 [182] Siriwardena AN, Asghar Z, Coupland CC. Influenza and pneumococcal vaccination and
1023 risk of stroke or transient ischaemic attack-matched case control study. *Vaccine*. 2014;32:1354-
1024 61.

1025 [183] Song JY, Cheong HJ, Hyun HJ, Seo YB, Lee J, Wie S-H, et al. Immunogenicity and safety
1026 of a 13-valent pneumococcal conjugate vaccine and an MF59-adjuvanted influenza vaccine after
1027 concomitant vaccination in 60-year-old adults. *Vaccine*. 2017;35:313-20.

1028 [184] Song JY, Cheong HJ, Noh JY, Choi MJ, Yoon JG, Lee SN, et al. Immunogenicity and
1029 safety of a tetanus-diphtheria vaccine and a 13-valent pneumococcal conjugate vaccine after
1030 concomitant vaccination in \geq 50-year-old adults. *BMC infectious diseases*. 2018;18:628.

1031 [185] Song JY, Cheong HJ, Tsai TF, Chang H-A, Choi MJ, Jeon JH, et al. Immunogenicity and
1032 safety of concomitant MF59-adjuvanted influenza vaccine and 23-valent pneumococcal
1033 polysaccharide vaccine administration in older adults. *Vaccine*. 2015a;33:4647-52.

1034 [186] Song JY, Cheong HJ, Tsai TF, Chang HA, Choi MJ, Jeon JH, et al. Immunogenicity and
1035 safety of concomitant MF59-adjuvanted influenza vaccine and 23-valent pneumococcal
1036 polysaccharide vaccine administration in older adults. *Vaccine*. 2015b;33:4647-52.

1037 [187] Stockwell MS, Broder K, LaRussa P, Lewis P, Fernandez N, Sharma D, et al. Risk of fever
1038 after pediatric trivalent inactivated influenza vaccine and 13-valent pneumococcal conjugate
1039 vaccine. *JAMA pediatrics*. 2014;168:211-9.

1040 [188] Stockwell MS, Broder KR, Lewis P, Jakob K, Iqbal S, Fernandez N, et al. Assessing Fever
1041 Frequency After Pediatric Live Attenuated Versus Inactivated Influenza Vaccination. *Journal of*
1042 *the Pediatric Infectious Diseases Society*. 2017;6:e7-e14.

1043 [189] Stowe J, Andrews N, Ladhani S, Miller E. The risk of intussusception following
1044 monovalent rotavirus vaccination in England: A self-controlled case-series evaluation. *Vaccine*.
1045 2016;34:3684-9.

1046 [190] Strezova A, Lal H, Enweonye I, Campora L, Beukelaers P, Segall N, et al. The adjuvanted
1047 recombinant zoster vaccine co-administered with a tetanus, diphtheria and pertussis vaccine in
1048 adults aged \geq 50 years: A randomized trial. *Vaccine*. 2019;37:5877-85.

1049 [191] Svensson T, Kattstrom M, Hammarlund Y, Roth D, Andersson PO, Svensson M, et al.
1050 Pneumococcal conjugate vaccine triggers a better immune response than pneumococcal
1051 polysaccharide vaccine in patients with chronic lymphocytic leukemia A randomized study by
1052 the Swedish CLL group. *Vaccine*. 2018;36:3701-7.

1053 [192] Tapiero B, Halperin SA, Dionne M, Meekison W, Diaz-Mitoma F, Zickler P, et al. Safety
1054 and immunogenicity of a hexavalent vaccine administered at 2, 4 and 6 months of age with or
1055 without a heptavalent pneumococcal conjugate vaccine: a randomized, open-label study.
1056 *Pediatric Infectious Disease Journal*. 2013;32:54-61.

1057 [193] Tate JE, Mwenda JM, Armah G, Jani B, Omoro R, Ademe A, et al. Evaluation of
1058 Intussusception after Monovalent Rotavirus Vaccination in Africa. *N Engl J Med*.
1059 2018;378:1521-8.

1060 [194] Tate JE, Yen C, Steiner CA, Cortese MM, Parashar UD. Intussusception Rates Before and
1061 After the Introduction of Rotavirus Vaccine. *Pediatrics*. 2016;138.

1062 [195] Thompson AR, Klein NP, Downey HJ, Patterson S, Sundaraiyer V, Watson W, et al.
1063 Coadministration of 13-valent pneumococcal conjugate and quadrivalent inactivated influenza
1064 vaccines in adults previously immunized with polysaccharide pneumococcal vaccine 23: a
1065 randomized clinical trial. *Human vaccines & immunotherapeutics*. 2019;15:444-51.

1066 [196] Timmermann CAG, Osuna CE, Steuerwald U, Weihe P, Poulsen LK, Grandjean P.
1067 Asthma and allergy in children with and without prior measles, mumps, and rubella vaccination.
1068 *Pediatric allergy and immunology : official publication of the European Society of Pediatric
1069 Allergy and Immunology*. 2015;26:742-9.

1070 [197] Tinoco JC, Pavia-Ruz N, Cruz-Valdez A, Aranza Doniz C, Chandrasekaran V, Dewe W, et
1071 al. Immunogenicity, reactogenicity, and safety of inactivated quadrivalent influenza vaccine
1072 candidate versus inactivated trivalent influenza vaccine in healthy adults aged ≥ 18 years: a
1073 phase III, randomized trial. *Vaccine*. 2014;32:1480-7.

1074 [198] Togashi T, Okada K, Yamaji M, Thompson A, Gurtman A, Cutler M, et al.
1075 Immunogenicity and Safety of a 13-Valent Pneumococcal Conjugate Vaccine Given With DTaP
1076 Vaccine in Healthy Infants in Japan. *The Pediatric infectious disease journal*. 2015;34:1096-104.

1077 [199] Treanor JT, Albano FR, Sawlwin DC, Graves Jones A, Airey J, Formica N, et al.
1078 Immunogenicity and safety of a quadrivalent inactivated influenza vaccine compared with two
1079 trivalent inactivated influenza vaccines containing alternate B strains in adults: A phase 3,
1080 randomized noninferiority study. *Vaccine*. 2017;35:1856-64.

1081 [200] Tregnaghi M, Lopez P, Stambouljian D, Grana G, Odrlic T, Bedell L, et al.
1082 Immunogenicity and safety of a quadrivalent meningococcal polysaccharide CRM conjugate
1083 vaccine in infants and toddlers. *International journal of infectious diseases : IJID : official
1084 publication of the International Society for Infectious Diseases*. 2014;26:22-30.

1085 [201] Tseng H-F, Sy LS, Ackerson BK, Hechter RC, Tartof SY, Haag M, et al. Safety of
1086 Quadrivalent Meningococcal Conjugate Vaccine in 11- to 21-Year-Olds. *Pediatrics*. 2017;139.

1087 [202] Tseng HF, Sy LS, Liu ILA, Qian L, Marcy SM, Weintraub E, et al. Postlicensure
1088 surveillance for pre-specified adverse events following the 13-valent pneumococcal conjugate
1089 vaccine in children. *Vaccine*. 2013;31:2578-83.

1090 [203] Tseng HF, Sy LS, Qian L, Liu I-LA, Mercado C, Lewin B, et al. Pneumococcal Conjugate
1091 Vaccine Safety in Elderly Adults. *Open forum infectious diseases*. 2018;5:ofy100.

1092 [204] Uhlig U, Kostev K, Schuster V, Koletzko S, Uhlig HH. Impact of rotavirus vaccination in
1093 Germany: rotavirus surveillance, hospitalization, side effects and comparison of vaccines. *The
1094 Pediatric infectious disease journal*. 2014;33:e299-304.

1095 [205] Uno Y, Uchiyama T, Kurosawa M, Aleksic B, Ozaki N. Early exposure to the combined
1096 measles-mumps-rubella vaccine and thimerosal-containing vaccines and risk of autism spectrum
1097 disorder. *Vaccine*. 2015;33:2511-6.

1098 [206] Vaarala O, Jokinen J, Lahdenkari M, Leino T. Rotavirus Vaccination and the Risk of
1099 Celiac Disease or Type 1 Diabetes in Finnish Children at Early Life. *The Pediatric infectious*
1100 *disease journal*. 2017;36:674-5.

1101 [207] Van Damme P, Meijer CJLM, Kieninger D, Schuyleman A, Thomas S, Luxembourg A, et
1102 al. A phase III clinical study to compare the immunogenicity and safety of the 9-valent and
1103 quadrivalent HPV vaccines in men. *Vaccine*. 2016;34:4205-12.

1104 [208] Vandecasteele SJ, De Bacquer D, Caluwe R, Ombelet S, Van Vlem B. Immunogenicity
1105 and safety of the 13-valent Pneumococcal Conjugate vaccine in 23-valent pneumococcal
1106 polysaccharide vaccine-naïve and pre-immunized patients under treatment with chronic
1107 haemodialysis: a longitudinal quasi-experimental phase IV study. *Clinical microbiology and*
1108 *infection : the official publication of the European Society of Clinical Microbiology and*
1109 *Infectious Diseases*. 2018;24:65-71.

1110 [209] Vesikari T, Brodski N, van Damme P, Diez-Domingo J, Icardi G, Petersen LK, et al. A
1111 Randomized, Double-Blind, Phase III Study of the Immunogenicity and Safety of a 9-Valent
1112 Human Papillomavirus L1 Virus-Like Particle Vaccine (V503) Versus Gardasil in 9-15-Year-
1113 Old Girls. *The Pediatric infectious disease journal*. 2015;34:992-8.

1114 [210] Vesikari T, Karvonen A, Prymula R, Schuster V, Tejedor J, Cohen R, et al. Efficacy of
1115 human rotavirus vaccine against rotavirus gastroenteritis during the first 2 years of life in
1116 European infants: randomised, double-blind controlled study. *Lancet*. 2007;370:1757-63.

1117 [211] Vesikari T, Wysocki J, Beeslaar J, Eiden J, Jiang Q, Jansen KU, et al. Immunogenicity,
1118 Safety, and Tolerability of Bivalent rLP2086 Meningococcal Group B Vaccine Administered
1119 Concomitantly With Diphtheria, Tetanus, and Acellular Pertussis and Inactivated Poliomyelitis
1120 Vaccines to Healthy Adolescents. *Journal of the Pediatric Infectious Diseases Society*.
1121 2016;5:180-7.

1122 [212] Vila-Corcoles A, Ochoa-Gondar O, de Diego C, Satue E, Aragón M, Vila-Rovira A, et al.
1123 Evaluating clinical effectiveness of 13-valent pneumococcal conjugate vaccination against
1124 pneumonia among middle-aged and older adults in Catalonia: results from the EPIVAC cohort
1125 study. *BMC Infectious Diseases*. 2018;18:196.

1126 [213] Villa M, Black S, Groth N, Rothman KJ, Apolone G, Weiss NS, et al. Safety of MF59-
1127 adjuvanted influenza vaccination in the elderly: results of a comparative study of MF59-
1128 adjuvanted vaccine versus nonadjuvanted influenza vaccine in northern Italy. *American journal*
1129 *of epidemiology*. 2013;178:1139-45.

1130 [214] Walter EB, Klein NP, Wodi AP, Rountree W, Todd CA, Wiesner A, et al. Fever after
1131 influenza, diphtheria-tetanus-acellular pertussis, and pneumococcal vaccinations. *Pediatrics*.
1132 2020;145.

1133 [215] Wang L, Chandrasekaran V, Domachowske JB, Li P, Innis BL, Jain VK. Immunogenicity
1134 and Safety of an Inactivated Quadrivalent Influenza Vaccine in US Children 6-35 Months of Age
1135 During 2013-2014: Results From A Phase II Randomized Trial. *Journal of the Pediatric*
1136 *Infectious Diseases Society*. 2016;5:170-9.

1137 [216] Wang SV, Abdurrob A, Spoendlin J, Lewis E, Newcomer SR, Fireman B, et al. Methods
1138 for addressing "innocent bystanders" when evaluating safety of concomitant vaccines.
1139 *Pharmacoepidemiology and drug safety*. 2018;27:405-12.

1140 [217] Yih WK, Lieu TA, Kulldorff M, Martin D, McMahonill-Walraven CN, Platt R, et al.
1141 Intussusception risk after rotavirus vaccination in U.S. infants. *The New England journal of*
1142 *medicine*. 2014;370:503-12.

1143 [218] Yung CF, Chan SP, Soh S, Tan A, Thoon KC. Intussusception and Monovalent Rotavirus
1144 Vaccination in Singapore: Self-Controlled Case Series and Risk-Benefit Study. *J Pediatr*.
1145 2015;167:163-8.e1.

1146 [219] Yung CF, Ma X, Cheung YB, Oh BK, Soh S, Thoon KC. Kawasaki Disease following
1147 administration of 13-valent pneumococcal conjugate vaccine in young children. *Scientific*
1148 *reports*. 2019;9:14705.

1149 [220] Zahid M, Singla I, Good CB, Stone RA, Kim S, Fine MJ, et al. Associations between
1150 Pneumococcal Vaccination and Adverse Outcomes in Patients with Suspected Acute Coronary
1151 Syndrome. *Adv Infect Dis*. 2012:122-34.

1152 [221] Erratum: v72P10 Meningococcal B Adolescent Vaccine Study group. Immunogenicity and
1153 tolerability of a multicomponent meningococcal serogroup B (4CMenB) vaccine in healthy
1154 adolescents in Chile: a phase 2b/3 randomised, observer-blind, placebo-controlled study (*The*
1155 *Lancet* (2012) 379 (617-624)). *Lancet*. 2015;385:1728.

1156 [222] AstraZeneca. A Phase3 Study to Evaluate the Efficacy and Safety of MEDI3250 in
1157 Healthy Japanese Children Age 7 Years Through 18 Years. 2015.

1158 [223] AZ Sint-Jan AV, Pfizer. 13 Valent Pneumococcal Conjugate Vaccine in Chronic Dialysis
1159 Patients. 2013.

1160 [224] Baker MA, Lieu TA, Li LL, Hua W, Qiang YD, Kawai AT, et al. A Vaccine Study Design
1161 Selection Framework for the Postlicensure Rapid Immunization Safety Monitoring Program.
1162 *American Journal of Epidemiology*. 2015;181:608-18.

1163 [225] Baxter R, Lewis E, Fireman B, DeStefano F, Gee J, Klein NP. Case-centered Analysis of
1164 Optic Neuritis After Vaccines. *Clin Infect Dis*. 2016;63:79-81.

1165 [226] Clinical Trials Registry-India. CTRI/2012/07/002820: A study to evaluate safety of
1166 Rotavirus vaccine in Healthy Adult Volunteers followed by Safety, Tolerability and
1167 Immunogenicity evaluation in healthy infants. July 31, 2013.

1168 [227] Colindres R, Wascotte V, Brex A, Clarke C, Hervé C, Kim JH, et al. Post hoc analysis of
1169 reactogenicity trends between dose 1 and dose 2 of the adjuvanted recombinant zoster vaccine in
1170 two parallel randomized trials. *Human Vaccines and Immunotherapeutics*. 2020.

1171 [228] Columbia University, Centers for Disease Control Prevention. FeverText: Assessing Fever
1172 Rates After Vaccination During the 2011-12 Influenza Season Using Text Messaging. 2011.

1173 [229] Columbia University, Centers for Disease Control Prevention. Assessing Fever Rates in
1174 Children Ages 24 to 59 Months After Live Attenuated Influenza Vaccine (LAIV) or Inactivated
1175 Influenza Vaccines (IIV) Using Text Messaging for U.S. Influenza Vaccines in 2012-2013 &
1176 2013-2014. 2013.

1177 [230] Cowling BJ, Thompson MG, Ng TWY, Fang VJ, Perera RAPM, Leung NHL, et al.
1178 Comparative Reactogenicity of Enhanced Influenza Vaccines in Older Adults. *The Journal of*
1179 *infectious diseases*. 2020;222:1383-91.

1180 [231] DeSilva M, Vazquez-Benitez G, Nordin JD, Lipkind HS, Romitti PA, DeStefano F, et al.
1181 Tdap Vaccination During Pregnancy and Microcephaly and Other Structural Birth Defects in
1182 Offspring. *JAMA*. 2016;316:1823-5.

1183 [232] Dezure A, Marechal C, Lal H, Poder A, Ferguson M, Enweonye I, et al. Immunogenicity
1184 and safety of an adjuvanted herpes zoster subunit vaccine candidate when coadministered with a
1185 23-valent pneumococcal polysaccharide vaccine in adults 50 years of age or older: a phase iii,
1186 randomized clinical trial. *Journal of the american pharmacists association*. 2018;58:e72- e3.

1187 [233] Dynavax Technologies Corporation. Safety and Immunogenicity Study of the Hepatitis B
1188 Virus (HBV) Vaccine, HEPLISAV Compared to Engerix-B Vaccine. 2010.

1189 [234] Eucr BE. Evaluation of the immunogenicity and safety of GlaxoSmithKline (GSK)
1190 Biologicals' Quadrivalent Influenza Vaccine Influxplit™ Tetra (Fluarix™ Tetra)
1191 (GSK2321138A) when co-administered with Pneumovax™ 23 in adults 50 years of age and
1192 older. <http://www.who.int/trialssearch/Trial2.aspx?TrialID=EUCTR2014-001118-24-BE>. 2014.
1193 [235] Eucr CZ. Immunogenicity and safety study of Boostrix in pregnant women.
1194 <http://www.who.int/trialssearch/Trial2.aspx?TrialID=EUCTR2014-001119-38-CZ>. 2015.
1195 [236] Everett D. Hepatitis B Vaccine (Recombinant), Adjuvanted (Heplisav-B): Review of
1196 Safety. Vaccines and Related Biological Products Advisory Committee Meeting: U.S. Food &
1197 Drug Administration; July 28, 2017.
1198 [237] Garland SM, Pitisuttithum P, Ngan HYS, Cho CH, Lee CY, Chen CA, et al. Efficacy,
1199 Immunogenicity, and Safety of a 9-Valent Human Papillomavirus Vaccine: Subgroup Analysis
1200 of Participants From Asian Countries. The Journal of infectious diseases. 2018;218:95-108.
1201 [238] Giuliano AR, Joura EA, Garland SM, Huh WK, Iversen O-E, Kjaer SK, et al. Nine-valent
1202 HPV vaccine efficacy against related diseases and definitive therapy: comparison with historic
1203 placebo population. Gynecologic oncology. 2019;154:110-7.
1204 [239] GlaxoSmithKline. To Test 2 Doses of GSK Biologicals' Oral Live Attenuated Human
1205 Rotavirus (HRV) Vaccine in Healthy Infants in Co-administration With Specific Childhood
1206 Vaccines. <https://ClinicalTrials.gov/show/NCT00140686>; 2004.
1207 [240] GlaxoSmithKline. Immunogenicity and Safety Study of a GSK Influenza Vaccine
1208 Candidate in Adults. 2008.
1209 [241] GlaxoSmithKline. Immunogenicity and Safety Study of GSK Biologicals' Herpes Zoster
1210 Vaccine With Various Formulations in Adults >= 50 Years. 2009.
1211 [242] GlaxoSmithKline. Immunogenicity and Safety Study of a GlaxoSmithKline Biologicals'
1212 Candidate Influenza Vaccine in Healthy Children.
1213 <https://ClinicalTrials.gov/show/NCT00985790>; 2009.
1214 [243] GlaxoSmithKline. Safety and Immunogenicity Study of GSK Biologicals' Seasonal
1215 Influenza Candidate Vaccine (GSK2321138A). 2010.
1216 [244] GlaxoSmithKline. A Study to Evaluate the Safety and Immunogenicity of GSK
1217 Biologicals' Seasonal Influenza Vaccine in Children. 2010.
1218 [245] GlaxoSmithKline. A Study to Evaluate the Safety and Immunogenicity of GSK
1219 Biologicals' Seasonal Influenza Vaccine in Adults. 2010.
1220 [246] GlaxoSmithKline. Study to Assess the Efficacy, Immunogenicity and Safety of Liquid
1221 Human Rotavirus Vaccine, in Healthy Chinese Infants. 2010.
1222 [247] GlaxoSmithKline. Study to Evaluate Efficacy, Safety and Immunogenicity of GSK
1223 Biologicals' Herpes Zoster (HZ) Vaccine GSK1437173A in Adults Aged 50 Years and Older.
1224 2010.
1225 [248] GlaxoSmithKline. Immunogenicity and Safety Study of GSK Biologicals' Influenza
1226 Vaccine When Administered in Children. <https://ClinicalTrials.gov/show/NCT01196988>; 2010.
1227 [249] GlaxoSmithKline. Study to Evaluate Immunogenicity and Safety of GlaxoSmithKline
1228 (GSK) Biologicals' Quadrivalent Influenza Vaccine GSK2282512A When Administered to
1229 Children 6 to 35 Months of Age. 2012.
1230 [250] GlaxoSmithKline. Immunogenicity and Safety Study of GlaxoSmithKline (GSK)
1231 Biologicals' Meningococcal Conjugate Vaccine (GSK134612) When Co-administered With
1232 Boostrix® in Subjects Between 11 and 25 Years of Age. 2014.

1233 [251] GlaxoSmithKline. Study to Evaluate Immunogenicity and Safety of GlaxoSmithKline
1234 (GSK) Biologicals' Quadrivalent Influenza Vaccine GSK2282512A When Administered to
1235 Children From 6 to 35 Months of Age. 2014.

1236 [252] GlaxoSmithKline. Study to Evaluate the Immunogenicity and Safety of GlaxoSmithKline
1237 (GSK) Biologicals' Herpes Zoster Vaccine GSK1437173A When Co-administered With GSK
1238 Biologicals' Seasonal Influenza Vaccine GSK2321138A in Adults Aged 50 Years and Older.
1239 2014.

1240 [253] GlaxoSmithKline. Study to Assess the Immunogenicity and Safety of GlaxoSmithKline
1241 (GSK) Biologicals' Herpes Zoster Subunit (HZ/su) Vaccine (GSK1437173A) When Co-
1242 administered With GSK Biologicals' Diphtheria, Tetanus and Pertussis Vaccine (Boostrix®) in
1243 Adults Aged 50 Years and Older. 2015.

1244 [254] GlaxoSmithKline. Evaluation of the Immunogenicity and Safety of GlaxoSmithKline
1245 (GSK) Biologicals' Quadrivalent Influenza Vaccine Influxplit™ Tetra (Fluarix™ Tetra)
1246 (GSK2321138A) When Co-administered With Pneumovax™ 23 in Adults 50 Years of Age and
1247 Older. 2015.

1248 [255] GlaxoSmithKline. Study to Evaluate Immunogenicity and Safety Study of GSK
1249 Biologicals' Herpes Zoster (HZ) Vaccine GSK1437173A When Co-administered With
1250 Pneumovax 23™ in Adults Aged 50 Years and Older. 2015.

1251 [256] GlaxoSmithKline. A Post-marketing, Observational, Retrospective Study to Assess the
1252 Safety of Refortrix™ (Tdap) When Administered During Pregnancy in a Maternal
1253 Immunization Program in Brazil. 2017.

1254 [257] GlaxoSmithKline. Immunogenicity and Safety Study of GlaxoSmithKline (GSK)
1255 Biologicals' Boostrix™ Vaccine in Pregnant Women. 2017.

1256 [258] GlaxoSmithKline. Study to Evaluate Efficacy, Safety and Immunogenicity of
1257 GlaxoSmithKline (GSK) Biologicals' Herpes Zoster (HZ) Vaccine GSK1437173A in Adults
1258 Aged 70 Years and Older. 2017.

1259 [259] GlaxoSmithKline. Immunogenicity and Safety Study of Infanrix Hexa in Healthy Infants
1260 Born to Mothers Vaccinated With Boostrix™ During Pregnancy or Immediately Post-delivery.
1261 2018.

1262 [260] GlaxoSmithKline. Evaluation of Immunogenicity and Safety of a Booster Dose of Infanrix
1263 Hexa™ in Healthy Infants Born to Mothers Vaccinated With Boostrix™ During Pregnancy or
1264 Immediately Post-delivery. 2019.

1265 [261] Guevara A, Cabello R, Woelber L, Moreira ED, Jr., Joura E, Reich O, et al. Antibody
1266 persistence and evidence of immune memory at 5years following administration of the 9-valent
1267 HPV vaccine. *Vaccine*. 2017;35:5050-7.

1268 [262] Guevara AM, Suarez E, Victoria A, Ngan HYS, Hirschberg AL, Fedrizzi E, et al. Maternal
1269 transfer of anti HPV 6 and 11 antibodies upon immunization with the 9-valent HPV vaccine.
1270 *Human Vaccines and Immunotherapeutics*. 2019;15:141-5.

1271 [263] Hambidge SJ, Newcomer SR, Narwaney KJ, Glanz JM, Daley MF, Xu S, et al. Timely
1272 versus delayed early childhood vaccination and seizures. *Pediatrics*. 2014;133:e1492-9.

1273 [264] Hedrick J, Christensen S, Chang LJ, Pan J, Jordanov E, Dhingra M. Study of the
1274 immunogenicity and safety of an investigational quadrivalent meningococcal conjugate vaccine
1275 (MENACYW-TT) when co-administered with other vaccines in healthy adolescents. *Open
1276 forum infectious diseases*. 2018;5:S570- .

1277 [265] Hedrick J, Simon MW, Christensen S, Anez G, Pan J, Jordanov E, et al. Immunogenicity
1278 and safety of a booster dose of a quadrivalent meningococcal conjugate vaccine (menacyw-TT)
1279 in adolescents and adults. *Open forum infectious diseases*. 2019;6:S959- .

1280 [266] Helsinki University Central Hospital, Tampere University Hospital. Immunogenicity of
1281 13-valent Pneumococcal Conjugate Vaccine Compared to the Pneumococcal Polysaccharide
1282 Vaccine in Adult Kidney and Liver Transplant Patients.
1283 <https://ClinicalTrials.gov/show/NCT01781871>; 2013.

1284 [267] Hospital Universitario Dr. Jose E. Gonzalez. Immunogenicity and Safety of an Acellular
1285 DPT Vaccine During Pregnancy. 2011.

1286 [268] Hyer R, McGuire DK, Xing B, Jackson S, Janssen R. Safety of a two-dose investigational
1287 hepatitis B vaccine, HBsAg-1018, using a toll-like receptor 9 agonist adjuvant in adults.
1288 *Vaccine*. 2018;36:2604-11.

1289 [269] Hyer RN, Janssen RS. Immunogenicity and safety of a 2-dose hepatitis B vaccine,
1290 HBsAg/CpG 1018, in persons with diabetes mellitus aged 60-70years. *Vaccine*. 2019;37:5854-
1291 61.

1292 [270] Jeanfreau R, Esteves-Jaramillo A, Neveu D, Jordanov E, Dhingra MS. Immunogenicity
1293 and safety of a quadrivalent meningococcal conjugate vaccine (MenACYW-TT) administered in
1294 individuals 56 years of age and older. *Open forum infectious diseases*. 2018;5.

1295 [271] Joura EA, Giuliano AR, Iversen O-E, Bouchard C, Mao C, Mehlsen J, et al. A 9-valent
1296 HPV vaccine against infection and intraepithelial neoplasia in women. *The New England journal*
1297 *of medicine*. 2015;372:711-23.

1298 [272] Kantso B, Halkjaer S, Thomsen O, Belard E, Gottschalck IB, Jorgensen CS, et al. Specific
1299 antibody response to two pneumococcal vaccines in crohn's disease patients treated with
1300 immunosuppressive drugs alone or in combination with biological therapy. *Gastroenterology*.
1301 2015;148:S176- S7.

1302 [273] Karolinska University Hospital. Pneumococcal Vaccine in Untreated CLL Patients. 2013.

1303 [274] Kharbanda EO, Vazquez-Benitez G, Lipkind HS, Klein NP, Cheetham TC, Naleway A, et
1304 al. Evaluation of the association of maternal pertussis vaccination with obstetric events and birth
1305 outcomes. *JAMA*. 2014;312:1897-904.

1306 [275] Korea University Guro Hospital. Influenza Vaccine and Pneumococcal Vaccine.
1307 <https://ClinicalTrials.gov/show/NCT02582047>; 2012.

1308 [276] Korea University Guro Hospital. Immunogenicity and Safety of a Tetanus-diphtheria
1309 Vaccine and a 13-valent Pneumococcal Conjugate Vaccine. 2013.

1310 [277] Korea University Guro Hospital, Novartis. MF59-adjuvanted Influenza Vaccine and 23-
1311 valent Pneumococcal Polysaccharide Vaccine. 2013.

1312 [278] Korea University Guro Hospital, Pfizer. Immunogenicity and Safety of PCV13 and Flud
1313 in Adults Aged ≥ 60 Years. 2014.

1314 [279] Laval University. Immunogenicity and Safety of Gardasil-9 and Cervarix. 2015.

1315 [280] Li R-C, Huang T, Li Y, Wang L-H, Tao J, Fu B, et al. Immunogenicity and reactogenicity
1316 of the human rotavirus vaccine, RIX4414 oral suspension, when co-administered with routine
1317 childhood vaccines in Chinese infants. *Human vaccines & immunotherapeutics*. 2016;12:785-93.

1318 [281] Li R-C, Li Y-P, Mo Z-J, Luo D, Huang T, Kong J-L, et al. Reactogenicity and safety of a
1319 liquid human rotavirus vaccine (RIX4414) in healthy adults, children and infants in China:
1320 randomized, double-blind, placebo-controlled Phase I studies. *Human vaccines &*
1321 *immunotherapeutics*. 2013;9:1638-42.

1322 [282] Lopez-Fauqued M, Campora L, Delannois F, El Idrissi M, Oostvogels L, De Looze FJ, et
1323 al. Safety profile of the adjuvanted recombinant zoster vaccine: Pooled analysis of two large
1324 randomised phase 3 trials. *Vaccine*. 2019;37:2482-93.

1325 [283] Luxembourg A, Brown D, Bouchard C, Giuliano AR, Iversen OE, Joura EA, et al. Phase II
1326 studies to select the formulation of a multivalent HPV L1 virus-like particle (VLP) vaccine.
1327 *Human Vaccines and Immunotherapeutics*. 2015;11:1313-22.

1328 [284] Mayrand MH, Bautista O, Moeller E, Ritter M, Luxembourg A. End of study efficacy,
1329 immunogenicity and safety of a novel 9-valent HPV 11 virus-like particle vaccine in 16-26 year
1330 old women. *International journal of gynecology and obstetrics (varpagings)*. 2015;131:E270.

1331 [285] McElhaney JE, Lal H, Cunningham AL, Levin MJ, Chlibek R, Diez-Domingo J, et al.
1332 Efficacy, immunogenicity and safety of an investigational subunit adjuvanted herpes zoster
1333 vaccine in adults aged 60 years and older: results from the zoe-50 and zoe-70 efficacy studies.
1334 *Open forum infectious diseases*. 2016;3.

1335 [286] MedImmune LLC. A Study to Evaluate the Immunogenicity of Quadrivalent LAIV in
1336 Adults 18 to 49 Years of Age. <https://ClinicalTrials.gov/show/NCT00860067>; 2009.

1337 [287] MedImmune LLC. A Study to Evaluate the Immunogenicity of Quadrivalent Live
1338 Attenuated Influenza Vaccine (LAIV) in Children.
1339 <https://ClinicalTrials.gov/show/NCT01091246>; 2010.

1340 [288] MedImmune LLC. Postmarketing Safety Study of Q/LAIV in Subjects 2 Through 49
1341 Years of Age. <https://ClinicalTrials.gov/show/NCT01985997>; 2014.

1342 [289] MedImmune LLC, AstraZeneca. Evaluate the Shedding and Immunogenicity of Different
1343 Formulations of FluMist in Children 24 to <48 Months of Age. 2017.

1344 [290] Merck Sharp, Dohme Corp. Efficacy, Safety, and Immunogenicity of V260 in Healthy
1345 Chinese Infants (V260-024). 2015.

1346 [291] Merck Sharp, Dohme Corp. Concomitant Use of Hepatitis A Vaccine, Inactivated With
1347 Haemophilus b Conjugate Vaccine (Meningococcal Protein Conjugate) and Diphtheria and
1348 Tetanus Toxoids and Acellular Pertussis Vaccine Adsorbed Given to Healthy Children 15
1349 Months of Age (V251-068). 2006.

1350 [292] Merck Sharp, Dohme Corp. Broad Spectrum HPV (Human Papillomavirus) Vaccine Study
1351 in 16-to 26-Year-Old Women (V503-001). 2007.

1352 [293] Merck Sharp, Dohme Corp. Immunogenicity and Safety Study of Proquad® and Infanrix®
1353 Hexa When Administered Concomitantly (V221-035). 2007.

1354 [294] Merck Sharp, Dohme Corp. Phase III Placebo-controlled Study of V260 (RotaTeq™) in
1355 Japanese Healthy Infants (V260-029)(COMPLETED).
1356 <https://ClinicalTrials.gov/show/NCT00718237>; 2008.

1357 [295] Merck Sharp, Dohme Corp. A Study of V503, a 9-valent Human Papillomavirus (9vHPV)
1358 Vaccine in Females 12-26 Years of Age Who Have Previously Received GARDASIL™ (V503-
1359 006). 2010.

1360 [296] Merck Sharp, Dohme Corp. Immunogenicity and Tolerability of V503 Versus GARDASIL
1361 (V503-009). 2011.

1362 [297] Merck Sharp, Dohme Corp. A Study to Compare Immune Response of V503 to Gardasil in
1363 16- to 26-year-old Men (V503-020). 2014.

1364 [298] Merck Sharp, Dohme Corp., MCM Vaccines B.V. A Study of V419 Given Concomitantly
1365 With Prevnar 13™ and RotaTeq™ (V419-006). 2011.

1366 [299] Merck Sharp, Dohme Corp., MCM Vaccines B.V. Safety, Tolerability, and
1367 Immunogenicity of V419 Given Concomitantly With Prevnar 13™ and RotaTeq™ (V419-005).
1368 2011.

1369 [300] Munoz FM, Bond NH, Maccato M. Safety and Immunogenicity of Tetanus Diphtheria and
1370 Acellular Pertussis (Tdap) Immunization During Pregnancy in Mothers and Infants: A
1371 Randomized Clinical Trial (vol 311, pg 1760, 2014). Jama-Journal of the American Medical
1372 Association. 2017;317:442-.

1373 [301] National Institute of Allergy Infectious Diseases. Pertussis Vaccine in Healthy Pregnant
1374 Women. 2009.

1375 [302] Novartis, Novartis Vaccines. Study to Evaluate the Safety of Novartis MenACWY
1376 Conjugate Vaccine When Administered With Routine Infant Vaccinations to Healthy Infants.
1377 2008.

1378 [303] Novartis, Novartis Vaccines. A Phase 3b, Randomized, Open-Label Study to Evaluate the
1379 Safety and Immunogenicity of Select Travel Vaccines When Administered Concomitantly With
1380 MenACWY in Adults. 2011.

1381 [304] Novartis, Novartis Vaccines. A Phase 4, Placebo-Controlled, Randomized Study to
1382 Evaluate the Immunogenicity and Safety of HPV and Tdap When Administered With
1383 MenACWY in Adolescents. 2011.

1384 [305] Novartis Vaccines, GlaxoSmithKline, Novartis. Study to Evaluate the Safety and
1385 Immunogenicity of Combined Hepatitis A/B Vaccine With MenACWY-CRM Conjugate
1386 Vaccine. 2011.

1387 [306] Novartis Vaccines, Novartis. Safety and Immunogenicity of Meningococcal ACWY
1388 Conjugate Versus Polysaccharide Vaccine in Children 2 to 10 Years of Age. 2006.

1389 [307] Novartis Vaccines, Novartis. A Study to Evaluate Safety and Immune Response of
1390 Novartis Meningococcal ACWY Vaccine In Infants. 2007.

1391 [308] Novartis Vaccines, Novartis. Safety, Tolerability and Immunogenicity of Novartis
1392 Meningococcal B Recombinant Vaccine Administered to Healthy Adolescents According to
1393 Different Vaccination Schedules. 2008.

1394 [309] Novartis Vaccines, Novartis. A Multi-center, Observer-blind, Placebo-controlled,
1395 Randomized Study to Evaluate the Immunogenicity and Safety of MenACWY in Adolescents
1396 and Adults in Korea. 2010.

1397 [310] Novartis Vaccines, Novartis. Safety and Immunogenicity of MF59C.1 Adjuvanted
1398 Trivalent Subunit Influenza Vaccine in Elderly Subjects. 2010.

1399 [311] Novartis Vaccines, Novartis. Safety and Immunogenicity of Three Influenza Vaccines
1400 Adults Ages 18 and Older. 2014.

1401 [312] Novartis Vaccines, Novartis. Safety and Immunogenicity of Three Influenza Vaccines in
1402 Children Aged 4 Years Old to Less Than 18 Years Old. 2014.

1403 [313] Novartis Vaccines, Novartis. Safety and Immunogenicity Study of Two Doses of Novartis
1404 Meningococcal Serogroup B Recombinant Vaccine in Adolescents Aged 11-17 Years. 2014.

1405 [314] Novartis Vaccines and Diagnostics S.r.l. A phase III, double-blind, randomized, placebo-
1406 controlled, multi-country and multi-center study to assess the efficacy and safety of two doses of
1407 GSK Biologicals' oral live attenuated human rotavirus (HRV) vaccine in healthy infants.\. 2014.

1408 [315] Oostvogels L, Heineman TC, Johnson RW, Levin MJ, McElhaney JE, Van den Steen P, et
1409 al. Medical conditions at enrollment do not impact efficacy and safety of the adjuvanted
1410 recombinant zoster vaccine: a pooled post-hoc analysis of two parallel randomized trials. Human
1411 vaccines & immunotherapeutics. 2019;15:2865-72.

1412 [316] Perrett K, Garcia IC, Halperin S, Nolan T, Virta M, Stranak Z, et al. Pertussis vaccination
1413 during pregnancy: a multicountry randomised placebo-controlled trial assessing immunogenicity
1414 and safety of dTpa in mothers and their infants. *Bjog*. 2019;126:173- 4.

1415 [317] Perrett KP, Halperin SA, Nolan T, Martinez AC, Martinon-Torres F, Garcia-Sicilia J, et al.
1416 Impact of tetanus-diphtheria-acellular pertussis immunization during pregnancy on subsequent
1417 infant immunization seroresponses: follow-up from a large randomized placebo-controlled trial.
1418 *Vaccine*. 2020;38:2105-14.

1419 [318] Petersen LK, Restrepo J, Moreira ED, Iversen OE, Pitisuttithum P, Van Damme P, et al.
1420 Impact of baseline covariates on the immunogenicity of the 9-valent HPV vaccine – A combined
1421 analysis of five phase III clinical trials. *Papillomavirus Research*. 2017;3:105-15.

1422 [319] Peterson J, Hedrick J, Pan J, Neveu D, Jordanov E, Dhingra MS. Immunogenicity and
1423 safety of a quadrivalent meningococcal conjugate vaccine (menacyw-TT) administered in adults
1424 18-55 years of age. *Open forum infectious diseases*. 2019;6:S957- .

1425 [320] Peterson J, Hedrick J, Pan J, Neveu D, Jordanov E, Dhingra MS. Immunogenicity and
1426 safety of a quadrivalent meningococcal conjugate vaccine (menacyw-TT) administered in
1427 adolescents 10-17 years of age. *Open forum infectious diseases*. 2019;6:S958- .

1428 [321] Pfizer. Study Evaluating the Impact of a 13-valent Pneumococcal Conjugate Vaccine on
1429 Nasopharyngeal Colonization. 2007.

1430 [322] Pfizer. Study Evaluating 13 Valent Pneumococcal Conjugate Vaccine With Trivalent
1431 Inactivated Influenza Vaccine. <https://ClinicalTrials.gov/show/NCT00521586>; 2007.

1432 [323] Pfizer. Study Evaluating Safety and Immunogenicity of 13-Valent Pneumococcal
1433 Conjugate Vaccine With Influenza Vaccine in Adults.
1434 <https://ClinicalTrials.gov/show/NCT00492557>; 2007.

1435 [324] Pfizer. Study Evaluating the Efficacy of a 13-Valent Pneumococcal Conjugate Vaccine
1436 (13vPnC) in Adults. 2008.

1437 [325] Pfizer. A Study Evaluating Safety And Immunogenicity Of Meningococcal B Rlp2086
1438 Vaccine In Adolescents. 2009.

1439 [326] Pfizer. Trial Evaluating a 13-valent Pneumococcal Conjugate Vaccine Given With
1440 Diphtheria, Tetanus, and Acellular Pertussis Vaccine (DTaP) in Healthy Japanese Infants. 2010.

1441 [327] Pfizer. A Trial to Assess the Safety, Tolerability and Immunogenicity of Repevax and
1442 rLP2086 Vaccine When Given Together in Healthy Subjects Aged ≥ 11 to < 19 Years. 2011.

1443 [328] Pfizer. A Global Phase 3 Safety Study of 120 mcg rLP2086 Vaccine in Adolescents and
1444 Young Adults Aged 10 to 25 Years. 2012.

1445 [329] Pfizer. Immunogenicity and Safety Study of 1 and 2 Doses of GlaxoSmithKline (GSK)
1446 Biologicals' Meningococcal Vaccine MenACWY-TT (GSK134612) in Toddlers, Persistence up
1447 to 5 Years After Vaccination and Co-administration With Pfizer's Prevenar 13™ Vaccine. 2013.

1448 [330] Pfizer. Concomitant Administration of 13-valent Pneumococcal Conjugate Vaccine
1449 (13vPnC) With Influenza Vaccine in 23-valent Pneumococcal Polysaccharide (23vPS) Pre-
1450 vaccinated Adults. 2015.

1451 [331] Pfizer Winawoso. Study Evaluating A 13-Valent Pneumococcal Conjugate Vaccine
1452 Administered To Infants In Korea. <https://ClinicalTrials.gov/show/NCT00689351>; 2008.

1453 [332] Protein Sciences Corporation. Protective Efficacy of Flublok® Quadrivalent Versus
1454 Licensed Inactivated Influenza Vaccine in Adults ≥ 50 Years of Age. 2015.

1455 [333] Protein Sciences Corporation, Syneos Health, Department of Health Human Services.
1456 Safety and Immunogenicity of Flublok Quadrivalent vs IIV4 in Adults 18-49 Years of Age.
1457 2015.

1458 [334] Rowhani-Rahbar A, Fireman B, Lewis E, Nordin J, Naleway A, Jacobsen SJ, et al. Effect
1459 of age on the risk of Fever and seizures following immunization with measles-containing
1460 vaccines in children. JAMA pediatrics. 2013;167:1111-7.

1461 [335] Ruiz-Sternberg AM, Moreira ED, Jr., Restrepo JA, Lazcano-Ponce E, Cabello R, Silva A,
1462 et al. Efficacy, immunogenicity, and safety of a 9-valent human papillomavirus vaccine in Latin
1463 American girls, boys, and young women. Papillomavirus research (Amsterdam, Netherlands).
1464 2018;5:63-74.

1465 [336] Sanofi Pasteur a Sanofi Company. Immune Lot Consistency, Immunogenicity, and Safety
1466 of an Investigational Quadrivalent Meningococcal Conjugate Vaccine.

1467 [337] Sanofi Pasteur a Sanofi Company, Merck Sharp, Dohme Corp., Sanofi. Study of PR5I, a
1468 Pediatric Combination Vaccine With Enhanced Hepatitis B Component Given Concomitantly
1469 With Prevnar®. 2006.

1470 [338] Sanofi Pasteur a Sanofi Company, Sanofi. 36-Month Post-marketing Surveillance and
1471 Analysis of Menactra Vaccine in 2-10 Year Olds. 2005.

1472 [339] Sanofi Pasteur a Sanofi Company, Sanofi. Descriptive, Post-marketing, Surveillance
1473 Safety Study of Menactra Vaccine. 2005.

1474 [340] Sanofi Pasteur a Sanofi Company, Sanofi. Database Surveillance Safety Study of
1475 PENTACEL® Vaccine. 2008.

1476 [341] Sanofi Pasteur a Sanofi Company, Sanofi. A Study of Influenza Virus Vaccines in
1477 Children and Adults. 2009.

1478 [342] Sanofi Pasteur a Sanofi Company, Sanofi. Study of Quadrivalent Influenza Vaccine
1479 Among Children. 2010.

1480 [343] Sanofi Pasteur a Sanofi Company, Sanofi. Study of Quadrivalent Influenza Vaccine
1481 Among Adults. <https://ClinicalTrials.gov/show/NCT01218646>; 2010.

1482 [344] Sanofi Pasteur a Sanofi Company, Sanofi. Post-licensure Safety Surveillance Study of
1483 Menactra Vaccine When Administered As a 2-dose Schedule to Children. 2011.

1484 [345] Sanofi Pasteur a Sanofi Company, Sanofi. Safety and Immunogenicity in Adults of
1485 Revaccination With Adacel® Vaccine 10 Years After a Previous Dose. 2011.

1486 [346] Sanofi Pasteur a Sanofi Company, Sanofi. Immunogenicity and Safety of an
1487 Investigational Quadrivalent Meningococcal Conjugate Vaccine in Healthy Adolescents. 2014.

1488 [347] Sanofi Pasteur a Sanofi Company, Sanofi. Immunogenicity and Safety of a Booster Dose
1489 of an Investigational Quadrivalent Meningococcal Conjugate Vaccine. 2016.

1490 [348] Sanofi Pasteur a Sanofi Company, Sanofi. Immunogenicity and Safety of an
1491 Investigational Quadrivalent Meningococcal Conjugate Vaccine in Adults 56 Years and Older.
1492 2016.

1493 [349] Sanofi Pasteur a Sanofi Company, Sanofi. Safety and Immunogenicity of High-Dose
1494 Quadrivalent Influenza Vaccine in Participants ≥ 65 Years in the US. 2017.

1495 [350] Sanofi Pasteur aSC, Sanofi. Study of a Quadrivalent Meningococcal Conjugate Vaccine in
1496 Subjects Aged 56 and Older. <https://ClinicalTrials.gov/show/NCT01732627>; 2012.

1497 [351] Sanofi Pasteur aSC, Sanofi. Study of an Investigational Quadrivalent Meningococcal
1498 Conjugate Vaccine Administered in Children Aged 2 to 9 Years.
1499 <https://ClinicalTrials.gov/show/NCT03077438>; 2017.

1500 [352] Scott Halperin, IWK Health Centre, Sanofi Pasteur a Sanofi Company, Dalhousie
1501 University. Pertussis Maternal Immunization Study. 2007.

1502 [353] Scott J. Statistical Analysis of Heplisav-B AMI Risk. Vaccines and Related Biological
1503 Products Advisory Committee Meeting: U.S. Food & Drug Administration; July 28, 2017.

1504 [354] Seqirus. Phase 3 Safety and Immunogenicity Study of aQIV in Elderly Adults.

1505 [355] Seqirus. A Study to Evaluate the Immunogenicity and Safety of bioCSL Quadrivalent

1506 Influenza Vaccine (QIV) in Adults Aged 18 Years and Above. 2014.

1507 [356] Simon MW, Brandon D, Christensen S, Baccarini C, Jordanov E, Dhingra MS. Safety and

1508 immunogenicity of a quadrivalent meningococcal conjugate vaccine (menacyw-TT)

1509 administered in healthy meningococcal vaccine-naïve children (2-9 years). Open forum

1510 infectious diseases. 2019;6:S958- S9.

1511 [357] Statens Serum Institut. Pneumococcal vaccination of Crohn patients - A randomized, non-

1512 blinded phase 4 clinical trial with the purpose of investigating the immune response against two

1513 different pneumococcal vaccines in patients with Crohn's disease. 2013.

1514 [358] Statens Serum Institut, Hvidovre University Hospital, Herlev Hospital. Pneumococcal

1515 Vaccination of Crohn Patients. 2013.

1516 [359] The University of Hong Kong, Centers for Disease Control Prevention. Immunogenicity of

1517 Alternative Annual Influenza Vaccination Strategies in Older Adults in Hong Kong. 2017.

1518 [360] U.S Food and Drug Administration. Clinical Review, November 9, 2017-Heplisav-B: BLA

1519 Clinical Review Memorandum. 2017.

1520 [361] University D, Centers for Disease Control Prevention, Kaiser Permanente. Fever After

1521 Simultaneous Versus Sequential Vaccination in Young Children. 2018.

1522 [362] University of Newcastle, National Health and Medical Research Council. Pneumococcal

1523 polysaccharide vaccine versus Normal Saline for primary prevention of heart attacks and strokes

1524 in at-risk Australians aged 55-60. 2015.

1525 [363] University of Siena, Ministry of Education Universities, Research Italy. Serological

1526 Response to Antipneumococcal Vaccination and Impact on Streptococcus Pneumoniae Nasal

1527 Carriage in HIV Adults. 2011.

1528 [364] van Werkhoven CH, Bonten MJM. The Community-Acquired Pneumonia immunization

1529 Trial in Adults (CAPiTA): what is the future of pneumococcal conjugate vaccination in elderly?

1530 Future microbiology. 2015;10:1405-13.

1531 [365] Vila-Corcoles A, Ochoa-Gondar O, Rodriguez-Blanco T, de Diego-Cabanes C, Satue-

1532 Gracia E, Vila-Rovira A, et al. Evaluating clinical effectiveness of pneumococcal vaccination in

1533 preventing stroke: the CAPAMIS Study, 3-year follow-up. Journal of stroke and cerebrovascular

1534 diseases : the official journal of National Stroke Association. 2014;23:1577-84.

1535 [366] Vila-Corcoles A, Ochoa-Gondar O, Rodriguez-Blanco T, Gutierrez-Perez A, Vila-Rovira

1536 A, Gomez F, et al. Clinical effectiveness of pneumococcal vaccination against acute myocardial

1537 infarction and stroke in people over 60 years: the CAPAMIS study, one-year follow-up. BMC

1538 Public Health. 2012;12:222.

1539 [367] Wyeth is now a wholly owned subsidiary of Pfizer. Study to Evaluate a 13-valent

1540 Pneumococcal Conjugate Vaccine in Elderly Subjects.

1541 <https://ClinicalTrials.gov/show/NCT00269672>; 2005.

1542 [368] Wyeth is now a wholly owned subsidiary of Pfizer. Study Evaluating the Safety,

1543 Tolerability and Immunogenicity of 13vPnC as a 2-Dose Regimen or With 23vPS.

1544 <https://ClinicalTrials.gov/show/NCT00574548>; 2007.

1545 [369] Yih K, Lieu T, Kulldorff M, Martin D, McMahonill-Walraven C, Platt R, et al.

1546 Intussusception Risk After Rotavirus Vaccination in U.S. Infants. Available at: [http://www.mini-](http://www.mini-sentinel.org/work_products/PRISM/Mini-Sentinel_PRISM_Rotavirus-and-intussusception-Report.pdf)

1547 [sentinel.org/work_products/PRISM/Mini-Sentinel_PRISM_Rotavirus-and-intussusception-](http://www.mini-sentinel.org/work_products/PRISM/Mini-Sentinel_PRISM_Rotavirus-and-intussusception-Report.pdf)

1548 [Report.pdf](http://www.mini-sentinel.org/work_products/PRISM/Mini-Sentinel_PRISM_Rotavirus-and-intussusception-Report.pdf). Mini-Sentinel Coordinating Center; June 2013.

1549 [370] Andrews N, Stowe J, Miller E, Svanstrom H, Johansen K, Bonhoeffer J, et al. A
1550 collaborative approach to investigating the risk of thrombocytopenic purpura after measles-
1551 mumps-rubella vaccination in England and Denmark. *Vaccine*. 2012;30:3042-6.

1552 [371] Andrews N, Stowe J, Wise L, Miller E. Post-licensure comparison of the safety profile of
1553 diphtheria/tetanus/whole cell pertussis/haemophilus influenza type b vaccine and a 5-in-1
1554 diphtheria/tetanus/acellular pertussis/haemophilus influenza type b/polio vaccine in the United
1555 Kingdom. *Vaccine*. 2010;28:7215-20.

1556 [372] Armah GE, Sow SO, Breiman RF, Dallas MJ, Tapia MD, Feikin DR, et al. Efficacy of
1557 pentavalent rotavirus vaccine against severe rotavirus gastroenteritis in infants in developing
1558 countries in sub-Saharan Africa: a randomised, double-blind, placebo-controlled trial. *Lancet*.
1559 2010;376:606-14.

1560 [373] Barbosa CM, Terreri MT, Rosario PO, de Moraes-Pinto MI, Silva CA, Hilario MO.
1561 Immune response and tolerability of varicella vaccine in children and adolescents with systemic
1562 lupus erythematosus previously exposed to varicella-zoster virus. *Clin Exp Rheumatol*.
1563 2012;30:791-8.

1564 [374] Barrett PN, Berezuk G, Fritsch S, Aichinger G, Hart MK, El-Amin W, et al. Efficacy,
1565 safety, and immunogenicity of a Vero-cell-culture-derived trivalent influenza vaccine: a
1566 multicentre, double-blind, randomised, placebo-controlled trial. *Lancet*. 2011;377:751-9.

1567 [375] Baxter R, Bakshi N, Fireman B, Lewis E, Ray P, Vellozzi C, et al. Lack of association of
1568 guillain-barre syndrome with vaccinations. *Clin Infect Dis*. 2013;57:197-204.

1569 [376] Baxter R, Toback SL, Sifakis F, Hansen J, Bartlett J, Aukes L, et al. A postmarketing
1570 evaluation of the safety of Ann Arbor strain live attenuated influenza vaccine in adults 18-49
1571 years of age. *Vaccine*. 2012;30:3053-60.

1572 [377] Baxter R, Toback SL, Sifakis F, Hansen J, Bartlett J, Aukes L, et al. A postmarketing
1573 evaluation of the safety of Ann Arbor strain live attenuated influenza vaccine in children 5
1574 through 17 years of age. *Vaccine*. 2012;30:2989-98.

1575 [378] Baxter R, Tran TN, Hansen J, Emery M, Fireman B, Bartlett J, et al. Safety of
1576 ZostavaxTM--a cohort study in a managed care organization. *Vaccine*. 2012;30:6636-41.

1577 [379] Benchimol EI, Hawken S, Kwong JC, Wilson K. Safety and utilization of influenza
1578 immunization in children with inflammatory bowel disease. *Pediatrics*. 2013;131:e1811-20.

1579 [380] Bernsen RMD, de Jongste JC, Koes BW, Aardoom HA, van der Wouden JC. Diphtheria
1580 tetanus pertussis poliomyelitis vaccination and reported atopic disorders in 8-12-year-old
1581 children. *Vaccine*. 2006;24:2035-42.

1582 [381] Bertuola F, Morando C, Menniti-Ippolito F, Da Cas R, Capuano A, Perilongo G, et al.
1583 Association between drug and vaccine use and acute immune thrombocytopenia in childhood: a
1584 case-control study in Italy. *Drug Safety*. 2010;33:65-72.

1585 [382] Bhatla N, Suri V, Basu P, Shastri S, Datta SK, Bi D, et al. Immunogenicity and safety of
1586 human papillomavirus-16/18 AS04-adjuvanted cervical cancer vaccine in healthy Indian
1587 women.[Erratum appears in *J Obstet Gynaecol Res*. 2010 Apr;36(2):466]. *J Obstet Gynaecol*
1588 *Res*. 2010;36:123-32.

1589 [383] Block SL, Brown DR, Chatterjee A, Gold MA, Sings HL, Meibohm A, et al. Clinical trial
1590 and post-licensure safety profile of a prophylactic human papillomavirus (Types 6, 11, 16, and
1591 18) L1 virus-like particle vaccine. *Pediatric Infectious Disease Journal*. 2010;29:95-101.

1592 [384] Block SL, Vesikari T, Goveia MG, Rivers SB, Adeyi BA, Dallas MJ, et al. Efficacy,
1593 immunogenicity, and safety of a pentavalent human-bovine (WC3) reassortant rotavirus vaccine
1594 at the end of shelf life. *Pediatrics*. 2007;119:11-8.

1595 [385] Buttery JP, Danchin MH, Lee KJ, Carlin JB, McIntyre PB, Elliott EJ, et al. Intussusception
1596 following rotavirus vaccine administration: post-marketing surveillance in the National
1597 Immunization Program in Australia. *Vaccine*. 2011;29:3061-6.

1598 [386] Capeding MRZ, Nohynek H, Pascual LG, Kayhty H, Sombrero LT, Eskola J, et al. The
1599 immunogenicity of three Haemophilus influenzae type B conjugate vaccines after a primary
1600 vaccination series in Philippine infants. *American Journal of Tropical Medicine and Hygiene*.
1601 1996;55:516-20.

1602 [387] Chang C-C, Chang M-H, Lin T-Y, Lee H-C, Hsieh W-S, Lee P-I. Experience of
1603 pentavalent human-bovine reassortant rotavirus vaccine among healthy infants in Taiwan.
1604 *Journal of the Formosan Medical Association*. 2009;108:280-5.

1605 [388] Chao C, Klein NP, Velicer CM, Sy LS, Slezak JM, Takhar H, et al. Surveillance of
1606 autoimmune conditions following routine use of quadrivalent human papillomavirus vaccine. *J*
1607 *Intern Med*. 2012;271:193-203.

1608 [389] Chen RT, Glasser JW, Rhodes PH, Davis RL, Barlow WE, Thompson RS, et al. Vaccine
1609 safety datalink project: A new tool for improving vaccine safety monitoring in the United States.
1610 *Pediatrics*. 1997;99:765-73.

1611 [390] Clark LR, Myers ER, Huh W, Joura EA, Paavonen J, Perez G, et al. Clinical trial
1612 experience with prophylactic human papillomavirus 6/11/16/18 vaccine in young black women.
1613 *Journal of Adolescent Health*. 2013;52:322-9.

1614 [391] Crawford NW, Cheng A, Andrews N, Charles PG, Clothier HJ, Day B, et al. Guillain-
1615 Barre syndrome following pandemic (H1N1) 2009 influenza A immunisation in Victoria: a self-
1616 controlled case series. *Med J Aust*. 2012;197:574-8.

1617 [392] De Carvalho N, Teixeira J, Roteli-Martins CM, Naud P, De Borba P, Zahaf T, et al.
1618 Sustained efficacy and immunogenicity of the HPV-16/18 AS04-adjuvanted vaccine up to 7.3
1619 years in young adult women. *Vaccine*. 2010;28:6247-55.

1620 [393] Dennehy PH, Brady RC, Halperin SA, Ward RL, Alvey JC, Fischer FH, Jr., et al.
1621 Comparative evaluation of safety and immunogenicity of two dosages of an oral live attenuated
1622 human rotavirus vaccine. *Pediatric Infectious Disease Journal*. 2005;24:481-8.

1623 [394] Dodds L, Macdonald N, Scott J, Spencer A, Allen VM, McNeil S. The association
1624 between influenza vaccine in pregnancy and adverse neonatal outcomes. *J Obstet Gynaecol Can*.
1625 2012;34:714-20.

1626 [395] Duderstadt SK, Rose CE, Jr., Real TM, Sabatier JF, Stewart B, Ma G, et al. Vaccination
1627 and risk of type 1 diabetes mellitus in active component U.S. Military, 2002-2008. *Vaccine*.
1628 2012;30:813-9.

1629 [396] Eder L, Law T, Chandran V, Shanmugarajah S, Shen H, Rosen CF, et al. Association
1630 between environmental factors and onset of psoriatic arthritis in patients with psoriasis. *Arthritis*
1631 *Care and Research*. 2011;63:1091-7.

1632 [397] Englund JA, Walter E, Black S, Blatter M, Nyberg J, Ruben FL, et al. Safety and
1633 immunogenicity of trivalent inactivated influenza vaccine in infants: a randomized double-blind
1634 placebo-controlled study. *Pediatric Infectious Disease Journal*. 2010;29:105-10.

1635 [398] Eurich DT, Johnstone JJ, Minhas-Sandhu JK, Marrie TJ, Majumdar SR. Pneumococcal
1636 vaccination and risk of acute coronary syndromes in patients with pneumonia: population-based
1637 cohort study. *Heart*. 2012;98:1072-7.

1638 [399] Farez MF, Ysrraelit MC, Fiol M, Correale J. H1N1 vaccination does not increase risk of
1639 relapse in multiple sclerosis: a self-controlled case-series study. *Multiple Sclerosis*. 2012;18:254-
1640 6.

1641 [400] Fell DB, Sprague AE, Liu N, Yasseen AS, 3rd, Wen S-W, Smith G, et al. H1N1 influenza
1642 vaccination during pregnancy and fetal and neonatal outcomes. *American Journal of Public*
1643 *Health*. 2012;102:e33-40.

1644 [401] Frey S, Vesikari T, Szymczakiewicz-Multanowska A, Lattanzi M, Izu A, Groth N, et al.
1645 Clinical efficacy of cell culture-derived and egg-derived inactivated subunit influenza vaccines
1646 in healthy adults. *Clinical Infectious Diseases*. 2010;51:997-1004.

1647 [402] Gallagher CM, Goodman MS. Hepatitis B Vaccination of Male Neonates and Autism
1648 Diagnosis, NHIS 1997-2002. *Journal of Toxicology and Environmental Health-Part a-Current*
1649 *Issues*. 2010;73:1665-77.

1650 [403] Garbe E, Andersohn F, Bronder E, Salama A, Klimpel A, Thomae M, et al. Drug-induced
1651 immune thrombocytopaenia: results from the Berlin Case-Control Surveillance Study. *European*
1652 *Journal of Clinical Pharmacology*. 2012;68:821-32.

1653 [404] Gee J, Naleway A, Shui I, Baggs J, Yin R, Li R, et al. Monitoring the safety of
1654 quadrivalent human papillomavirus vaccine: findings from the Vaccine Safety Datalink.
1655 *Vaccine*. 2011;29:8279-84.

1656 [405] Gilbertson DT, Guo H, Arneson TJ, Collins AJ. The association of pneumococcal
1657 vaccination with hospitalization and mortality in hemodialysis patients. *Nephrology Dialysis*
1658 *Transplantation*. 2011;26:2934-9.

1659 [406] Giuliano AR, Palefsky JM, Goldstone S, Moreira ED, Jr., Penny ME, Aranda C, et al.
1660 Efficacy of quadrivalent HPV vaccine against HPV Infection and disease in males.[Erratum
1661 appears in *N Engl J Med*. 2011 Apr 14;364(15):1481]. *New England Journal of Medicine*.
1662 2011;364:401-11.

1663 [407] Glanz JM, Newcomer SR, Hambidge SJ, Daley MF, Narwaney KJ, Xu S, et al. Safety of
1664 trivalent inactivated influenza vaccine in children aged 24 to 59 months in the vaccine safety
1665 datalink. *Archives of Pediatrics & Adolescent Medicine*. 2011;165:749-55.

1666 [408] Gold M, Dugdale S, Woodman RJ, McCaul KA. Use of the Australian Childhood
1667 Immunisation Register for vaccine safety data linkage. *Vaccine*. 2010;28:4308-11.

1668 [409] Gotoh K, Ito Y, Suzuki E, Kaneko K, Kiuchi T, Ando H, et al. Effectiveness and safety of
1669 inactivated influenza vaccination in pediatric liver transplant recipients over three influenza
1670 seasons. *Pediatric Transplantation*. 2011;15:112-6.

1671 [410] Greene SK, Rett M, Weintraub ES, Li L, Yin R, Amato AA, et al. Risk of Confirmed
1672 Guillain-Barre Syndrome Following Receipt of Monovalent Inactivated Influenza A (H1N1) and
1673 Seasonal Influenza Vaccines in the Vaccine Safety Datalink Project, 2009-2010. *American*
1674 *Journal of Epidemiology*. 2012;175:1100-9.

1675 [411] Greenhawt MJ, Spergel JM, Rank MA, Green TD, Masnoor D, Sharma H, et al. Safe
1676 administration of the seasonal trivalent influenza vaccine to children with severe egg allergy.
1677 *Ann Allergy Asthma Immunol*. 2012;109:426-30.

1678 [412] Grimaldi-Bensouda L, Alperovitch A, Besson G, Vial C, Cuisset J-M, Papeix C, et al.
1679 Guillain-Barre syndrome, influenzalike illnesses, and influenza vaccination during seasons with
1680 and without circulating A/H1N1 viruses. *American Journal of Epidemiology*. 2011;174:326-35.

1681 [413] Groves FD, Gridley G, Wacholder S, Shu XO, Robison LL, Neglia JP, et al. Infant
1682 vaccinations and risk of childhood acute lymphoblastic leukaemia in the USA. *Br J Cancer*.
1683 1999;81:175-8.

1684 [414] Gruber C, Warner J, Hill D, Bauchau V, Group ES. Early atopic disease and early
1685 childhood immunization--is there a link? *Allergy*. 2008;63:1464-72.

1686 [415] Gwini SM, Coupland CAC, Siriwardena AN. The effect of influenza vaccination on risk of
1687 acute myocardial infarction: self-controlled case-series study. *Vaccine*. 2011;29:1145-9.

1688 [416] Halasa N, Englund JA, Nachman S, Weinberg GA, Huber VC, Allison K, et al. Safety of
1689 live attenuated influenza vaccine in mild to moderately immunocompromised children with
1690 cancer. *Vaccine*. 2011;29:4110-5.

1691 [417] Hambidge SJ, Ross C, Glanz J, McClure D, Daley MF, Xu S, et al. Trivalent inactivated
1692 influenza vaccine is not associated with sickle cell crises in children. *Pediatrics*. 2012;129:e54-9.

1693 [418] Hambidge SJ, Ross C, McClure D, Glanz J, team VSD. Trivalent inactivated influenza
1694 vaccine is not associated with sickle cell hospitalizations in adults from a large cohort. *Vaccine*.
1695 2011;29:8179-81.

1696 [419] Hedlund J, Christenson B, Lundbergh P, Ortqvist A. Effects of a large-scale intervention
1697 with influenza and 23-valent pneumococcal vaccines in elderly people: A 1-year follow-up.
1698 *Vaccine*. 2003;21:3906-11.

1699 [420] Hummel M, Fuchtenbusch M, Schenker M, Ziegler AG. No major association of breast-
1700 feeding, vaccinations, and childhood viral diseases with early islet autoimmunity in the German
1701 BABYDIAB study. *Diabetes Care*. 2000;23:969-74.

1702 [421] Hurst FP, Lee JJ, Jindal RM, Agodoa LY, Abbott KC. Outcomes associated with influenza
1703 vaccination in the first year after kidney transplantation. *Clin J Am Soc Nephrol*. 2011;6:1192-7.

1704 [422] Huu TN, Toan NT, Tuan HM, Viet HL, Le Thanh Binh P, Yu TW, et al. Safety and
1705 reactogenicity of primary vaccination with the 10-valent pneumococcal non-typeable
1706 *Haemophilus influenzae* protein D conjugate vaccine in Vietnamese infants: a randomised,
1707 controlled trial. *BMC Infectious Diseases*. 2013;13:95.

1708 [423] Iorio A, Basileo M, Marcucci M, Guercini F, Camilloni B, Paccamiccio E, et al. Influenza
1709 vaccination and vitamin K antagonist treatment: a placebo-controlled, randomized, double-blind
1710 crossover study. *Archives of Internal Medicine*. 2010;170:609-16.

1711 [424] Irving SA, Kieke BA, Donahue JG, Mascola MA, Baggs J, DeStefano F, et al. Trivalent
1712 inactivated influenza vaccine and spontaneous abortion. *Obstet Gynecol*. 2013;121:159-65.

1713 [425] Isai A, Durand J, Le Meur S, Hidalgo-Simon A, Kurz X. Autoimmune disorders after
1714 immunisation with Influenza A/H1N1 vaccines with and without adjuvant: EudraVigilance data
1715 and literature review. *Vaccine*. 2012;30:7123-9.

1716 [426] Italian Multicenter Study Group for D, Vaccine Safety in C. Effectiveness and safety of the
1717 A-H1N1 vaccine in children: a hospital-based case-control study. *BMJ Open*. 2011;1:e000167.

1718 [427] Jackson LA, Gaglani MJ, Keyserling HL, Balser J, Bouveret N, Fries L, et al. Safety,
1719 efficacy, and immunogenicity of an inactivated influenza vaccine in healthy adults: a
1720 randomized, placebo-controlled trial over two influenza seasons. *BMC Infectious Diseases*.
1721 2010;10:71.

1722 [428] Johnstone J, Loeb M, Teo KK, Gao P, Dyal L, Liu L, et al. Influenza vaccination and
1723 major adverse vascular events in high risk patients. *Circulation*. 2012;126:21.

1724 [429] Kang S, Kim KH, Kim YT, Kim YH, Song YS, et al. Safety and immunogenicity
1725 of a vaccine targeting human papillomavirus types 6, 11, 16 and 18: a randomized, placebo-
1726 controlled trial in 176 Korean subjects. *Int J Gynecol Cancer*. 2008;18:1013-9.

1727 [430] Kawamura N, Tokoeda Y, Oshima M, Okahata H, Tsutsumi H, Van Doorn LJ, et al.
1728 Efficacy, safety and immunogenicity of RIX4414 in Japanese infants during the first two years
1729 of life. *Vaccine*. 2011;29:6335-41.

1730 [431] Kelly H, Carcione D, Dowse GK, Effler P. The vaccine-attributable risk for febrile
1731 convulsions following influenza vaccine. *Pediatric Infectious Disease Journal*. 2012;31:792.

1732 [432] Kerdpanich A, Chokephaibulkit K, Watanaveeradej V, Vanprapar N, Simasathien S,
1733 Phavichitr N, et al. Immunogenicity of a live-attenuated human rotavirus RIX4414 vaccine with
1734 or without buffering agent. *Human Vaccines*. 2010;6:254-62.

1735 [433] Khalil M, Al-Mazrou Y, Findlow H, Chadha H, Bosch Castells V, Johnson DR, et al.
1736 Safety and immunogenicity of a meningococcal quadrivalent conjugate vaccine in five- to eight-
1737 year-old Saudi Arabian children previously vaccinated with two doses of a meningococcal
1738 quadrivalent polysaccharide vaccine. *Clin Vaccine Immunol*. 2012;19:1561-6.

1739 [434] Khatun S, Akram Hussain SM, Chowdhury S, Ferdous J, Hossain F, Begum SR, et al.
1740 Safety and immunogenicity profile of human papillomavirus-16/18 AS04 adjuvant cervical
1741 cancer vaccine: a randomized controlled trial in healthy adolescent girls of Bangladesh. *Japanese*
1742 *Journal of Clinical Oncology*. 2012;42:36-41.

1743 [435] Kim DS, Lee TJ, Kang JH, Kim J-H, Lee JH, Ma SH, et al. Immunogenicity and safety of
1744 a pentavalent human-bovine (WC3) reassortant rotavirus vaccine in healthy infants in Korea.
1745 *Pediatric Infectious Disease Journal*. 2008;27:177-8.

1746 [436] Kim JS, Bae CW, Lee KY, Park MS, Choi YY, Kim KN, et al. Immunogenicity,
1747 reactogenicity and safety of a human rotavirus vaccine (RIX4414) in Korean infants: A
1748 randomized, double-blind, placebo-controlled, phase IV study. *Human Vaccines and*
1749 *Immunotherapeutics*. 2012;8:806-12.

1750 [437] Kim SC, Song YS, Kim Y-T, Kim YT, Ryu K-S, Gunapalaiah B, et al. Human
1751 papillomavirus 16/18 AS04-adjuvanted cervical cancer vaccine: immunogenicity and safety in
1752 15-25 years old healthy Korean women. *Journal of Gynecologic Oncology*. 2011;22:67-75.

1753 [438] Klein NP, Aukes L, Lee J, Fireman B, Shapira SK, Slade B, et al. Evaluation of
1754 immunization rates and safety among children with inborn errors of metabolism. *Pediatrics*.
1755 2011;127:e1139-e46.

1756 [439] Klein NP, Hansen J, Chao C, Velicer C, Emery M, Slezak J, et al. Safety of quadrivalent
1757 human papillomavirus vaccine administered routinely to females. *Archives of Pediatrics &*
1758 *Adolescent Medicine*. 2012;166:1140-8.

1759 [440] Klein NP, Massolo ML, Greene J, Dekker CL, Black S, Escobar GJ. Risk factors for
1760 developing apnea after immunization in the neonatal intensive care unit. *Pediatrics*.
1761 2008;121:463-9.

1762 [441] Klein NP, Reisinger KS, Johnston W, Odrlic T, Gill CJ, Bedell L, et al. Safety and
1763 immunogenicity of a novel quadrivalent meningococcal CRM-conjugate vaccine given
1764 concomitantly with routine vaccinations in infants.[Erratum appears in *Pediatr Infect Dis J*. 2012
1765 Oct;31(10):1105]. *Pediatric Infectious Disease Journal*. 2012;31:64-71.

1766 [442] Langley JM, Aoki F, Ward BJ, McGeer A, Angel JB, Stiver G, et al. A nasally
1767 administered trivalent inactivated influenza vaccine is well tolerated, stimulates both mucosal
1768 and systemic immunity, and potentially protects against influenza illness. *Vaccine*.
1769 2011;29:1921-8.

1770 [443] Lee GM, Greene SK, Weintraub ES, Baggs J, Kulldorff M, Fireman BH, et al. H1N1 and
1771 seasonal influenza vaccine safety in the vaccine safety datalink project. *American Journal of*
1772 *Preventive Medicine*. 2011;41:121-8.

1773 [444] Lee S, Park WB, Shin K-H, Ahn DH, Yoon SH, Cho J-Y, et al. Immunogenicity and safety
1774 of a single intramuscular dose of a diphtheria-tetanus toxoid (Td) vaccine (GC1107) in Korean
1775 adults. *Vaccine*. 2011;29:7638-43.

1776 [445] Levin MJ, Moscicki A-B, Song L-Y, Fenton T, Meyer WA, 3rd, Read JS, et al. Safety and
1777 immunogenicity of a quadrivalent human papillomavirus (types 6, 11, 16, and 18) vaccine in
1778 HIV-infected children 7 to 12 years old. *J Acquir Immune Defic Syndr*. 2010;55:197-204.

1779 [446] Li R, Li Y, Radley D, Liu Y, Huang T, Singhs HL, et al. Safety and immunogenicity of a
1780 vaccine targeting human papillomavirus types 6, 11, 16 and 18: a randomized, double-blind,
1781 placebo-controlled trial in Chinese males and females. *Vaccine*. 2012;30:4284-91.

1782 [447] Lin TH, Lin SY, Lin CH, Lin RI, Lin HC, Chiu TH, et al. AdimFlu-S((registered
1783 trademark)) influenza A (H1N1) vaccine during pregnancy: The Taiwanese Pharmacovigilance
1784 Survey. *Vaccine*. 2012;30:2671-5.

1785 [448] Ma X, Does MB, Metayer C, Russo C, Wong A, Buffler PA. Vaccination history and risk
1786 of childhood leukaemia. *Int J Epidemiol*. 2005;34:1100-9.

1787 [449] Macaladad N, Marcano T, Guzman M, Moya J, Jurado F, Thompson M, et al. Safety and
1788 immunogenicity of a zoster vaccine in varicella-zoster virus seronegative and low-seropositive
1789 healthy adults. *Vaccine*. 2007;25:2139-44.

1790 [450] MacArthur AC, McBride ML, Spinelli JJ, Tamaro S, Gallagher RP, Theriault GP. Risk of
1791 childhood leukemia associated with vaccination, infection, and medication use in childhood: the
1792 Cross-Canada Childhood Leukemia Study. *Am J Epidemiol*. 2008;167:598-606.

1793 [451] Madhi SA, Cunliffe NA, Steele D, Witte D, Kirsten M, Louw C, et al. Effect of human
1794 rotavirus vaccine on severe diarrhea in African infants. *New England Journal of Medicine*.
1795 2010;362:289-98.

1796 [452] Madhi SA, Dittmer S, Kuwanda L, Venter M, Cassim H, Lazarus E, et al. Efficacy and
1797 immunogenicity of influenza vaccine in HIV-infected children: a randomized, double-blind,
1798 placebo controlled trial. *Aids*. 2013;27:369-79.

1799 [453] Madhi SA, Maskew M, Koen A, Kuwanda L, Besselaar TG, Naidoo D, et al. Trivalent
1800 inactivated influenza vaccine in African adults infected with human immunodeficient virus:
1801 double blind, randomized clinical trial of efficacy, immunogenicity, and safety. *Clinical*
1802 *Infectious Diseases*. 2011;52:128-37.

1803 [454] Mallory RM, Malkin E, Ambrose CS, Bellamy T, Shi L, Yi T, et al. Safety and
1804 Immunogenicity Following Administration of a Live, Attenuated Monovalent 2009 H1N1
1805 Influenza Vaccine to Children and Adults in Two Randomized Controlled Trials. *PLoS ONE*
1806 [Electronic Resource]. 2010;5:e13755.

1807 [455] Matheson MC, Haydn Walters E, Burgess JA, Jenkins MA, Giles GG, Hopper JL, et al.
1808 Childhood immunization and atopic disease into middle-age--a prospective cohort study. *Pediatr*
1809 *Allergy Immunol*. 2010;21:301-6.

1810 [456] Mills R, Tying SK, Levin MJ, Parrino J, Li X, Coll KE, et al. Safety, tolerability, and
1811 immunogenicity of zoster vaccine in subjects with a history of herpes zoster. *Vaccine*.
1812 2010;28:4204-9.

1813 [457] Mommers M, Weishoff-Houben M, Swaen GM, Creemers H, Freund H, Dott W, et al.
1814 Infant immunization and the occurrence of atopic disease in Dutch and German children: a
1815 nested case-control study. *Pediatr Pulmonol*. 2004;38:329-34.

1816 [458] Moreira Jr ED, Palefsky JM, Giuliano AR, Goldstone S, Aranda C, Jessen H, et al. Safety
1817 and reactogenicity of a quadrivalent human papillomavirus (types 6, 11, 16, 18) L1 viral-like-
1818 particle vaccine in older adolescents and young adults. *Human Vaccines*. 2011;7:768-75.

1819 [459] Morgan TM, Schlegel C, Edwards KM, Welch-Burke T, Zhu Y, Sparks R, et al. Vaccines
1820 are not associated with metabolic events in children with urea cycle disorders. *Pediatrics*.
1821 2011;127:e1147-e53.

1822 [460] Mullooly JP, Schuler R, Barrett M, Maher JE. Vaccines, antibiotics, and atopy.
1823 *Pharmacoepidemiology and Drug Safety*. 2007;16:275-88.

1824 [461] Mullooly JP, Schuler R, Mesa J, Drew L, DeStefano F, team VSD. Wheezing lower
1825 respiratory disease and vaccination of premature infants. *Vaccine*. 2011;29:7611-7.

1826 [462] Murray AV, Reisinger KS, Kerzner B, Stek JE, Sausser TA, Xu J, et al. Safety and
1827 tolerability of zoster vaccine in adults ≥ 60 years old. *Human Vaccines*. 2011;7:1130-6.

1828 [463] Nakajima K, Dharmage SC, Carlin JB, Wharton CL, Jenkins MA, Giles GG, et al. Is
1829 childhood immunisation associated with atopic disease from age 7 to 32 years? *Thorax*.
1830 2007;62:270-5.

1831 [464] Narang A, Bose A, Pandit AN, Dutta P, Kang G, Bhattacharya SK, et al. Immunogenicity,
1832 reactogenicity and safety of human rotavirus vaccine (RIX4414) in Indian infants. *Human*
1833 *Vaccines*. 2009;5:414-9.

1834 [465] Ngan HYS, Cheung ANY, Tam KF, Chan KKL, Tang HW, Bi D, et al. Human
1835 papillomavirus-16/18 AS04-adjuvanted cervical cancer vaccine: Immunogenicity and safety in
1836 healthy Chinese women from Hong Kong. *Hong Kong Medical Journal*. 2010;16:171-9.

1837 [466] Nordin JD, Kharbanda EO, Benitez GV, Nichol K, Lipkind H, Naleway A, et al. Maternal
1838 safety of trivalent inactivated influenza vaccine in pregnant women. *Obstet Gynecol*.
1839 2013;121:519-25.

1840 [467] O'Leary ST, Glanz JM, McClure DL, Akhtar A, Daley MF, Nakasato C, et al. The risk of
1841 immune thrombocytopenic purpura after vaccination in children and adolescents. *Pediatrics*.
1842 2012;129:248-55.

1843 [468] Omenaca F, Sarlangue J, Szenborn L, Nogueira M, Suryakiran PV, Smolenov IV, et al.
1844 Safety, reactogenicity and immunogenicity of the human rotavirus vaccine in preterm European
1845 Infants: a randomized phase IIIb study. *Pediatric Infectious Disease Journal*. 2012;31:487-93.

1846 [469] Omer SB, Goodman D, Steinhoff MC, Rochat R, Klugman KP, Stoll BJ, et al. Maternal
1847 influenza immunization and reduced likelihood of prematurity and small for gestational age
1848 births: a retrospective cohort study. *PLoS Med*. 2011;8:e1000441.

1849 [470] Pagaoa MA, Okcu MF, Bondy ML, Scheurer ME. Associations between vaccination and
1850 childhood cancers in Texas regions. *Journal of Pediatrics*. 2011;158:996-1002.

1851 [471] Pahud BA, Rowhani-Rahbar A, Glaser C, Gavali S, Salibay CJ, Fireman B, et al. Lack of
1852 association between childhood immunizations and encephalitis in California, 1998-2008.
1853 *Vaccine*. 2012;30:247-53.

1854 [472] Patel MM, Lopez-Collada VR, Bulhoes MM, De Oliveira LH, Bautista Marquez A,
1855 Flannery B, et al. Intussusception risk and health benefits of rotavirus vaccination in Mexico and
1856 Brazil. *New England Journal of Medicine*. 2011;364:2283-92.

1857 [473] Phua KB, Lim FS, Lau YL, Nelson EAS, Huang LM, Quak SH, et al. Rotavirus vaccine
1858 RIX4414 efficacy sustained during the third year of life: A randomized clinical trial in an Asian
1859 population. *Vaccine*. 2012;30:4552-7.

1860 [474] Phua KB, Quak SH, Lee BW, Emmanuel SC, Goh P, Han HH, et al. Evaluation of
1861 RIX4414, A live, attenuated rotavirus vaccine, in a randomized, double-blind, placebo-controlled
1862 phase 2 trial involving 2464 Singaporean infants. *Journal of Infectious Diseases*. 2005;192:S6-
1863 S16.

1864 [475] Ray P, Black S, Shinefield H, Dillon A, Carpenter D, Lewis E, et al. Risk of rheumatoid
1865 arthritis following vaccination with tetanus, influenza and hepatitis B vaccines among persons
1866 15-59 years of age. *Vaccine*. 2011;29:6592-7.

1867 [476] Richards JL, Hansen C, Bredfeldt C, Bednarczyk RA, Steinhoff MC, Adjaye-Gbewonyo
1868 D, et al. Neonatal outcomes after antenatal influenza immunization during the 2009 H1N1
1869 influenza pandemic: impact on preterm birth, birth weight, and small for gestational age birth.
1870 *Clinical Infectious Diseases*. 2013;56:1216-22.

1871 [477] Roteli-Martins CM, Naud P, De Borba P, Teixeira JC, De Carvalho NS, Zahaf T, et al.
1872 Sustained immunogenicity and efficacy of the HPV-16/18 AS04-adjuvanted vaccine: Up to 8.4
1873 years of follow-up. *Human Vaccines and Immunotherapeutics*. 2012;8:381-8.

1874 [478] Rowhani-Rahbar A, Klein NP, Lewis N, Fireman B, Ray P, Rasgon B, et al. Immunization
1875 and bell's palsy in children: A case-centered analysis. *American Journal of Epidemiology*.
1876 2012;175:878-85.

1877 [479] Ruiz-Palacios GM, Perez-Schael I, Velazquez FR, Abate H, Breuer T, Clemens SC, et al.
1878 Safety and efficacy of an attenuated vaccine against severe rotavirus gastroenteritis. *New*
1879 *England Journal of Medicine*. 2006;354:11-22.

1880 [480] Santosham M, Wolff M, Reid R, Hohenboken M, Bateman M, Goepf J, et al. The efficacy
1881 in Navajo infants of a conjugate vaccine consisting of Haemophilus influenzae type b
1882 polysaccharide and Neisseria meningitidis outer-membrane protein complex. *New England*
1883 *Journal of Medicine*. 1991;324:1767-72.

1884 [481] Schmader KE, Levin MJ, Gnann JW, Jr., McNeil SA, Vesikari T, Betts RF, et al. Efficacy,
1885 safety, and tolerability of herpes zoster vaccine in persons aged 50-59 years. *Clinical Infectious*
1886 *Diseases*. 2012;54:922-8.

1887 [482] Schwarz TF, Huang LM, Medina DM, Valencia A, Lin TY, Behre U, et al. Four-year
1888 follow-up of the immunogenicity and safety of the HPV-16/18 AS04-adjuvanted vaccine when
1889 administered to adolescent girls aged 10-14 years. *Journal of Adolescent Health*. 2012;50:187-
1890 94.

1891 [483] Shui IM, Baggs J, Patel M, Parashar UD, Rett M, Belongia EA, et al. Risk of
1892 intussusception following administration of a pentavalent rotavirus vaccine in US infants.
1893 *JAMA*. 2012;307:598-604.

1894 [484] Siberry GK, Williams PL, Lujan-Zilbermann J, Warshaw MG, Spector SA, Decker MD, et
1895 al. Phase I/II, open-label trial of safety and immunogenicity of meningococcal (groups A, C, Y,
1896 and W-135) polysaccharide diphtheria toxoid conjugate vaccine in human immunodeficiency
1897 virus-infected adolescents. *Pediatric Infectious Disease Journal*. 2010;29:391-6.

1898 [485] Simberkoff MS, Arbeit RD, Johnson GR, Oxman MN, Boardman KD, Williams HM, et al.
1899 Safety of herpes zoster vaccine in the shingles prevention study: a randomized trial. *Annals of*
1900 *Internal Medicine*. 2010;152:545-54.

1901 [486] Siriwardena AN, Gwini SM, Coupland CAC. Influenza vaccination, pneumococcal
1902 vaccination and risk of acute myocardial infarction: matched case-control study. *CMAJ*
1903 *Canadian Medical Association Journal*. 2010;182:1617-23.

1904 [487] Sow PS, Watson-Jones D, Kiviat N, Changalucha J, Mbaye KD, Brown J, et al. Safety and
1905 immunogenicity of human papillomavirus-16/18 AS04-adjuvanted vaccine: a randomized trial in
1906 10-25-year-old HIV-Seronegative African girls and young women. *Journal of Infectious*
1907 *Diseases*. 2013;207:1753-63.

1908 [488] Steele AD, Reynders J, Scholtz F, Bos P, de Beer MC, Tumbo J, et al. Comparison of 2
1909 different regimens for reactogenicity, safety, and immunogenicity of the live attenuated oral
1910 rotavirus vaccine RIX4414 coadministered with oral polio vaccine in South African infants.
1911 *Journal of Infectious Diseases*. 2010;202 Suppl:S93-100.

1912 [489] Stowe J, Andrews N, Bryan P, Seabroke S, Miller E. Risk of convulsions in children after
1913 monovalent H1N1 (2009) and trivalent influenza vaccines: A database study. *Vaccine*.
1914 2011;29:9467-72.

1915 [490] Sun YL, Christensen J, Hviid A, Li J, Vedsted P, Olsen J, et al. Risk of Febrile Seizures
1916 and Epilepsy After Vaccination With Diphtheria, Tetanus, Acellular Pertussis, Inactivated
1917 Poliovirus, and Haemophilus Influenzae Type b. *Jama-Journal of the American Medical*
1918 *Association*. 2012;307:823-31.

1919 [491] Talaat KR, Greenberg ME, Lai MH, Hartel GF, Wichems CH, Rockman S, et al. A single
1920 dose of unadjuvanted novel 2009 H1N1 vaccine is immunogenic and well tolerated in young and
1921 elderly adults. *Journal of Infectious Diseases*. 2010;202:1327-37.

1922 [492] Tanner K, Roy N, Merrill RM, Sauder C, Houtz DR, Smith ME. Case-control study of risk
1923 factors for spasmodic dysphonia: A comparison with other voice disorders. *Laryngoscope*.
1924 2012;122:1082-92.

1925 [493] Thomson JA, Widjaja C, Darmaputra AAP, Lowe A, Matheson MC, Bennett CM, et al.
1926 Early childhood infections and immunisation and the development of allergic disease in
1927 particular asthma in a high-risk cohort: A prospective study of allergy-prone children from birth
1928 to six years. *Pediatr Allergy Immunol*. 2010;21:1076-85.

1929 [494] Ting SCH, Crooks SW, South G. The effect of influenza vaccination on the incidence of
1930 chronic obstructive pulmonary disease exacerbations in the immediate postvaccination period. *J*
1931 *Epidemiol Community Health*. 2011;65:157-9.

1932 [495] Tokars JI, Lewis P, DeStefano F, Wise M, Viray M, Morgan O, et al. The risk of Guillain-
1933 Barre syndrome associated with influenza A (H1N1) 2009 monovalent vaccine and 2009-2010
1934 seasonal influenza vaccines: results from self-controlled analyses. *Pharmacoepidemiol Drug Saf*.
1935 2012;21:546-52.

1936 [496] Treanor JJ, El Sahly H, King J, Graham I, Izikson R, Kohberger R, et al. Protective
1937 efficacy of a trivalent recombinant hemagglutinin protein vaccine (FluBlok[REGISTERED])
1938 against influenza in healthy adults: a randomized, placebo-controlled trial. *Vaccine*.
1939 2011;29:7733-9.

1940 [497] Tse A, Tseng HF, Greene SK, Vellozzi C, Lee GM, Group VSDRCAIW. Signal
1941 identification and evaluation for risk of febrile seizures in children following trivalent inactivated
1942 influenza vaccine in the Vaccine Safety Datalink Project, 2010-2011. *Vaccine*. 2012;30:2024-31.

1943 [498] Tseng HF, Liu A, Sy L, Marcy SM, Fireman B, Weintraub E, et al. Safety of zoster
1944 vaccine in adults from a large managed-care cohort: a Vaccine Safety Datalink study. *J Intern*
1945 *Med*. 2012;271:510-20.

1946 [499] Tseng HF, Slezak JM, Quinn VP, Sy LS, Van den Eeden SK, Jacobsen SJ. Pneumococcal
1947 vaccination and risk of acute myocardial infarction and stroke in men. *JAMA*. 2010;303:1699-
1948 706.

1949 [500] Uno Y, Uchiyama T, Kurosawa M, Aleksic B, Ozaki N. The combined measles, mumps,
1950 and rubella vaccines and the total number of vaccines are not associated with development of
1951 autism spectrum disorder: The first case-control study in Asia. *Vaccine*. 2012;30:4292-8.

1952 [501] Velazquez FR, Colindres RE, Grajales C, Hernandez MT, Mercadillo MG, Torres FJ, et al.
1953 Postmarketing surveillance of intussusception following mass introduction of the attenuated
1954 human rotavirus vaccine in Mexico. *Pediatric Infectious Disease Journal*. 2012;31:736-44.

1955 [502] Velentgas P, Amato AA, Bohn RL, Chan KA, Cochrane T, Funch DP, et al. Risk of
1956 Guillain-Barre syndrome after meningococcal conjugate vaccination. *Pharmacoepidemiol Drug*
1957 *Saf*. 2012;21:1350-8.

1958 [503] Vermeulen JN, Lange JMA, Tying SK, Peters PH, Nunez M, Poland G, et al. Safety,
1959 tolerability, and immunogenicity after 1 and 2 doses of zoster vaccine in healthy adults ≥ 60
1960 years of age. *Vaccine*. 2012;30:904-10.

1961 [504] Vesikari T, Clark HF, Offit PA, Dallas MJ, DiStefano DJ, Goveia MG, et al. Effects of the
1962 potency and composition of the multivalent human-bovine (WC3) reassortant rotavirus vaccine
1963 on efficacy, safety and immunogenicity in healthy infants. *Vaccine*. 2006;24:4821-9.

1964 [505] Vesikari T, Karvonen A, Bouckenoghe A, Suryakiran PV, Smolenov I, Han HH.
1965 Immunogenicity, reactogenicity and safety of the human rotavirus vaccine RIX4414 oral
1966 suspension (liquid formulation) in Finnish infants. *Vaccine*. 2011;29:2079-84.

1967 [506] Vesikari T, Karvonen A, Korhonen T, Espo M, Lebacqz E, Forster J, et al. Safety and
1968 immunogenicity of RIX4414 live attenuated human rotavirus vaccine in adults, toddlers and
1969 previously uninfected infants. *Vaccine*. 2004;22:2836-42.

1970 [507] Vesikari T, Karvonen A, Puustinen L, Zeng S-Q, Szakal ED, Delem A, et al. Efficacy of
1971 RIX4414 live attenuated human rotavirus vaccine in Finnish infants. *Pediatric Infectious Disease*
1972 *Journal*. 2004;23:937-43.

1973 [508] Vesikari T, Matson DO, Dennehy P, Van Damme P, Santosham M, Rodriguez Z, et al.
1974 Safety and efficacy of a pentavalent human-bovine (WC3) reassortant rotavirus vaccine. *New*
1975 *England Journal of Medicine*. 2006;354:23-33.

1976 [509] Wang IK, Lin CL, Lin PC, Liang CC, Liu YL, Chang CT, et al. Effectiveness of influenza
1977 vaccination in patients with end-stage renal disease receiving hemodialysis: a population-based
1978 study. *PLoS ONE [Electronic Resource]*. 2013;8:e58317.

1979 [510] Weinberg A, Levin MJ, Macgregor RR. Safety and immunogenicity of a live attenuated
1980 varicella vaccine in VZV-seropositive HIV-infected adults. *Human Vaccines*. 2010;6:318-21.

1981 [511] Whitehouse AJO, Maybery M, Wray JA, Hickey M. No association between early
1982 gastrointestinal problems and autistic-like traits in the general population. *Dev Med Child*
1983 *Neurol*. 2011;53:457-62.

1984 [512] Wilson K, Hawken S, Kwong JC, Deeks S, Crowcroft NS, Van Walraven C, et al. Adverse
1985 events following 12 and 18 month vaccinations: a population-based, self-controlled case series
1986 analysis. *PLoS ONE [Electronic Resource]*. 2011;6:e27897.

1987 [513] Wise ME, Viray M, Sejvar JJ, Lewis P, Baughman AL, Connor W, et al. Guillain-Barre
1988 Syndrome During the 2009-2010 H1N1 Influenza Vaccination Campaign: Population-based
1989 Surveillance Among 45 Million Americans. *American Journal of Epidemiology*. 2012;175:1110-
1990 9.

1991 [514] Xu R, Luo Y, Chambers C. Assessing the effect of vaccine on spontaneous abortion using
1992 time-dependent covariates Cox models. *Pharmacoepidemiology and Drug Safety*. 2012.

1993 [515] Yih WK, Lee GM, Lieu TA, Ball R, Kulldorff M, Rett M, et al. Surveillance for Adverse
1994 Events Following Receipt of Pandemic 2009 H1N1 Vaccine in the Post-Licensure Rapid
1995 Immunization Safety Monitoring (PRISM) System, 2009-2010. *American Journal of*
1996 *Epidemiology*. 2012;175:1120-8.

1997 [516] Yu O, Bohlke K, Hanson CA, Delaney K, Rees TG, Zavitkovsky A, et al. Hepatitis B
1998 vaccine and risk of autoimmune thyroid disease: A Vaccine Safety Datalink study.
1999 *Pharmacoepidemiology and Drug Safety*. 2007;16:736-45.

2000 [517] Zaman K, Sack DA, Yunus M, Arifeen SE, Podder G, Azim T, et al. Successful co-
2001 administration of a human rotavirus and oral poliovirus vaccines in Bangladeshi infants in a 2-
2002 dose schedule at 12 and 16 weeks of age. *Vaccine*. 2009;27:1333-9.

2003 [518] Zhang J, Xie F, Delzell E, Chen L, Winthrop KL, Lewis JD, et al. Association between
2004 vaccination for herpes zoster and risk of herpes zoster infection among older patients with
2005 selected immune-mediated diseases. JAMA. 2012;308:43-9.

2006 [519] Christie CDC, Duncan ND, Thame KA, Onorato MT, Smith HD, Malcolm LG, et al.
2007 Pentavalent rotavirus vaccine in developing countries: safety and health care resource utilization.
2008 Pediatrics. 2010;126:e1499-506.

2009 [520] EU/EEA EO. Study to evaluate the efficacy, safety and immunogenicity of two or three
2010 doses of GSK Biologicals' oral live attenuated human rotavirus (HRV) vaccine given
2011 concomitantly with routine EPI vaccinations in healthy infants.
2012 <http://www.who.int/trialsearch/Trial2.aspx?TrialID=EUCTR2015-001485-26-Outside-EU/EEA>.
2013 2015.

2014 [521] EU/EEA EO. A phase III study to assess the efficacy, immunogenicity and safety of GSK
2015 Biologicals' human rotavirus (HRV) vaccine given concomitantly with routine expanded
2016 program on immunisation (EPI) vaccinations including oral poliovirus vaccine (OPV) in healthy
2017 infants across 6 countries in Latin America.
2018 <http://www.who.int/trialsearch/Trial2.aspx?TrialID=EUCTR2015-001540-10-Outside-EU/EEA>.
2019 2015.

2020 [522] Euctr Outside EU/EEA. A phase II study to assess the safety and immunogenicity of
2021 GlaxoSmithKline Biologicals' rotavirus vaccine, RIX4414 when administered to HIV infected
2022 infants in South Africa. [http://www.who.int/trialsearch/Trial2.aspx?TrialID=EUCTR2015-001484-](http://www.who.int/trialsearch/Trial2.aspx?TrialID=EUCTR2015-001484-39-Outside-EU/EEA)
2023 [39-Outside-EU/EEA](http://www.who.int/trialsearch/Trial2.aspx?TrialID=EUCTR2015-001484-39-Outside-EU/EEA). 2015.

2024 [523] Euctr Outside EU/EEA. Study to evaluate the efficacy and safety of two doses of GSK
2025 Biologicals' oral live attenuated human rotavirus (HRV) vaccine in healthy infants. 2015.

2026 [524] Euctr Outside EU/EEA. Study to evaluate immunogenicity, reactogenicity and safety of
2027 Rotarix™ vaccine in Korean infants.
2028 <http://www.who.int/trialsearch/Trial2.aspx?TrialID=EUCTR2015-001545-81-Outside-EU/EEA>.
2029 2015.

2030 [525] Euctr Outside EU/EEA. Study to evaluate the efficacy and safety of two doses of GSK
2031 Biologicals' oral live attenuated human rotavirus (HRV) vaccine in healthy infants.
2032 <http://www.who.int/trialsearch/Trial2.aspx?TrialID=EUCTR2015-001541-92-Outside-EU/EEA>.
2033 2015.

2034 [526] GlaxoSmithKline. A Study to Test 2 Doses of GSK Biologicals' Oral Live Attenuated
2035 Human Rotavirus (HRV) Vaccine in Healthy Infants.
2036 <https://ClinicalTrials.gov/show/NCT00197210>; 2003.

2037 [527] GlaxoSmithKline. Study of 2 Doses of HRV Vaccine Given Concomitantly With Routine
2038 EPI Vaccinations Including OPV in Healthy Infants. 2003.

2039 [528] GlaxoSmithKline. Year 3 Extension for Efficacy Follow-up in Subjects Vaccinated in
2040 Studies Rota-028, 029 or 030 (NCT00197210). <https://ClinicalTrials.gov/show/NCT00329745>;
2041 2007.

2042 [529] GlaxoSmithKline Biologicals. **A phase III, double-blind, randomized, placebo-**
2043 **controlled, multi-country and multi-center study to assess the efficacy and safety of two**
2044 **doses of GSK Biologicals' oral live attenuated human rotavirus (HRV) vaccine in healthy**
2045 **infants** 2015.

2046 [530] GlaxoSmithKline Biologicals. Study to evaluate immunogenicity, reactogenicity and
2047 safety of Rotarix™ vaccine in Korean infants.

2048 <http://www.who.int/trialsearch/Trial2.aspx?TrialID=EUCTR2015-001545-81-Outside-EU/EEA>.
2049 2015.

2050 [531] Goveia MG, Rodriguez ZM, Dallas MJ, Itzler RF, Boslego JW, Heaton PM, et al. Safety
2051 and efficacy of the pentavalent human-bovine (WC3) reassortant rotavirus vaccine in healthy
2052 premature infants. *Pediatric Infectious Disease Journal*. 2007;26:1099-104.

2053 [532] Grant LR, Watt JP, Weatherholtz RC, Moulton LH, Reid R, Santosham M, et al. Efficacy
2054 of a pentavalent human-bovine reassortant rotavirus vaccine against rotavirus gastroenteritis
2055 among American Indian children. *Pediatric Infectious Disease Journal*. 2012;31:184-8.

2056 [533] Guillermo M, et al. RIX4414 (Rotarix™) has demonstrated efficacy during the first 2 years
2057 of life in infants from 11 Latin American countries. 10th International Rotavirus Symposium
2058 (IRS). Bangkok, Thailand2012.

2059 [534] Hemming-Harlo M, Lahdeaho M-L, Maki M, Vesikari T. Rotavirus Vaccination Does Not
2060 Increase Type 1 Diabetes and May Decrease Celiac Disease in Children and Adolescents. *The*
2061 *Pediatric infectious disease journal*. 2019;38:539-41.

2062 [535] Laserson KF, Nyakundi D, Feikin DR, Nyambane G, Cook E, Oyieko J, et al. Safety of the
2063 pentavalent rotavirus vaccine (PRV), RotaTeq([REGISTERED]), in Kenya, including among
2064 HIV-infected and HIV-exposed infants. *Vaccine*. 2012;30 Suppl 1:A61-70.

2065 [536] Lau YL, Nelson EA, Poon KH, Chan PK, Chiu S, Sung R, et al. Efficacy, safety and
2066 immunogenicity of a human rotavirus vaccine (RIX4414) in Hong Kong children up to three
2067 years of age: a randomized, controlled trial. *Vaccine*. 2013;31:2253-9.

2068 [537] Lee KJ, Carlin JB. Fractional polynomial adjustment for time-varying covariates in a self-
2069 controlled case series analysis. *Stat Med*. 2014;33:105-16.

2070 [538] Oxman MN, Levin MJ, Johnson GR, Schmader KE, Straus SE, Gelb LD, et al. A vaccine
2071 to prevent herpes zoster and postherpetic neuralgia in older adults. *N Engl J Med*.
2072 2005;352:2271-84.

2073 [539] Phua KB, et al. Human rotavirus vaccine RIX4414 (Rotarix™) is highly efficacious in
2074 infants from Asia during the first two years of life. 13th International Congress on Infectious
2075 Diseases (ICID). Kuala Lumpur, Malaysia2008.

2076 [540] Phua KB, et al. Human rotavirus vaccine RIX4414 (Rotarix™) is highly efficacious in
2077 Asian infants during the first three years of life. 13th Asian Pacific Congress of Pediatrics
2078 (APCP). Shanghai, China2009.

2079 [541] Phua KB, et al. Human rotavirus vaccine RIX4414 is highly efficacious in Asian infants
2080 during the third year of life. 27th Annual Meeting of the European Society for Paediatric
2081 Infectious Diseases (ESPID). Brussels, Belgium2009.

2082 [542] Phua KB, et al. Efficacy of rotavirus vaccine RIX4414 during the first 3 years of life: a
2083 randomised, double-blind, placebo-controlled study in infants from Hong Kong, Singapore and
2084 Taiwan. 10th International Rotavirus Symposium (IRS). Bangkok, Thailand2012.

2085 [543] Phua KB, Lim FS, Lau YL, Nelson EAS, Huang LM, Quak SH, et al. Safety and efficacy
2086 of human rotavirus vaccine during the first 2 years of life in Asian infants: randomised, double-
2087 blind, controlled study. *Vaccine*. 2009;27:5936-41.

2088 [544] Popmihajlov Z, Pang L, Brown E, Joshi A, Su SC, Kaplan SS, et al. A post hoc analysis
2089 utilizing the FDA toxicity grading scale to assess injection site adverse events following
2090 immunization with the live attenuated Zoster Vaccine (ZVL). *Human Vaccines and*
2091 *Immunotherapeutics*. 2018;14:2916-20.

2092 [545] Rodriguez ZM, Goveia MG, Stek JE, Dallas MJ, Boslego JW, DiNubile MJ, et al.
2093 Concomitant use of an oral live pentavalent human-bovine reassortant rotavirus vaccine with

2094 licensed parenteral pediatric vaccines in the United States. *Pediatric Infectious Disease Journal*.
2095 2007;26:221-7.

2096 [546] Sow SO, Tapia M, Haidara FC, Ciarlet M, Diallo F, Kodio M, et al. Efficacy of the oral
2097 pentavalent rotavirus vaccine in Mali. *Vaccine*. 2012;30 Suppl 1:A71-8.

2098 [547] Tregnaghi MW, Abate HJ, Valencia A, Lopez P, Da Silveira TR, Rivera L, et al. Human
2099 rotavirus vaccine is highly efficacious when coadministered with routine expanded program of
2100 immunization vaccines including oral poliovirus vaccine in Latin America. *Pediatric Infectious
2101 Disease Journal*. 2011;30:e103-e8.

2102 [548] Zaman K, Dang DA, Victor JC, Shin S, Yunus M, Dallas MJ, et al. Efficacy of pentavalent
2103 rotavirus vaccine against severe rotavirus gastroenteritis in infants in developing countries in
2104 Asia: a randomised, double-blind, placebo-controlled trial. *Lancet*. 2010;376:615-23.

2105 [549] Zaman K, Yunus M, El Arifeen S, Azim T, Faruque ASG, Huq E, et al. Methodology and
2106 lessons-learned from the efficacy clinical trial of the pentavalent rotavirus vaccine in
2107 Bangladesh. *Vaccine*. 2012;30 Suppl 1:A94-100.

2108 [550] Dudley MZ, Halsey NA, Omer SB, Orenstein WA, O'Leary ST, Limaye RJ, et al. The state
2109 of vaccine safety science: systematic reviews of the evidence. *Lancet Infect Dis*. 2020;20:e80-e9.

2110 [551] Lu H-L, Ding Y, Goyal H, Xu H-G. Association Between Rotavirus Vaccination and Risk
2111 of Intussusception Among Neonates and Infants: A Systematic Review and Meta-analysis.
2112 *JAMA network open*. 2019;2:e1912458.

2113 [552] Health Resources and Services Administration. *Vaccine Injury Table*.

2114 [553] Hartung DM, Zarin DA, Guise JM, McDonagh M, Paynter R, Helfand M. Reporting
2115 discrepancies between the ClinicalTrials.gov results database and peer-reviewed publications.
2116 *Ann Intern Med*. 2014;160:477-83.

2117 [554] Tang E, Ravaud P, Riveros C, Perrodeau E, Dechartres A. Comparison of serious adverse
2118 events posted at ClinicalTrials.gov and published in corresponding journal articles. *BMC Med*.
2119 2015;13:189.

2120 [555] HealthIT.gov. *Interoperability Proving Ground*.

2121

Table 1. Included vaccines, populations, and recent changes (within five years)

Vaccine (abbreviation; brand name)	Populations recommended for routine use	Recent changes to formulation, age indication, or dosing (within last five years)
9-valent human papillomavirus (HPV9; Gardasil 9®)	Adults, children	Gardasil 9 approval expanded to include use in women and men 27 through 45 years of age in 2018. Gardasil 9 approved as a two-dose series if first dose initiated 9-14 years of age (otherwise three-dose series as before) in 2016. Catch-up HPV vaccination recommended for all persons through age 26 years in 2019.
13-valent pneumococcal conjugate (PCV13; Prevnar 13®)	Adults, children	Age indications were expanded from younger than 18 years and older than 50 years to include adults aged 18-49 years in 2016.
23-valent pneumococcal polysaccharide (PPSV23; Pneumovax®)	Adults, children	None
Diphtheria, tetanus, and acellular pertussis (DTaP; Daptacel®, Infanrix®)	Children	None
<i>Haemophilus influenzae</i> type b (Hib; ActHIB®, Hiberix®, PedvaxHIB®)	Children	Hiberix approved in 2016 as a three-dose primary series at ages 2, 4, and 6 months (initially approved only as a booster dose for ages 15 months through 4 years).
Hepatitis A (HepA; Havrix®, Vaqta®)	Adults, children	None
Hepatitis B (HepB; Engerix-B®, Recombivax HB®, HEPLISAV-B®)	Adults, children, pregnant women (except for HEPLISAV-B, which is not recommended for children and pregnant women)	HEPLISAV-B approved in 2017.
Hepatitis A-Hepatitis B (HepA-HepB; Twinrix®)	Adults	None
Inactivated poliovirus (IPV; IPOL®)	Children	None
Influenza, inactivated (IIV; Afluria Quadrivalent®, Fluarix Quadrivalent®, Flucelvax Quadrivalent®, Flulaval Quadrivalent®, Fluzone High Dose Quadrivalent®, Fluzone Quadrivalent®)	Adults, children, pregnant women (except for Fluzone High Dose Quadrivalent, which is for adults aged 65 years and older)	Afluria Quadrivalent and Flucelvax Quadrivalent approved in 2016. Fluzone High Dose Quadrivalent approved in 2019. Flulaval Quadrivalent expanded use to 6 months of age and older in 2016. Afluria Quadrivalent and Fluarix Quadrivalent expanded use to 6 months of age and older in 2018. Fluzone Quadrivalent dose for children aged 6 through 35 months was updated to be either 0.25 mL or 0.5 mL in 2018.
Influenza, inactivated, adjuvanted (aIIV; Fluad®, Fluad Quadrivalent®)	Adults aged 65 years and older	Fluad approved in 2015; Fluad Quadrivalent approved in 2020. Changes to influenza strains for vaccine made annually.
Influenza, recombinant (RIV; Flublok Quadrivalent®)	Adults, pregnant women	Flublok Quadrivalent approved in 2017. Changes to influenza strains for vaccine made annually.
Influenza, live attenuated (LAIV; FluMist Quadrivalent®)	Adults (through 49 years of age), children	Changes to influenza vaccine strains made annually.
Measles, mumps, rubella (MMR; M-M-R II®)	Adults, children	None
Serogroup A, C, W, and Y meningococcal (MenACWY-D, Menactra®; Men-ACWY-CRM,	Adults, children	MenQuadFi (MenACWY-TT) was approved in 2020.

Vaccine (abbreviation; brand name)	Populations recommended for routine use	Recent changes to formulation, age indication, or dosing (within last five years)
Menveo®; MenACWY-TT, MenQuadFi®)		
Serogroup B meningococcal (MenB-FHbp, Trumenba®; MenB-4C, Bexsero®)	Adults, children	None.
Rotavirus (RV; Rotarix®, RotaTeq®)	Children	None
Tetanus, diphtheria, and acellular pertussis (Tdap; Adacel®, Boostrix®)	Children, adults, pregnant women	Adacel approved for repeat dose in people 10 through 64 years of age in 2019. ACIP recommendation updated to allow for use of Tdap or Td as decennial booster, wound prophylaxis, and catch up vaccination in 2020.
Tetanus, diphtheria (Td; TDVAX®, Tenivac®)	Adults	None
Varicella (VAR; Varivax®)	Children, adults	None
Zoster recombinant (RZV; Shingrix®)	Adults	Shingrix was approved in 2017. (Use of live zoster vaccine [Zostavax] was discontinued in November 2020.)

2123 Abbreviations: aIIV—Adjuvanted inactivated influenza vaccine; DTaP—Diphtheria and tetanus toxoids and acellular
2124 pertussis vaccine; HepA—Hepatitis A vaccine; HepB—Hepatitis B vaccine; HepA-HepB—Hepatitis A and Hepatitis B
2125 vaccines; Hib—*Haemophilus influenzae* type b vaccine; HPV9—9-valent human papillomavirus vaccine; IIV—
2126 Inactivated influenza vaccine; IPV—Inactivated poliovirus vaccine; LAIV—Live attenuated influenza vaccine;
2127 MenACWY—Serogroups A, C, W, and Y meningococcal vaccine; MenB—Serogroup B meningococcal vaccine;
2128 MMR—Measles, mumps, and rubella vaccine; MMR-V—Measles, mumps, rubella, and varicella vaccine; PCV13—
2129 13-valent pneumococcal conjugate vaccine; PPSV23—23-valent pneumococcal polysaccharide vaccine; RIV—
2130 Recombinant influenza vaccine; RV—Rotavirus vaccine; RZV—Recombinant zoster vaccine; Td—Tetanus and
2131 diphtheria toxoids; Tdap—Tetanus and diphtheria toxoids and acellular pertussis vaccine; VAR—Varicella vaccine
2132 Note: Combination vaccines that incorporate existing vaccines (e.g., DTaP-IPV/Hib) were also assessed, and are
2133 summarized in the Appendix.
2134

2135 **Table 2. Strength of Evidence (SoE) for safety of vaccines in children**

Vaccine (abbreviation; brand name[s])	Synthesis of SoE* and findings for vaccines currently in use in children
9-valent human papillomavirus (HPV9; Gardasil 9®)	Low: No evidence of increased risk of autoimmune disease, birth defects, death, reproductive system events, seizures, spontaneous abortion
13-valent pneumococcal conjugate (PCV13; Prevnar 13®)	Low: Increased risk of febrile seizures Moderate: No evidence of increased risk of death Low: No evidence of increased risk of asthma, cardiovascular events, intussusception, meningitis, reproductive system events, seizures
23-valent pneumococcal polysaccharide (PPSV23; Pneumovax®)	Insufficient evidence to draw conclusions about key adverse events
Diphtheria, tetanus, and acellular pertussis (DTaP; Daptacel®, Infanrix®)	Moderate: No evidence of increased risk of type 1 diabetes mellitus Low: No evidence of increased risk of asthma or death
Tetanus, diphtheria, and acellular pertussis (Tdap; Adacel®, Boostrix®)	Low: No evidence of increased risk of cardiovascular events, death
<i>Haemophilus influenzae</i> type b (Hib; PedvaxHIB®, ActHIB®, Hiberix®)	Moderate: No evidence of increased risk of serious adverse events in short term
Hepatitis A (HepA; Havrix®, Vaqta®)	Moderate: Increased risk of idiopathic thrombocytopenic purpura
Hepatitis B (HepB; Engerix-B®, Recombivax HB®)	Moderate: No evidence of increased risk of multiple sclerosis
Inactivated poliovirus (IPV; IPOL®)	Insufficient evidence to draw conclusions about key adverse events
Influenza, inactivated (IIV; Afluria Quadrivalent®, Fluarix Quadrivalent®, Flulaval Quadrivalent®, Fluzone Quadrivalent®, Flucelvax Quadrivalent®)	Moderate: No evidence of increased risk of death Low: No evidence of increased risk of anaphylaxis or systemic allergic reaction, asthma, autoimmune disease, cardiovascular events, febrile seizures, seizures
Influenza, live attenuated (LAIV; FluMist Quadrivalent®)	Low: No evidence of increased risk of death or seizures
Measles, mumps, and rubella (MMR; M-M-R II®)	High: No evidence of increased risk of autism High: Increased risk of anaphylaxis in children with allergies; increased risk of febrile seizures Moderate: Increased risk of idiopathic thrombocytopenic purpura Low: No evidence of increased risk for asthma

Vaccine (abbreviation; brand name[s])	Synthesis of SoE* and findings for vaccines currently in use in children
Meningococcal, A, C, W, and Y (MenACWY; MenACWY-D [Menactra®], MenACWY-CRM [Menveo®], MenACWY-TT [MenQuadFi®])	Moderate: No evidence of increased risk of cardiovascular events, diabetes, febrile seizures, intussusception, idiopathic thrombocytopenic purpura, Kawasaki disease, seizures Moderate: Increased risk of anaphylaxis in children with allergies Low: No evidence of increased risk of acute disseminated encephalomyelitis, anaphylaxis or systemic allergic reaction, asthma, autoimmune disease, death, encephalitis/encephalopathy, meningitis, multiple sclerosis, reproductive system events, transverse myelitis
Meningococcal B (MenB; MenB-4C [Bexsero®], MenB-FHbp [Trumenba®])	Moderate: No evidence of increased risk of anaphylaxis or systemic allergic reaction, reproductive system events Low: No evidence of increased risk of asthma, death, seizures
Rotavirus (RV; Rotarix®, RotaTeq®)	High: No evidence of increased risk of diabetes Moderate: No evidence of increased risk of intussusception (moderate SoE for increased risk from prior report was not confirmed when combining all available trials, though some observational studies showed increased risk). No evidence of increased risk of asthma, autoimmune disease, death, encephalitis/encephalopathy, febrile seizures, idiopathic thrombocytopenic purpura, seizures, stroke Low: No evidence of increased risk of anaphylaxis or systemic allergic reaction, autoimmune thyroiditis (Hashimoto's disease), Kawasaki disease, meningitis, reproductive system events
Varicella (VAR; Varivax®)	High: Increased risk of anaphylaxis Moderate: Increased risk of idiopathic thrombocytopenic purpura among children aged 11 to 17 years

2136 *Please see Appendix Table 5a for a description of the SoE and findings from the prior 2014 report (including
2137 adverse events not examined as key adverse events in the update), the update, and the synthesis across the report and
2138 update (including for combination vaccines).
2139

2140 **Table 3. Strength of Evidence (SoE) for safety of vaccines in adults**

Vaccine (abbreviation; brand name[s])	Synthesis of SoE and findings for vaccines currently in use in adults
9-valent human papillomavirus (HPV9; Gardasil 9®)	Insufficient evidence to draw conclusions; see Table 2 for studies that combined children and adults
13-valent pneumococcal conjugate (PCV13; Prevnar 13®)	Moderate: No evidence of increased risk of cardiovascular events, herpes zoster, myocardial infarction, reproductive system events, stroke Low: No evidence of increased risk of acute disseminated encephalomyelitis, anaphylaxis or systemic allergic reaction, asthma, autoimmune disease, death, encephalitis/encephalopathy, herpes zoster, idiopathic thrombocytopenic purpura, meningitis, seizures
23-valent pneumococcal polysaccharide (PPSV23; Pneumovax®)	High: No evidence of increased risk of cardiovascular or cerebrovascular events in adults aged 65 years and older Moderate: No evidence of increased risk of death
Hepatitis A (HepA; Havrix®, Vaqta®)	Insufficient evidence to draw conclusions about key adverse events

Vaccine (abbreviation; brand name[s])	Synthesis of SoE and findings for vaccines currently in use in adults
Hepatitis B (HepB; Engerix-B®, Recombivax HB®, HEPLISAV-B®)	<p>Moderate: No evidence of increased risk of multiple sclerosis (for hepatitis B vaccines except HEPLISAV-B, for which there was insufficient evidence)</p> <p>Moderate: No evidence of increased risk of diabetes (across all hepatitis B vaccines)</p> <p>Moderate: Increased risk of anaphylaxis in patients allergic to yeast (for hepatitis B vaccines except HEPLISAV-B, for which there were no studies)</p> <p>Low: No evidence of increased risk of asthma, autoimmune disease, cardiovascular events, death, herpes zoster, reproductive system events; stroke for HEPLISAV-B</p>
Influenza, inactivated (IIV; Afluria Quadrivalent®, Flucelvax Quadrivalent®, Fluarix Quadrivalent®, Flulaval Quadrivalent®, Fluzone High Dose Quadrivalent®, Fluzone Quadrivalent®)	<p>Low: No evidence of increased risk of asthma, cardiovascular events, death, myocardial infarction, reproductive system events, seizures, stroke</p>
Influenza, inactivated, adjuvanted (aIIV; Fluad®, Fluad Quadrivalent®)	<p>Moderate: No evidence of increased risk of stroke</p> <p>Low: No evidence of increased risk of asthma, autoimmune disease, cardiovascular events, death, encephalitis/encephalopathy, Guillain-Barré syndrome, idiopathic thrombocytopenic purpura, myocardial infarction, seizures</p>
Influenza, recombinant (RIV; Flublok Quadrivalent®)	<p>Low: No evidence of increased risk of cardiovascular events, death, encephalitis/encephalopathy, myocardial infarction, reproductive system events, stroke</p>
Influenza, live attenuated (LAIV; FluMist Quadrivalent®)	<p>Insufficient evidence to draw conclusions about key adverse events</p>
Measles, mumps, and rubella (MMR; M-M-R II®)	<p>Moderate: No evidence of increased risk of type 1 diabetes mellitus</p>
Meningococcal A, C, W, and Y (MenACWY; MenACWY-D [Menactra®], MenACWY-CRM [Menveo®], MenACWY-TT [MenQuadFi®])	<p>Moderate: No evidence of increased risk of death</p> <p>Low: No evidence of increased risk of cardiovascular events, myocardial infarction, stroke</p>
Meningococcal B (MenB; MenB-4C [Bexsero®], MenB-FHbp [Trumenba®])	<p>Insufficient evidence to draw conclusions about key adverse events; see Table 2 for studies that combined children and adults</p>
Tetanus, diphtheria, and acellular pertussis (Tdap; Adacel®, Boostrix®) and tetanus and diphtheria (Td; TDVAX, Tenivac®)	<p>High: Increased risk of anaphylaxis</p>
Varicella (VAR; Varivax®)	<p>Insufficient evidence to draw conclusions about key adverse events</p>
Zoster recombinant (RZV; Shingrix®)	<p>High: No evidence of increased risk of herpes zoster</p> <p>Moderate: No evidence of increased risk of amyotrophic lateral sclerosis, anaphylaxis or systemic allergic reaction, asthma, cardiovascular events, death, diabetes, encephalitis/encephalopathy, Guillain-Barré syndrome, idiopathic</p>

Vaccine (abbreviation; brand name[s])	Synthesis of SoE and findings for vaccines currently in use in adults
	<p>thrombocytopenic purpura, meningitis, myocardial infarction, reproductive system events, seizures, stroke</p> <p>Low: No evidence of increased risk of acute disseminated encephalomyelitis, angioedema, ataxia, autoimmune disease, autoimmune thyroiditis (Hashimoto's disease)</p>

2141 *Please see Appendix Table 6a for a description of the SoE and findings from the prior 2014 report (including
2142 adverse events not examined as key adverse events in the update), the update, and the synthesis across the report and
2143 update (including for combination vaccines).

2144

2145 **Table 4. Strength of Evidence (SoE) for safety of vaccines in pregnant women**

Vaccine (abbreviation; brand name[s])	Synthesis of SOE* and findings for vaccines currently in use in pregnant women
Hepatitis B (HepB; Engerix-B®, Recombivax HB®)	Insufficient evidence to draw conclusions about key adverse events
Influenza, inactivated (IIV; Afluria Quadrivalent®, Flucelvax Quadrivalent®, Fluarix Quadrivalent®, Flulaval Quadrivalent®, Fluzone Quadrivalent®)	Insufficient evidence to draw conclusions about key adverse events
Influenza, recombinant (RIV; Flublok Quadrivalent®)	Insufficient evidence to draw conclusions about key adverse events
Tetanus, diphtheria, and acellular pertussis (Tdap; Adacel®, Boostrix®)	<p>Moderate: No evidence of increased risk of maternal cardiovascular events, maternal death, maternal diabetes, eclampsia/pre-eclampsia, preterm labor, maternal reproductive system events, stillbirth, cardiovascular events in infants, death in infants, encephalitis/encephalopathy in infants, seizures in infants</p> <p>Low: No evidence of increased risk of maternal encephalitis/encephalopathy, autism in infants, birth defects in infants, febrile seizures in infants</p>

2146 *Please see Appendix Table 7a for a description of the SoE and findings from the prior 2014 report, the update, and
 2147 the synthesis across the report and update.
 2148