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Nimenrix[™]

1. NAME OF THE MEDICINAL PRODUCT

Nimenrix™

Meningococcal polysaccharide serogroups A, C, W-135, and Y conjugate vaccine

2. QUALITATIVE AND QUANTITATIVE COMPOSITION

After reconstitution, 1 dose (0.5 ml) contains 5 micrograms of polysaccharide for *Neisseria* meningitidis groups A¹, C¹, W-135¹ and Y¹.

¹ conjugated to tetanus toxoid carrier protein 44 micrograms

3. PHARMACEUTICAL FORM

White cake or powder. After reconstitution with diluent: clear and colourless liquid.

4. CLINICAL PARTICULARS

4.1 Therapeutic indications

Active immunization of individuals from 6 weeks of age against invasive meningococcal disease caused by *Neisseria meningitidis* groups A, C, W-135, and Y (see section 5.1).

4.2 Posology and method of administration

Posology

Nimenrix[™] should be used in accordance with available official recommendations.

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Table 1: Posology

Age Group	Primary Immunization	Booster
Infants from 6 weeks to less	Two doses, each of 0.5 ml, with	At 12 months of age
than 6 months of age*	the first dose given from 6	
	weeks of age, with an interval	
	of 2 months between doses	
Unvaccinated infants from	One dose of 0.5 ml given from	At 12 months of age with a
6 months to less than	6 months of age	minimum interval of at least
12 months of age**		2 months after the primary
		dose
Children from 12 months of	One dose of 0.5 ml	Not routinely administered
age, adolescents and adults**		

^{*} See section 5.1 for further information.

Long-term antibody persistence data following vaccination with **Nimenrix**[™] are available up to 10 years after vaccination (see sections 4.4 and 5.1).

Nimenrix[™] may be given as a booster dose to individuals who have previously received primary vaccination with **Nimenrix**[™] or other conjugated or plain polysaccharide meningococcal vaccines (see section 5.1).

Special populations

Individuals who have underlying conditions predisposing them to meningococcal infection due to anatomic or functional asplenia (such as sickle cell disease) may receive at least one dose of **Nimenrix**TM (see sections 4.8 and 5.1).

Method of administration

Nimenrix[™] is for intramuscular injection only.

In infants, the recommended injection site is the anterolateral aspect of the thigh.

^{**}In some situations, consideration may be given to administering an additional primary dose or a booster dose of **Nimenrix**[™] (see sections 4.4 and 5.1 for further information).

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In individuals from 1 year of age, the recommended injection site is the anterolateral aspect of the

thigh or deltoid muscle (see sections 4.4 and 4.5).

For instructions on reconstitution of the medicinal product before administration, see section 6.6.

4.3 Contraindications

Nimenrix[™] should not be administered to subjects with hypersensitivity to the active substances

or to any of the excipients contained in the vaccine.

4.4 Special warnings and precautions for use

Nimenrix[™] should under no circumstances be administered intravascularly, intradermally or

subcutaneously.

It is good clinical practice to precede vaccination by a review of the medical history (especially

with regard to previous vaccination and possible occurrence of undesirable effects) and a clinical

examination.

As with all injectable vaccines, appropriate medical treatment and supervision should always be

readily available in case of a rare anaphylactic event following the administration of the vaccine.

Intercurrent illness

As with other vaccines, vaccination with **Nimenrix**[™] should be postponed in subjects suffering

from an acute severe febrile illness. The presence of a minor infection, such as a cold, should not

result in the deferral of vaccination.

Syncope

Syncope (fainting) can occur following, or even before, any vaccination as a psychogenic

response to the needle injection. It is important that procedures are in place to avoid injury from

faints.

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Thrombocytopenia and coagulation disorders

As with other vaccines administered intramuscularly, **Nimenrix**TM should be given with caution to

individuals with thrombocytopenia or any coagulation disorder since bleeding may occur following

an intramuscular administration to these subjects.

<u>Immunodeficiency</u>

It may be expected that in patients receiving immunosuppressive treatment or patients with

immunodeficiency, an adequate immune response may not be elicited.

Persons with certain complement deficiencies and persons receiving treatment that inhibits

terminal complement activation (for example, eculizumab) are at increased risk for invasive

disease caused by Neisseria meningitidis groups A, C, W-135, and Y even if they develop

antibodies following vaccination with Nimenrix™.

Special populations

Limited data are available on the safety and immunogenicity in individuals with increased

susceptibility to meningococcal infection due to anatomic or functional asplenia (such as sickle cell

disease) (see sections 4.2, 4.8 and 5.1).

Protection against meningococcal disease

Nimenrix[™] will only confer protection against *Neisseria meningitidis* groups A, C, W-135, and Y.

The vaccine will not protect against other Neisseria meningitidis groups.

As with any vaccine, a protective immune response may not be elicited in all vaccinees.

Immune response in infants aged 6 months to less than 12 months

A single-dose administered at 6 months was associated with lower human complement serum

bactericidal assay (hSBA) titres to groups W-135 and Y compared with three doses administered

at 2, 4, and 6 months (see section 5.1). The clinical relevance of this observation is unknown. If

an infant aged 6 months to less than 12 months is expected to be at immediate risk of invasive

meningococcal disease due to exposure to groups W-135 and/or Y, consideration may be given to

administering a second primary dose of **Nimenrix**TM after an interval of 2 months.

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Immune responses in toddlers aged 12-14 months

At 1 month post vaccination, toddlers aged 12-14 months had similar rSBA titres to groups A, C, W-135, and Y following one dose of **Nimenrix**[™] or two doses of **Nimenrix**[™] given 2 months apart. At 1 year post vaccination, the rSBA titres to groups A, C, W-135, and Y were similar in both the one and the two dose groups (see section 5.1).

Measured with a serum bactericidal assay using human complement (hSBA), 1 month post vaccination, responses to groups W-135 and Y were lower after a single dose than after 2 doses given 2 months apart, while responses to groups A and C were similar in the two groups (see section 5.1). The clinical relevance of these observations is unknown. If a toddler is expected to be at immediate risk of invasive meningococcal disease due to the exposure to groups W-135 and/or Y, consideration may be given to administering a second primary dose after an interval of 2 months. At 1 year post vaccination, the hSBA responses for groups A, C, W-135, and Y were similar in both the one and the two dose groups (see section 5.1). Regarding waning of antibody against group A or group C after a first dose of **Nimenrix**™ in children aged 12-23 months, see under Persistence of serum bactericidal antibody titres.

Persistence of serum bactericidal antibody titres

Persistence of antibodies has been evaluated up to 10 years after vaccination. The persistence studies with **Nimenrix**TM have shown a waning of serum bactericidal antibody titres against group A when using human complement in the assay (hSBA) (see section 5.1). The clinical relevance of this observation is unknown. However, if an individual is expected to be at particular risk of exposure to group A and received a dose of **Nimenrix**TM more than approximately 1 year previously, consideration may be given to administering a booster dose.

Similar to the monovalent Men C comparator, a decline in antibody titres over time has been observed. The clinical relevance of this observation is unknown. A booster dose might be considered in individuals remaining at high risk of exposure to meningococcal disease caused by groups A, C, W-135, and Y (see section 5.1).

Although **Nimenrix**[™] contains tetanus toxoid, this vaccine does not substitute for tetanus immunization.

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4.5 Interactions with other medicinal products and other forms of interaction

In infants, **Nimenrix**[™] can be given concomitantly with combined diphtheria, tetanus, acellular

pertussis, hepatitis B, inactivated poliovirus and Haemophilus influenzae type b vaccines

(DTaP/IPV/Hib/HepB), as well as 10-valent pneumococcal conjugate vaccine.

From age 1 year and above, Nimenrix[™] can be given concomitantly with any of the following

vaccines: hepatitis A (HAV) and hepatitis B (HBV) vaccines, measles - mumps - rubella (MMR)

vaccine, measles - mumps - rubella - varicella (MMRV) vaccine, 10-valent pneumococcal

conjugate vaccine or unadjuvanted seasonal influenza vaccine.

Nimenrix[™] can also be given concomitantly with combined diphtheria - tetanus - acellular

pertussis vaccines, including combination DTaP vaccines with hepatitis B, inactivated poliovirus or

Haemophilus influenzae type b, such as DTaP/IPV/Hib/HepB vaccine and 13-valent pneumococcal

conjugate vaccine in the second year of life.

In individuals aged 9 to 25 years, Nimenrix[™] can be given concomitantly with human

papillomavirus bivalent [Type 16 and 18] vaccine, recombinant (HPV2).

Safety and immunogenicity of Nimenrix[™] was evaluated when sequentially administered or co-

administered with a DTaP/IPV/Hib/HepB vaccine in the second year of life. The administration of

NimenrixTM 1 month after the DTaP/IPV/Hib/HepB vaccine resulted in lower MenA, MenC and

MenW-135 Geometric Mean Titres (GMTs) as measured with a serum bactericidal assay using

rabbit complement (rSBA). The clinical relevance of this observation is unknown, since at least

99.4% of subjects (N=178) had rSBA titres ≥8 for each group (A, C, W-135, and Y). Whenever

possible, Nimenrix[™] and a tetanus toxoid (TT) containing vaccine, such as DTaP/IPV/Hib/HepB

vaccine, should be co-administered or **Nimenrix**[™] should be administered at least 1 month before

the TT-containing vaccine.

One month after co-administration with a 10-valent pneumococcal conjugate vaccine, in toddlers

aged 12-23 months, lower Geometric Mean antibody Concentrations (GMCs) and

opsonophagocytic assay (OPA) antibody GMTs were observed for one pneumococcal serotype

(18C conjugated to tetanus toxoid carrier protein). The clinical relevance of this observation is

unknown. There was no impact of co-administration on the other nine pneumococcal serotypes.

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One month after co-administration with a combined tetanus toxoid, reduced diphtheria toxoid and

acellular pertussis vaccine, adsorbed (Tdap) in subjects aged 9 to 25 years, lower GMCs were

observed to each pertussis antigen (pertussis toxoid [PT], filamentous haemagglutinin [FHA] and

pertactin [PRN]). More than 98% of subjects had anti-PT, FHA or PRN concentrations above the

assay cut-off thresholds. The clinical relevance of these observations is unknown. There was no

impact of co-administration on immune responses to **Nimenrix**[™] or the tetanus or diphtheria

antigens included in Tdap.

If Nimenrix[™] is to be given at the same time as another injectable vaccine, the vaccines should

always be administered at different injection sites.

As with other vaccines, it may be expected that in patients receiving immunosuppressive

treatment an adequate response may not be elicited.

4.6 Fertility, pregnancy and lactation

Pregnancy

There is limited experience with use of **Nimenrix**[™] in pregnant women.

Animal studies with **Nimenrix**[™] do not indicate direct or indirect harmful effects with respect to

fertility, pregnancy, embryo/foetal development, parturition or post-natal development (see section

5.3).

Nimenrix[™] should be used during pregnancy only when clearly needed, and the possible

advantages outweigh the potential risks for the foetus.

Lactation

The safety of **Nimenrix**[™] when administered to breast-feeding women has not been evaluated. It

is unknown whether **Nimenrix**[™] is excreted in human breast milk.

Nimenrix[™] should only be used during breast-feeding when the possible advantages outweigh

the potential risks.

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4.7 Effects on ability to drive and use machines

No studies on the effects of **Nimenrix**[™] on the ability to drive and use machines have been performed.

4.8 Undesirable effects

Clinical Trial Data

The safety profile presented in Table 2 is based on two data sets:

- a pooled analysis in more than 9,000 subjects from the age of 1 year on, who have been vaccinated with 1 dose of NimenrixTM in clinical studies.
- data from approximately 1,000 infants (6 weeks to 12 months of age) who have been primed and boosted with NimenrixTM.

Table 2: Adverse Reactions by System Organ Class and Council for International Organizations of Medical Science (CIOMS) Frequency Category Listed in Order of Decreasing Medical Seriousness or Clinical Importance Within Each Frequency Category and SOC

System Organ Class	Very Common ≥1/10	Common ≥1/100 to <1/10	Uncommon ≥1/1,000 to <1/100	Rare ≥1/10,000 to <1/1,000	Very Rare <1/10,000	Frequency Not Known (cannot be estimated from the available data)
Metabolism and nutrition disorders	Appetite lost					
Psychiatric disorders	Irritability		Insomnia Crying			
Nervous system disorders	Drowsiness Headache ¹		Hypoaesthesia ¹ Dizziness ¹			
Gastrointestinal disorders		Gastrointestinal symptoms (including diarrhoea,				

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Table 2: Adverse Reactions by System Organ Class and Council for International Organizations of Medical Science (CIOMS) Frequency Category Listed in Order of Decreasing Medical Seriousness or Clinical Importance Within Each Frequency Category and SOC

System Organ	Very	Common	Uncommon	Rare	Very Rare	Frequency Not
Class	Common	≥1/100 to	≥1/1,000 to	≥1/10,000	<1/10,000	Known
	≥1/10	<1/10	<1/100	to		(cannot be
				<1/1,000		estimated from
						the available
						data)
		vomiting and				
		nausea ²)				
Skin and			Pruritus ¹			
subcutaneous			Rash ³			
tissue disorders						
Musculoskeletal			Myalgia ¹			
and connective			Pain in			
tissue disorders			extremity ¹			
General disorders	Fever	Injection site	Malaise			Extensive limb
and	Injection site	haematoma ²	Injection site			swelling at the
administration site	swelling		reaction			injection site,
conditions	Injection site		(including			frequently
	pain		induration,			associated with
	Injection site		pruritus,			erythema,
	redness		warmth,			sometimes
	Fatigue ¹		anaesthesia)			involving the
						adjacent joint or
						swelling of the
						entire injected
						limb*

^{*}Adverse Reaction identified post-marketing.

¹ Not reported in the infant clinical study (MenACWY-TT-083)

^{2.} Occurred at a frequency of Uncommon in infants

^{3.} Occurred at a frequency of Common in infants

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Local and general adverse reactions

In all age groups, the local adverse reactions of pain, redness and swelling at the injection site were reported at a very common frequency after vaccination.

In the infant and toddler groups, the general adverse reactions of drowsiness, fever, irritability/fussiness and loss of appetite were reported at a very common frequency after vaccination.

In a separate infant study, 554 infants were primed with one or three doses of **Nimenrix**[™] and 508 received booster doses in the second year of life. Local and general adverse reactions in this study were similar in frequency to the larger infant study.

In the 12-14 months age group who received two doses of **Nimenrix**[™] given 2 months apart, the first and second doses were associated with similar local and systemic reactogenicity.

In an additional clinical study of age matched subjects who were either healthy or at increased risk of meningococcal disease due to anatomical or functional asplenia (such as sickle cell disease), the safety profile of **Nimenrix**[™] in at-risk children and adolescents was generally similar to that observed in the non-asplenic population (see section 5.1).

The 2–5 year group reported general adverse reactions at a frequency ranging from common (irritability, loss of appetite and fever) to very common (drowsiness).

In the 6-10, 11-17 and ≥18 years age groups, the general adverse reactions were reported at a frequency ranging from common (gastrointestinal symptoms and fever) to very common (headache and fatique).

In a clinical study of 11 to 25 year old subjects co-administered **Nimenrix**[™] and Tdap or given the vaccines separately, the local reactions (injection site pain, redness, and swelling) and general reactions (fatigue and headache) occurred at a similar frequency in both groups and in the subjects in the pooled analysis (very common). The general reactions gastrointestinal events (nausea, vomiting, diarrhoea, abdominal pain) occurred more frequently (very common) and fever occurred less frequently (common) compared to subjects in the pooled analysis, but occurred at a similar frequency in subjects co-administered the vaccines and subjects given the vaccines separately in the study.

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In a clinical study of female subjects 9 to 25 years old, the local reactions (pain, redness, and

swelling at the **Nimenrix**[™] injection site) and general reactions (headache, fever, and fatigue)

occurred at a similar frequency in subjects co-administered Nimenrix[™], Tdap and HPV2 and in

subjects given Nimenrix[™] alone, as they did in subjects in the pooled analysis (very common).

The general reactions gastrointestinal events (nausea, vomiting, diarrhoea, abdominal pain) and

myalgia occurred at a similar frequency in the two groups but more frequently than in the pooled

analysis (very common), as did the general reaction rash (common).

The local and general adverse reaction profile of a booster dose of **Nimenrix**TM given to subjects

from 12 months of age after primary vaccination with **Nimenrix**TM or other conjugated or plain

polysaccharide meningococcal vaccines, was similar to the local and general adverse reaction

profile observed after primary vaccination with Nimenrix[™], except gastrointestinal symptoms

(including diarrhoea, vomiting, and nausea) which ranged from common to very common among

subjects 6 years of age and older (versus common after primary vaccination).

4.9 Overdose

No cases of overdose have been reported.

5. PHARMACOLOGICAL PROPERTIES

5.1 Pharmacodynamics Properties

Pharmacotherapeutic group: bacterial vaccines, ATC code J07AH08

Mechanism of action

Anti-capsular meningococcal antibodies protect against meningococcal disease via complement

mediated bactericidal killing. Nimenrix[™] induces the production of bactericidal antibodies against

capsular polysaccharides of Neisseria meningitidis groups A, C, W-135, and Y when measured by

assays using either rSBA or hSBA. By conjugating capsular polysaccharide to a protein carrier

that contains T-cell epitopes, meningococcal conjugate vaccines like Nimenrix[™] change the

nature of immune response to capsular polysaccharide from T-cell independent to T-cell

dependent.

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Immunogenicity in infants

In Study MenACWY-TT-083, the immunogenicity of a 2-dose primary vaccination schedule administered at 2 and 4 months of age was evaluated. Routinely used infant vaccines DTaP/IPV/Hib/HepB and a 10-valent pneumococcal vaccine were co-administered. For group C, rSBA and hSBA titres elicited by **Nimenrix**[™] were compared to a 2-dose priming with licensed monovalent meningococcal conjugate group C vaccines, MenC-CRM and MenC-TT vaccines. **Nimenrix**[™] elicited rSBA and hSBA titres against the four meningococcal groups. The response against group C was non-inferior to the one elicited by the licensed MenC-CRM and MenC-TT vaccines in terms of the percentage of subjects with rSBA titres ≥8 at 1 month after the second dose.

For subjects initially vaccinated in infancy with **Nimenrix**TM at 2 and 4 months of age and receiving a **Nimenrix**TM booster dose at 12 months of age, the increase in rSBA and hSBA titres 1 month post-booster dose ranged between 15 and 80-fold for all groups and more than 99.0% of all infants achieved post-booster titres above 8 for both assays. The observed booster response for group C was similar to that observed in subjects primed and boosted with a monovalent MenC conjugate vaccine (TT or CRM conjugated). Results are shown in Table 3.

Table 3: rSBA and hSBA titres following two doses of Nimenrix[™] (or MenC-CRM or MenC-TT) given 2 months apart with the first dose administered to infants 6-12 weeks of age and following a booster at 12 months of age (Study MenACWY-TT-083)

Mening				rSBA*			hSBA	* *
o-coccal	Vaccine	Time		≥8	GMT		≥8	GMT
group	group	point	N	(95% CI)	(95% CI)	N	(95% CI)	(95% CI)
		MO	450	97.4%	203	000	96.5%	157
	Nimenrix ^T	M3	456	(95.4; 98.6)	(182; 227)	202	(93.0; 98.6)	(131; 188)
Α	М		400	99.6%	1561	044	99.5%	1007
		M11	462	(98.4; 99.9)	(1412; 1725)	214	(97.4;100)	(836;1214)
		MO	450	98.7%	612	040	98.6%	1308
	Nimenrix ^T	M3	456	(97.2; 99.5)	(540; 693)	218	(96.0; 99.7)	(1052; 1627)
	М		400	99.8%	1177	004	99.5%	4992
С		M11	463	(98.8; 100)	(1059; 1308)	221	(97.5; 100)	(4086; 6100)
				99.6%	958		100%	3188
		M3	455	(98.4; 99.9)	(850; 1079)	202	(98.2; 100)	(2646; 3841)

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	MenC- CRM vaccine	M11	446	98.4% (96.8; 99.4)	1051 (920; 1202)	216	100% (98.3; 100)	5438 (4412; 6702)
	MenC-TT	M3 MenC-TT		100% (99.2; 100)	1188 (1080; 1307)	226	100% (98.4; 100)	2626 (2219; 3109)
	vaccine	M11	459	100% (99.2; 100)	1960 (1776; 2163)		100% (98.3; 100)	5542 (4765; 6446)
	Nimenrix ^T	М3	455	99.1% (97.8; 99.8)	1605 (1383; 1862)	217	100% (98.3; 100)	753 (644; 882)
W-135	М	M11	462	99.8% (98.8; 100)	2777 (2485; 3104)	218	100% (98.3; 100)	5123 (4504; 5826)
	Y Nimenrix ^T M3 M1		456	98.2% (96.6; 99.2)	483 (419; 558)	214	97.7% (94.6; 99.2)	328 (276; 390)
Y			462	99.4% (99.1; 99.9)	881 (787; 986)	217	100% (98.3; 100)	2954 (2498; 3493)

The analysis of immunogenicity was conducted on the primary according-to-protocol (ATP) cohort.

M3 = post-primary vaccination at Month 3

M11 = post-booster vaccination at Month 11

In Study MenACWY-TT-087, infants received either a single primary dose at 6 months followed by a booster dose at 15-18 months or three primary doses at 2, 4, and 6 months followed by a booster dose at 15-18 months. All subjects also received DTaP-IPV/Hib and 10-valent pneumococcal conjugate vaccines at all time points. A single primary dose administered at 6 months of age elicited robust rSBA titres to the four meningococcal groups, as measured by the percentage of subjects with rSBA titres ≥8, that were comparable to responses after the last dose of a three-dose primary series. A booster dose produced robust responses, comparable between the two dosing groups, against all four meningococcal groups. Results are shown in Table 4.

^{*}rSBA analysis performed at Public Health England (PHE) laboratories in UK

^{**}hSBA analysis performed at GSK laboratories

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Table 4: rSBA and hSBA titres following a single dose of Nimenrix[™] in infants at 6 months of age and pre- and post-booster at 15-18 months of age (Study MenACWY-TT-087)

Meningo-			rSB <i>A</i>	*		hSB	A**
coccal	Time point		≥8	GMT		≥8	GMT
group		N	(95% CI)	(95% CI)	N	(95% CI)	(95% CI)
	Post dose	400	98.8%	1333		98.3%	271
	1 ⁽¹⁾	163	(95.6; 99.9)	(1035; 1716)	59	(90.9; 100)	(206; 355)
		404	81.7%	125		66.2%	20.8
A	Pre booster	131	(74; 87.9)	(84.4; 186)	71	(54; 77)	(13.5; 32.2)
	Post	400	99.3%	2762		100%	1416
	booster ⁽¹⁾	139	(96.1; 100)	(2310; 3303)	83	(95.7; 100)	(1140; 1758)
	Post dose		99.4%	592		100%	523
	1 ⁽¹⁾	163	(96.6; 100)	(482; 726)	66	(94.6;100)	(382; 717)
		101	65.6%	27.4		96.2%	151
С	Pre booster	131	(56.9; 73.7)	(20.6; 36.6)	78	(89.2; 99.2)	(109; 210)
	Post		99.3%	2525	00	100%	13360
	booster ⁽¹⁾	139	(96.1; 100)	(2102; 3033)	92	(96.1; 100)	(10953; 16296)
	Post dose	400	93.9%	1256	47	87.2%	137
	1 ⁽¹⁾	163	(89; 97)	(917; 1720)	47	(74.3; 95.2)	(78.4; 238)
		404	77.9%	63.3		100%	429
W-135	Pre booster	131	(69.8; 84.6)	(45.6; 87.9)	53	(93.3; 100)	(328; 559)
	Post	400	100%	3145	5 0	100%	9016
	booster ⁽¹⁾	139	(97.4; 100)	(2637; 3750)	59	(93.9; 100)	(7045; 11537)
	Post dose	400	98.8%	1470	5 0	92.3%	195
	1 ⁽¹⁾	163	(95.6; 99.9)	(1187; 1821)	52	(81.5; 97.9)	(118; 323)
.,		464	88.5%	106	6.4	98.4%	389
Y	Pre booster	131	(81.8; 93.4)	(76.4; 148)	61	(91.2; 100)	(292; 518)
	Post	400	100%	2749	00	100%	5978
	booster ⁽¹⁾	139	(97.4; 100)	(2301; 3283)	69	(94.8; 100)	(4747; 7528)

The analysis of immunogenicity was conducted on the primary ATP cohort.

^{*}rSBA analysis performed at PHE laboratories UK

^{**}hSBA analysis performed at Neomed, Canada

⁽¹⁾ blood sampling performed 1-month post vaccination

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Measurement of hSBA titres was a secondary endpoint in Study MenACWY-TT-087. Although similar responses to groups A and C were observed with both dosing schedules, a single primary dose in infants at 6 months was associated with lower hSBA titres to groups W-135 and Y as measured by the percentage of subjects with hSBA titres ≥8 [87.2% (95% CI: 74.3, 95.2) and 92.3% (95% CI: 81.5, 97.9), respectively] compared with three primary doses at 2, 4, and 6 months of age [100% (95% CI: 96.6, 100) and 100% (95% CI: 97.1, 100), respectively] (see section 4.4). After a booster dose, hSBA titres to all four meningococcal groups were comparable between the two dosing schedules (Table 4).

Immunogenicity in toddlers aged 12-23 months

In clinical studies MenACWY-TT-039 and MenACWY-TT-040, a single dose of **Nimenrix**[™] elicited SBA titres against the four meningococcal groups, with group C rSBA titres that were comparable to those elicited by a licensed MenC-CRM vaccine in terms of the percentage of subjects with rSBA titres ≥8. In Study MenACWY-TT-039, hSBA was also measured as a secondary endpoint. Results are shown in Table 5.

Table 5: SBA* titres following a single dose of Nimenrix[™] (or MenC-CRM) in toddlers aged 12-23 months (Studies MenACWY-TT-039/040)

Menin				Study MenAC	WY-TT	-039 ⁽¹⁾		S	tudy MenACW	/Y-TT-040 ⁽²⁾
go-	Vaccine		rSBA	*	hSBA*				rSBA	*
coccal	group	N	≥8	GMT	N	≥8	GMT	N	≥8	GMT
group		IN	(95% CI)	(95% CI)	IN	(95% CI)	(95% CI)	IN	(95% CI)	(95% CI)
A	Nimenrix™	354	99.7%	2205	338	77.2%	19.0	183	98.4%	3170
	Milliellix	334	(98.4; 100)	(2008; 2422)	330	(72.4; 81.6)	(16.4; 22.1)	100	(95.3; 99.7)	(2577; 3899)
	TM	254	99.7%	478	244	98.5%	196	100	97.3%	829
	Nimenrix [™]	354	(98.4; 100)	(437; 522)	341	(96.6; 99.5)	(175; 219)	183	(93.7; 99.1)	(672; 1021)
С	MenC-									
	CRM	121	97.5%	212	116	81.9%	40.3	114	98.2%	691
		121	(92.9; 99.5)	(170; 265)	110	(73.7; 88.4)	(29.5; 55.1)	114	(93.8; 99.8)	(521; 918)
	vaccine									
W-135	Nimenrix™	354	100%	2682	336	87.5%	48.9	186	98.4%	4022
11-133	Millellix	004	(99.0; 100)	(2453; 2932)	550	(83.5; 90.8)	(41.2; 58.0)	100	(95.4; 99.7)	(3269; 4949)
		054	100%	2729	000	79.3%	30.9	405	97.3%	3168
Y	Nimenrix [™]	354	(99.0; 100)	(2473; 3013)	329	(74.5; 83.6)	(25.8; 37.1)	185	(93.8; 99.1)	(2522; 3979)

The analysis of immunogenicity was conducted on the ATP cohorts.

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In Study MenACWY-TT-104, **Nimenrix**[™] elicited rSBA titres against all four meningococcal groups following one or two doses administered 2 months apart that were similar in terms of the percentage of subjects with rSBA titres ≥8 and GMT. Results are shown in Table 6.

Table 6: rSBA and hSBA titres following one or two doses of Nimenrix[™] with the first dose administered to toddlers aged 12-14 months (Study MenACWY-TT-104)

Meningoc	Nimenrix [™]			rSBA	•		hSBA*	*
occal group	dose group	Time point ⁽¹⁾	N	≥8 (95%CI)	GMT (95% CI)	N	≥8 (95% CI)	GMT (95% CI)
9.04	9.046			, ,	,		, ,	, ,
		1 Month	180	97.8%	1437	74	95.9%	118
	1 dose	Post dose 1		(94.4; 99.4)	(1118; 1847)		(88.6; 99.2)	(86.8; 161)
	i dose	1 Year	167	63.5%	62.7	70	35.7%	6.1
		Post dose 1	107	(55.7; 70.8)	(42.6; 92.2)	70	(24.6; 48.1)	(4.1; 8.9)
		1 Month	450	96.8%	1275	00	97.0%	133
Α		Post dose 1	158	(92.8; 99.0)	(970; 1675)	66	(89.5; 99.6)	(98.1; 180)
		1 Month		98.0%	1176		97.0%	170
	2 doses	Post dose 2	150	(94.3; 99.6)	(922; 1501)	66	(89.5; 99.6)	(126; 230)
		1 Year	1.10	70.6%	76.6		35.5%	6.4
		Post dose 2	143	(62.4; 77.9)	(50.7; 116)	62	(23.7; 48.7)	(4.2; 10.0)
		1 Month	4=0	95.0%	452		98.7%	152
		Post dose 1	179	(90.7; 97.7)	(346; 592)	78	(93.1; 100)	(105; 220)
	1 dose	1 Year		49.1%	16.2	7.4	80.3%	35.2
		Post dose 1	167	(41.3; 56.9)	(12.4; 21.1)	71	(69.1; 88.8)	(22.5; 55.2)
		1 Month		95.5%	369		95.7%	161
С		Post dose 1	157	(91.0; 98.2)	(281; 486)	70	(88.0; 99.1)	(110; 236)
		1 Month		98.7%	639		100%	1753
	2 doses	Post dose 2	150	(95.3; 99.8)	(522; 783)	69	(94.8; 100)	(1278; 2404)
		1 Year		55.2%	21.2		90.5%	73.4
		Post dose 2	143	(46.7; 63.6)	(15.6; 28.9)	63	(80.4; 96.4)	(47.5; 113)

⁽¹⁾ blood sampling performed 42 to 56 days post vaccination

⁽²⁾ blood sampling performed 30 to 42 days post vaccination

^{*} SBA analyses performed at GSK laboratories

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Meningoc	Nimenrix [™]			rSBA'	*		hSBA*	*
occal	dose	Time point ⁽¹⁾		≥8	GMT		≥8	GMT
group	group		N	(95%CI)	(95% CI)	N	(95% CI)	(95% CI)
		1 Month	100	95.0%	2120	70	62.5%	27.5
	1 dose	Post dose 1	180	(90.8; 97.7)	(1601; 2808)	72	(50.3; 73.6)	(16.1; 46.8)
	1 dose	1 Year	407	65.3%	57.2	70	95.8%	209
		Post dose 1	167	(57.5; 72.5)	(39.9; 82.0)	72	(88.3; 99.1)	(150; 291)
W-135		1 Month	450	94.9%	2030	04	68.9%	26.2
		Post dose 1	158	(90.3; 97.8)	(1511; 2728)	61	(55.7; 80.1)	(16.0; 43.0)
		1 Month	450	100%	3533	70	97.1%	757
	2 doses	Post dose 2	150	(97.6; 100)	(2914; 4283)	70	(90.1; 99.7)	(550; 1041)
		1 Year	4.40	77.6%	123	0.5	98.5%	233
		Post dose 2	143	(69.9; 84.2)	(82.7; 183)	65	(91.7; 100)	(168; 321)
		1 Month		92.8%	952	74	67.6%	41.2
		Post dose 1	180	(88.0; 96.1)	(705; 1285)	71	(55.5; 78.2)	(23.7; 71.5)
	1 dose	1 Year	407	73.1%	76.8	00	91.9%	144
		Post dose 1	167	(65.7; 79.6)	(54.2; 109)	62	(82.2; 97.3)	(97.2; 215)
		1 Month	457	93.6%	933	50	64.3%	31.9
Y		Post dose 1	157	(88.6; 96.9)	(692; 1258)	56	(50.4; 76.6)	(17.6; 57.9)
	2 40000	1 Month Post		99.3%	1134		95.3%	513
	2 doses	dose 2	150	(96.3; 100)	(945; 1360)	64	(86.9; 99.0)	(339; 775)
		1 Year		79.7%	112		87.9%	144
		Post dose 2	143	(72.2; 86.0)	(77.5; 163)	58	(76.7; 95.0)	(88.5; 234)

The analysis of immunogenicity was conducted on the ATP cohort.

In Study MenACWY-TT-104, hSBA titres were measured as a secondary endpoint. In terms of the percentage of subjects with hSBA titres ≥8, at 1 month post vaccination, hSBA titres against groups W-135 and Y were higher after two doses of **Nimenrix**TM than after one dose, while the hSBA titres against groups A and C were similar in the two dose groups. At 1 year post vaccination, the percentages of subjects with hSBA titres ≥8 for all four meningococcal groups were similar in both the one and two dose groups (Table 6).

⁽¹⁾ blood sampling performed 21 to 48 days post vaccination and 44 to 60 weeks post vaccination

^{*} rSBA analysis performed at PHE laboratories

^{**}hSBA analysis performed at GSK laboratories

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In children vaccinated at toddler age, the persistence of rSBA and hSBA titres was evaluated up to 4 years in Study MenACWY-TT-048. Results are shown in Table 7.

Table 7: rSBA and hSBA titres up to 4 years following Nimenrix[™] (or MenC-CRM) in toddlers aged 12-23 months (Study MenACWY-TT-048)

Meningo-	W :	Time-		rSB <i>A</i>	\ *		hSBA**	
coccal group	Vaccine group	point (Years)	N	≥8 (95% CI)	GMT (95% CI)	N	≥8 (95% CI)	GMT (95% CI)
	Nimenrix	3	262	59.9% (53.7; 65.9)	19.3 (15.7; 23.6)	251	35.9% (29.9; 42.1)	5.8 (4.8; 7.0)
Α	ТМ	4	224	74.1% (67.9; 79.7)	107 (77.6; 148)	198	28.8% (22.6; 35.6)	4.9 (4.0; 6.0)
	Nimenrix	3	262	35.9% (30.1; 42.0)	9.8 (8.1; 11.7)	253	78.3% (72.7; 83.2)	37.8 (29.4; 48.6)
	тм	4	225	40.4% (34.0; 47.2)	12.3 (9.8; 15.3)	209	73.2% (66.7; 79.1)	32.0 (23.8; 43.0)
С	MenC-	3	46	13.0% (4.9; 26.3)	5.7 (4.2; 7.7)	31	41.9% (24.5; 60.9)	6.2 (3.7; 10.3)
	CRM vaccine	4	45	35.6% (21.9; 51.2)	13.5 (7.4; 24.5)	32	46.9% (29.1; 65.3)	11.3 (4.9; 25.6)
	Nimenrix	3	261	49.8% (43.6; 56.0)	24.9 (19.2; 32.4)	254	82.3% (77.0; 86.8)	52.0 (41.4; 65.2)
W-135	ТМ	4	225	49.3% (42.6; 56.1)	30.5 (22.4; 41.5)	165	80.6% (73.7; 86.3)	47.1 (35.7; 62.2)
Y	Nimenrix	3	262	53.8% (47.6; 60.0)	22.3 (17.6; 28.4)	250	72.0% (66.0; 77.5)	33.2 (25.9; 42.5)
	ТМ	4	225	58.2% (51.5; 64.7)	36.2 (27.1; 48.4)	130	65.4% (56.5; 73.5)	29.8 (20.2; 44.1)

The analysis of immunogenicity was conducted on the ATP cohort for persistence adapted for each time-point.

^{*} rSBA analysis performed at PHE laboratories in UK

^{**} hSBA analysis performed at GSK laboratories

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rSBA and hSBA titres were determined over a period of 10 years in children initially vaccinated with one dose of **Nimenrix**[™] or MenC-CRM at 12 to 23 months of age in Study MenACWY-TT-027. Persistence of SBA titres was evaluated in two extension studies: MenACWY-TT-032 (up to 5 years) and MenACWY-TT-100 (up to 10 years). Study MenACWY-TT-100 also evaluated the response to a single booster dose of **Nimenrix**[™] administered 10 years following the initial vaccination with **Nimenrix**[™] or MenC-CRM. Results are shown in Table 8 (see section 4.4).

Table 8: rSBA and hSBA titres following a single dose of Nimenrix[™] (or MenC-CRM) in toddlers aged 12-23 months, persistence up to 10 years, and post-booster administered 10 years following initial vaccination (Studies MenACWY-TT-027/032/100)

Meningo-				rSB/	*		hSB <i>A</i>	\ **
coccal group	Vaccine group	Time point	N	≥8 (95% CI)	GMT (95% CI)	N	≥8 (95% CI)	GMT (95% CI)
		Month 1 ⁽¹⁾	222	100% (98.4; 100)	3707 (3327; 4129)	217	91.2% (86.7; 94.6)	59.0 (49.3; 70.6)
	Nimenrix™	Year 4 ⁽²⁾	45	64.4% (48.8; 78.1)	35.1 (19.4; 63.4)	44	52.3% (36.7; 67.5)	8.8 (5.4; 14.2)
A		Year 5 ⁽²⁾	49	73.5% (58.9; 85.1)	37.4 (22.1; 63.2)	45	35.6% (21.9; 51.2)	5.2 (3.4; 7.8)
		Year 10 ⁽³⁾	62	66.1%	28.9	59	25.4%	4.2
		(Pre-booster)	62	98.4%	(16.4; 51.0) 5122	62	(15.0; 38.4) 100%	(3.0; 5.9)
		booster) ^(3,4) Month 1 ⁽¹⁾	220	(91.3; 100) 100%	(3726; 7043) 879	221	(94.2; 100) 99.1%	(1112; 2117) 190
		IVIORUT T	220	(98.3; 100) 97.8%	(779; 991) 110	221	(96.8; 99.9) 97.8%	(165; 219) 370
		Year 4 ⁽²⁾	45	(88.2; 99.9)	(62.7; 192)	45	(88.2; 99.9)	(214; 640)
С	Nimenrix [™]	Year 5 ⁽²⁾	49	77.6% (63.4; 88.2)	48.9 (28.5; 84.0)	48	91.7% (80.0; 97.7)	216 (124; 379)
		Year 10 ⁽³⁾	62	82.3%	128	60	91.7%	349
		(Pre-booster)	62	(70.5; 90.8) 100%	(71.1; 231) 7164	59	(81.6; 97.2) 100%	(197; 619)
		booster)(3,4)	02	(94.2; 100)	(5478; 9368)	199	(93.9; 100)	(23890; 48274)

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Table 8: rSBA and hSBA titres following a single dose of Nimenrix[™] (or MenC-CRM) in toddlers aged 12-23 months, persistence up to 10 years, and post-booster administered 10 years following initial vaccination (Studies MenACWY-TT-027/032/100)

Meningo-				rSB.	4 *		hSB <i>A</i>	\ **
coccal	Vaccine	Time point		≥8	GMT		≥8	GMT
group	group		N	(95% CI)	(95% CI)	N	(95% CI)	(95% CI)
				98.5%	415		72.1%	21.2
		Month 1 ⁽¹⁾	68	(92.1; 100)	(297; 580)	68	(59.9; 82.3)	(13.9; 32.3)
		4(2)	40	80.0%	137	40	70.0%	91.9
		Year 4 ⁽²⁾	10	(44.4; 97.5)	(22.6; 832)	10	(34.8; 93.3)	(9.8; 859)
	MenC-	V F(2)	44	63.6%	26.5	4.4	90.9%	109
	CRM	Year 5 ⁽²⁾	11	(30.8; 89.1)	(6.5; 107)	11	(58.7; 99.8)	(21.2; 557)
	vaccine	Year 10 ⁽³⁾	16	87.5%	86.7	15	93.3%	117
		(Pre-booster)	16	(61.7; 98.4)	(29.0; 259)	15	(68.1; 99.8)	(40.0; 344)
		(Post-	16	100%	5793	15	100%	42559
		booster)(3,4)	16	(79.4; 100)	(3631; 9242)	15	(78.2; 100)	(20106; 90086)
		Month 1 ⁽¹⁾	222	100%	5395	177	79.7%	38.8
		WOTH I	222	(98.4; 100)	(4870; 5976)	177	(73.0; 85.3)	(29.7; 50.6)
		Year 4 ⁽²⁾	45	60.0%	50.8	45	84.4%	76.9
		Teal 4	40	(44.3; 74.3)	(24.0; 108)	43	(70.5; 93.5)	(44.0; 134)
W-135	Nimenrix [™]	Vear 5 ⁽²⁾	49	34.7%	18.2	46	82.6%	59.7
77-133	Millellix	Teal 5	70	(21.7; 49.6)	(9.3; 35.3)	40	(68.6; 92.2)	(35.1; 101)
		Year 10 ⁽³⁾	62	30.6%	15.8	52	44.2%	7.7
		(Pre-booster)	02	(19.6; 43.7)	(9.1; 27.6)		(30.5; 58.7)	(4.9; 12.2)
		(Post-	62	100%	25911	62	100%	11925
		booster)(3,4)	02	(94.2; 100)	(19120; 35115)	02	(94.2; 100)	(8716; 16316)
		Month 1 ⁽¹⁾	222	100%	2824	201	66.7%	24.4
		WOTTET 1		(98.4; 100)	(2529; 3153)	201	(59.7; 73.1)	(18.6; 32.1)
		Year 4 ⁽²⁾	45	62.2%	44.9	41	87.8%	74.6
Y	Nimenrix™	10011	10	(46.5; 76.2)	(22.6; 89.3)		(73.8; 95.9)	(44.5; 125)
•		Year 5 ⁽²⁾	49	42.9%	20.6	45	80.0%	70.6
				(28.8; 57.8)	(10.9; 39.2)		(65.4; 90.4)	(38.7; 129)
		Year 10 ⁽³⁾	62	45.2%	27.4	56	42.9%	9.1
		(Pre-booster)	52	(32.5; 58.3)	(14.7; 51.0)	3	(29.7; 56.8)	(5.5; 15.1)

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Table 8: rSBA and hSBA titres following a single dose of Nimenrix[™] (or MenC-CRM) in toddlers aged 12-23 months, persistence up to 10 years, and post-booster administered 10 years following initial vaccination (Studies MenACWY-TT-027/032/100)

Meningo-	., .		rSBA*			hSBA**		
coccal	Vaccine	Time point	N	≥8	GMT	N	≥8	GMT
group	group		(95% CI)		(95% CI)	IN	(95% CI)	(95% CI)
		(Post-	00	98.4%	7661	0.4	100%	12154
		booster)(3,4)	62	(91.3; 100)	(5263; 11150)	61	(94.1; 100)	(9661; 15291

The analysis of immunogenicity was conducted on the ATP cohorts for 1 month and 5 years post vaccination and the booster ATP cohort.

- (1) Study MenACWY-TT-027
- (2) Study MenACWY-TT-032
- (3) Study MenACWY-TT-100
- (4) Blood sampling was performed 1 month after a booster dose at Year 10.

*rSBA analysis performed at GSK laboratories for 1 month post-primary vaccination samples and at PHE laboratories in UK for subsequent sampling time points.

**hSBA analysis performed at GSK laboratories and at Neomed in Canada for time points in Study MenACWY-TT-100.

Persistence of booster response

Study MenACWY-TT-102 evaluated the persistence of SBA titres up to 6 years after a booster dose of **Nimenrix**TM or MenC-CRM₁₉₇ administered in Study MenACWY-TT-048 to children who initially received the same vaccine at 12 to 23 months of age in Study MenACWY-TT-039. A single booster dose was administered 4 years after the initial vaccination. Results are shown in Table 9 (see section 4.4).

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Table 9: rSBA and hSBA titres following a single dose of Nimenrix[™] (or MenC-CRM) in toddlers aged 12-23 months, persistence at 4 years and response following a booster 4 years after initial vaccination, and persistence up to 6 years following booster vaccination (Studies MenACWY-TT-039/048/102)

Meningo-				rSBA*			hSBA*	r*
coccal	Vaccine	Time point		≥8	GMT		<u> </u> <	GMT
group	group		N	(95% CI)	(95% CI)	N	(95% CI)	(95% CI)
		A (1)	054	99.7%	2205	000	77.2%	19.0
		Month 1 ⁽¹⁾	354	(98.4; 100)	(2008; 2422)	338	(72.4; 81.6)	(16.4; 22.1)
		Year 4 ⁽²⁾		74.50/	440		00.00/	4.0
		(Pre-Nimenrix	212	74.5%	112	187	28.9%	4.8
		booster)		(68.1; 80.2)	(80.3; 156)		(22.5; 35.9)	(3.9; 5.9)
Α	Nimenrix TM	(Post-	044	100%	7173	000	99.5%	1343
		booster)(2,3)	214	(98.3; 100)	(6389; 8054)	202	(97.3; 100)	(1119; 1612)
		5 years after	407	89.8%	229	405	53.3%	13.2
		booster dose ⁽⁴⁾	137		(44.6; 62.0)	(9.6; 18.3)		
		6 years after	404	92.5%	297	400	58.5%	14.4
		booster dose ⁽⁴⁾	134	(86.7; 96.4)	(214; 413)	130	(49.5; 67.0)	(10.5; 19.7)
		NA (1 4 (1)	054	99.7%	478	341	98.5%	196
		Month 1 ⁽¹⁾	354	(98.4; 100)	(437; 522)	341	(96.6; 99.5)	(175; 219)
		Year 4 ⁽²⁾		39.9%	12.1		73.0%	31.2
		(Pre-Nimenrix	213			200		
		booster)		(33.3; 46.8)	(9.6; 15.2)		(66.3; 79.0)	(23.0; 42.2)
	Nimenrix	(Post-		100%	4512		100%	15831
С	ТМ	booster) ^(2,3)	215	(98.3; 100)	(3936; 5172)	209	(98.3; 100)	(13626;
		booster)		(90.3, 100)	(3930, 3172)		(90.3, 100)	18394)
		5 years after	137	80.3%	66.0	136	99.3%	337
		booster dose ⁽⁴⁾	131	(72.6; 86.6)	(48.1; 90.5)	130	(96.0; 100)	(261; 435)
		6 years after	134	71.6%	39.6	130	97.7%	259
		booster dose ⁽⁴⁾	104	(63.2; 79.1)	(28.6; 54.6)	130	(93.4; 99.5)	(195; 345)
	MenC-	Month 1 ⁽¹⁾	121	97.5%	212	116	81.9%	40.3
	CRM	MOHUL 157	141	(92.9; 99.5)	(170; 265)	110	(73.7; 88.4)	(29.5; 55.1)

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Table 9: rSBA and hSBA titres following a single dose of Nimenrix[™] (or MenC-CRM) in toddlers aged 12-23 months, persistence at 4 years and response following a booster 4 years after initial vaccination, and persistence up to 6 years following booster vaccination (Studies MenACWY-TT-039/048/102)

Meningo-	-			rSBA*			hSBA*	r*
coccal	Vaccine	Time point		≥8	GMT		≥8	GMT
group	group		N	(95% CI)	(95% CI)	N	(95% CI)	(95% CI)
	vaccine	Year 4 ⁽²⁾						
		(Pre-MenC-	42	37.2%	14.3	24	48.4%	11.9
		CRM ₁₉₇	43	(23.0; 53.3)	(7.7; 26.5)	31	(30.2; 66.9)	(5.1; 27.6)
		booster)						
		(Post-	42	100%	3718	22	100%	8646
		booster) ^(2,3)	43	(91.8; 100)	(2596; 5326)	33	(89.4; 100)	(5887; 12699)
		5 years after	23	78.3%	47.3	23	100%	241
		booster dose ⁽⁴⁾	23	(56.3; 92.5)	(19.0; 118)	23	(85.2; 100)	(139; 420)
		6 years after	23	65.2%	33.0	23	95.7%	169
		booster dose ⁽⁴⁾	23	(42.7; 83.6)	(14.7; 74.2)	23	(78.1; 99.9)	(94.1; 305)
		Month 1 ⁽¹⁾	354	100%	2682	336	87.5%	48.9
		WOTHT	304	(99.0; 100)	(2453; 2932)	330	(83.5; 90.8)	(41.2; 58.0)
		Year 4 ⁽²⁾		48.8%	30.2	158	81.6%	48.3
		(Pre-Nimenrix	213	(41.9; 55.7)	(21.9; 41.5)		(74.7; 87.3)	(36.5; 63.9)
		booster)		(11.0, 00.1)	(21.0, 11.0)		(7 1.17, 07 .0)	(00.0, 00.0)
W-135	Nimenrix	(Post-		100%	10950		100%	14411
	ТМ	booster) ^(2,3)	215	(98.3; 100)	(9531;	192	(98.1; 100)	(12972;
				(====, ===,	12579)		(====, ===,	16010)
		5 years after	137	88.3%	184	136	100%	327
		booster dose ⁽⁴⁾		(81.7; 93.2)	(130; 261)		(97.3; 100)	(276; 388)
		6 years after	134	85.8%	172	133	98.5%	314
		booster dose ⁽⁴⁾		(78.7; 91.2)	(118; 251)		(94.7; 99.8)	(255; 388)
	Nimenrix	Month 1 ⁽¹⁾	354	100%	2729	329	79.3%	30.9
				(99.0; 100)	(2473; 3013)		(74.5; 83.6)	(25.8; 37.1)
Y		Year 4 ⁽²⁾		58.2%	37.3		65.9%	30.2
		(Pre-Nimenrix	213	(51.3; 64.9)	(27.6; 50.4)	123	(56.8; 74.2)	(20.2; 45.0)
		booster)		, ,	(27.6; 50.4)		, ,	, ,

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Table 9: rSBA and hSBA titres following a single dose of Nimenrix[™] (or MenC-CRM) in toddlers aged 12-23 months, persistence at 4 years and response following a booster 4 years after initial vaccination, and persistence up to 6 years following booster vaccination (Studies MenACWY-TT-039/048/102)

Meningo-				rSBA*		hSBA**			
coccal	Vaccine	Time point	N	≥8	GMT	N	≥8	GMT	
group	group		IN	(95% CI)	(95% CI)	IN	(95% CI)	(95% CI)	
		(Post-	215	100%	4585	470	100%	6776	
		booster)(2,3)	215	(98.3; 100)	(4129; 5093)	173	(97.9; 100)	(5961; 7701)	
		5 years after	137	92.7%	265	137	97.8%	399	
		booster dose ⁽⁴⁾			(191; 368)	107	(93.7; 99.5)	(321; 495)	
		6 years after	S years after		260	131	97.7%	316	
		booster dose ⁽⁴⁾	134	(88.6; 97.4)	(189; 359)	131	(93.5; 99.5)	(253; 394)	

The analysis of immunogenicity was conducted on the ATP cohort for each time point.

- (1) Study MenACWY-TT-039
- (2) Study MenACWY-TT-048
- (3) Blood sampling was performed 1 month after a booster dose at Year 4.
- (4) Study MenACWY-TT-102

*rSBA analysis performed at GSK laboratories for 1 month post primary vaccination samples and at PHE laboratories in UK for the subsequent sampling time points.

**hSBA analysis performed at GSK laboratories and at Neomed in Canada for time points in Study MenACWY-TT-102.

Immune memory

In Study MenACWY-TT-014, the induction of immune memory was assessed 1 month after the administration of a fifth of the dose of ACWY-PS vaccine (10 µg of each polysaccharide) to children in the third year of life initially vaccinated in Study MenACWY-TT-013 with **Nimenrix**[™] or a licensed MenC-CRM vaccine at the age of 12 to 14 months.

One month after the challenge dose, the GMTs elicited by the initial vaccination with **Nimenrix**[™] increased by 6.5 to 8 fold for groups A, C, W-135, and Y, indicating that **Nimenrix**[™] induces immune memory to all four meningococcal groups. The post-challenge rSBA-MenC GMT was similar in both study groups, indicating that **Nimenrix**[™] induces an analogous immune memory to group C as the licensed MenC-CRM vaccine. Results are shown in Table 10.

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Table 10: rSBA* titres 1 month after a challenge vaccination in subjects initially vaccinated with Nimenrix[™] or a MenC-CRM vaccine at the age of 12 to 14 months (Study MenACWY-TT-014)

Meningo-			Pre-challenge		Post-challenge		
coccal	Vaccine group		GMT		GMT		
group		N	(95% CI)	N	(95% CI)		
Α	TM		544		3322		
	Nimenrix [™]	32	(325; 911)	25	(2294; 4810)		
	TM		174		5966		
	Nimenrix [™]	31	31 (105; 289) 32 (4128;				
С	MenC-CRM		34.4		5265		
	vaccine	28	(15.8; 75.3)	30	(3437; 8065)		
W-135	TM		644		11058		
	Nimenrix [™]	32	(394; 1052)	32	(8587; 14240)		
Y	TM		440		5737		
	Nimenrix [™]	32	(274; 706)	32	(4216; 7806)		

The analysis of immunogenicity was conducted on the ATP cohort.

Immunogenicity in children aged 2-10 years

In two comparative studies conducted in subjects aged 2-10 years, one group of subjects received a dose of **Nimenrix**[™] and a second group a dose of either a licensed MenC-CRM vaccine (Study MenACWY-TT-081) or the licensed ACWY-PS vaccine (Study MenACWY-TT-038) as comparator.

In Study MenACWY-TT-038, a single dose of **Nimenrix**[™] was demonstrated to be non-inferior to the licensed ACWY-PS vaccine in terms of vaccine response to the four meningococcal groups as shown in Table 11.

^{*} rSBA analysis performed at GSK laboratories

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Table 11: rSBA* titres following a single dose of to Nimenrix[™] (or ACWY-PS) in children aged 2-10 years (Study MenACWY-TT-038)

Mening		Nimen	rix ^{TM (1)}		ACWY-PS v	vaccine ⁽¹⁾
o-coccal group	N	VR (95% CI)	GMT (95% CI)	N	VR (95% CI)	GMT (95% CI)
group		, ,	<u> </u>		, ,	
Α	594	89.1%	6343	192	64.6%	2283
Α	JJ-4	(86.3; 91.5)	(5998; 6708)	132	(57.4; 71.3)	(2023; 2577)
		96.1%	4813		89.7%	1317
С	691	(94.4; 97.4)	(4342; 5335)	234	(85.1; 93.3)	(1043; 1663)
		97.4%	11543		82.6%	2158
W-135	691	(95.9; 98.4)	(10873; 12255)	236	(77.2; 87.2)	(1815; 2565)
V	700	92.7%	10825	0.40	68.8%	2613
Y	723	(90.5; 94.5)	(10233; 11452)	240	(62.5; 74.6)	(2237; 3052)

The analysis of immunogenicity was conducted on the ATP cohort.

VR: vaccine response defined as the proportion of subjects with:

- rSBA titres ≥32 for initially seronegative subjects (i.e., pre-vaccination rSBA titre <8)
- at least a 4-fold increase in rSBA titres from pre- to post vaccination for initially seropositive subjects (i.e., pre-vaccination rSBA titre ≥8)

In Study MenACWY-TT-081, a single dose of **Nimenrix[™]** (N=268) was demonstrated to be non-inferior to another licensed MenC-CRM vaccine (N=92) in terms of vaccine response to group C [94.8% (95% CI: 91.4; 97.1) and 95.7% (95% CI: 89.2; 98.8), respectively], GMTs were lower for the **Nimenrix[™]** group [2795 (95% CI: 2393; 3263)] versus the MenC-CRM vaccine [5292 (95% CI: 3815; 7340)].

In Study MenACWY-TT-088, the persistence of SBA titres was evaluated up to 68 months after vaccination in children 2-10 years of age initially vaccinated in Study MenACWY-TT-081. Results are shown in Table 12 (see section 4.4).

⁽¹⁾ Blood sampling performed 1 month post vaccination

^{*}rSBA analysis performed at GSK laboratories

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Table 12: rSBA and hSBA titres up to 68 months following Nimenrix[™] (or MenC-CRM) in children aged 2-10 years at time of vaccination (Study MenACWY-TT-088)

Meningo-				rSBA*			hSBA*	*
coccal group	Vaccine group	Time-point (months)	N	≥8 (95% CI)	GMT (95% CI)	N***	≥8 (95% CI)	GMT (95% CI)
	TM	32	193	86.5% (80.9; 91.0)	196 (144; 267)	90	25.6% (16.9; 35.8)	4.6 (3.3; 6.3)
A	Nimenrix™	68	178	86.5% (80.6; 91.2)	129 (93.5; 179)	170	40.6% (33.1; 48.4)	6.9 (5.4; 8.9)
	тм	32	192	64.6% (57.4; 71.3)	34.8 (26.0; 46.4)	90	95.6% (89.0; 98.8)	75.9 (53.4; 108)
	Nimenrix [™]	68	178	39.9% (32.6; 47.5)	14.2 (10.8; 18.7)	172	75.6% (68.5; 81.8)	28.4 (21.2; 37.9)
С	MenC-CRM	32	69	76.8% (65.1; 86.1)	86.5 (47.3; 158)	33	90.9% (75.7; 98.1)	82.2 (34.6; 196)
	vaccine	68	61	62.3% (49.0; 74.4)	44.5 (23.7; 83.6)	57	75.4% (62.2; 85.9)	34.3 (19.0; 61.9)
	TM	32	193	77.2% (70.6; 82.9)	214 (149; 307)	86	84.9% (75.5; 91.7)	69.9 (48.2; 101)
W-135	Nimenrix™	68	178	52.8% (45.2; 60.3)	59.2 (39.3; 89.2)	159	78.6% (71.4; 84.7)	56.7 (41.5; 77.3)
		32	193	81.3% (75.1; 86.6)	227 (165; 314)	91	81.3% (71.8; 88.7)	79.2 (52.5; 119)
Y	Y Nimenrix [™]	68	178	71.3% (64.1; 77.9)	139 (96.0; 202)	159	73.0% (65.3; 79.7)	56.3 (39.5; 80.3)

The analysis of immunogenicity was conducted on the ATP cohort for persistence adapted for each time-point.

In Study MenACWY-TT-028, the persistence of hSBA titres was evaluated 1 year after vaccination in children aged 6-10 years who were initially vaccinated in Study MenACWY-TT-027. Results are shown in Table 13

^{*} rSBA analysis performed at PHE laboratories in UK

^{**} hSBA analysis performed at GSK laboratories

^{***} at Month 32, a subset of subjects has been tested for hSBA

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Table 13: hSBA* titres following a single dose of Nimenrix[™] (or ACWY-PS) in children aged 6-10 years and persistence 1 year following vaccination (Studies MenACWY-TT-027/028)

Meningo	-		1 month post vac	ccination		1 year pers	sistence
	Vaccine	(Study MenACW\	/-TT-027)		(Study MenAC	WY-TT-028)
-coccal	group	N	≥8	GMT	N.	≥8	GMT
group		N	(95% CI)	(95% CI)	N	(95% CI)	(95% CI)
	Nimenrix™	105	80.0%	53.4	104	16.3%	3.5
	Milliellix	103	(71.1; 87.2)	(37.3; 76.2)	104	(9.8; 24.9)	(2.7; 4.4)
Α	ACWY-PS	35	25.7%	4.1	35	5.7%	2.5
	vaccine	33	(12.5; 43.3)	(2.6; 6.5)	33	(0.7; 19.2)	(1.9; 3.3)
	Nimenrix™	101	89.1%	156	105	95.2%	129
		101	(81.3; 94.4)	(99.3; 244)	103	(89.2; 98.4)	(95.4; 176)
С	ACWY-PS	38	39.5%	13.1	31	32.3%	7.7
	vaccine	30	(24.0; 56.6)	(5.4; 32.0)	31	(16.7; 51.4)	(3.5; 17.3)
	Nimenrix™	103	95.1%	133	103	100%	257
	Nimennx	103	(89.0; 98.4)	(99.9; 178)	103	(96.5; 100)	(218; 302)
W-135	ACWY-PS	25	34.3%	5.8	24	12.9%	3.4
	vaccine	35	(19.1; 52.2)	(3.3; 9.9)	31	(3.6; 29.8)	(2.0; 5.8)
	N: TM	00	83.1%	95.1	400	99.1%	265
	Nimenrix [™]	89	(73.7; 90.2)	(62.4; 145)	106	(94.9; 100)	(213; 330)
Y	ACWY-PS	20	43.8%	12.5	26	33.3%	9.3
	vaccine	32	(26.4; 62.3)	(5.6; 27.7)	36	(18.6; 51.0)	(4.3; 19.9)

The analysis of immunogenicity was conducted on the ATP cohort for persistence at Year 1.

hSBA analysis was not performed for children aged 2 to <6 years (at time of vaccination).

SBA titres were determined over a period of 10 years in children initially vaccinated with one dose of **Nimenrix**[™] or ACWY-PS at 2 to 10 years of age in Study MenACWY-TT-027. Persistence of SBA titres was evaluated in two extension studies: MenACWY-TT-032 (up to 5 years) and MenACWY-TT-100 (up to 10 years). Study MenACWY-TT-100 also evaluated the response to a single booster dose of **Nimenrix**[™] administered 10 years following the initial vaccination with **Nimenrix**[™] or ACWY-PS. Results are shown in Table 14 (see section 4.4).

^{*} hSBA analysis performed at GSK laboratories

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Table 14: rSBA and hSBA titres following a single dose of Nimenrix[™] (or ACWY-PS) in children aged 2-10 years, persistence up to 10 years, and post-booster administered 10 years following initial vaccination (Studies MenACWY-TT-027/032/100)

Meningo-	•	ludies MeliAC		rSB			hSBA	**
coccal	Vaccine	Time point		≥8	GMT		≥8	GMT
group	group		N	(95% CI)	(95% CI)	N	(95% CI)	(95% CI)
				100%	7301	4.4.(5)	81.1%	57.0
		Month 1 ⁽¹⁾	225	(98.4; 100)	(6586; 8093)	111 ⁽⁵⁾	(72.5; 87.9)	(40.3; 80.6)
		(2)	00	90.8%	141	, (6)		
		Year 5 ⁽²⁾	98	(83.3; 95.7)	(98.2; 203)	n/a ⁽⁶⁾		
	N: TM	V C(3)	00	79.6%	107	00	41.1%	6.5
	Nimenrix [™]	Year 6 ⁽³⁾	98	(70.3; 87.1)	(66.0; 174)	90	(30.8; 52.0)	(4.8; 8.8)
		Year 10 ⁽³⁾	70	89.0%	96.3	00	33.9%	4.5
		(Pre-booster)	73	(79.5; 95.1)	(57.1; 163)	62	(22.3; 47.0)	(3.3; 6.2)
		(Post-		95.9%	4626	72	100%	1213
		booster)(3,4)	74	(88.6; 99.2)	(3041; 7039)	73	(95.1; 100)	(994; 1481)
Α		Month 1 ⁽¹⁾	75	100%	2033	35 ⁽⁵⁾	25.7%	4.1
		WOTH IN	75	(95.2; 100)	(1667; 2480)	35.7	(12.5; 43.3)	(2.6; 6.5)
		Year 5 ⁽²⁾	13	15.4%	4.7	n/a ⁽⁶⁾		
		rear 5.7		(1.9; 45.4)	(3.7; 6.0)	II/a**		
	ACWY-PS	Year 6 ⁽³⁾	24	12.5%	5.8	21	33.3%	5.9
	vaccine	Teal 0	24	(2.7; 32.4)	(3.5; 9.6)	21	(14.6; 57.0)	(3.0; 11.7)
		Year 10 ⁽³⁾	17	23.5%	8.0	17	29.4%	6.2
		(Pre-booster)	.,	(6.8; 49.9)	(3.3; 19.3)	.,	(10.3; 56.0)	(2.4; 15.7)
		(Post-	17	100%	6414	17	100%	211
		booster)(3,4)	1,	(80.5; 100)	(3879; 10608)	1,	(80.5; 100)	(131; 340)
		Month 1 ⁽¹⁾	225	100%	2435	107 ⁽⁵⁾	89.7%	155
		WOITH 1	220	(98.4; 100)	(2106; 2816)	107	(82.3; 94.8)	(101; 237)
		Year 5 ⁽²⁾	98	90.8%	79.7	n/a ⁽⁶⁾		
С	Nimenrix [™]		30	(83.3; 95.7)	(56.0; 113)	11/4		
	Annemia	Year 6 ⁽³⁾	98	82.7%	193	97	93.8%	427
		1 501 0		(73.7; 89.6)	(121; 308)	0,	(87.0; 97.7)	(261; 700)
		Year 10 ⁽³⁾	74	85.1%	181	73	91.8%	222
		(Pre-booster)	′+	(75.0; 92.3)	(106; 310)	73	(83.0; 96.9)	(129; 380)

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Table 14: rSBA and hSBA titres following a single dose of Nimenrix[™] (or ACWY-PS) in children aged 2-10 years, persistence up to 10 years, and post-booster administered 10 years following initial vaccination (Studies MenACWY-TT-027/032/100)

Meningo-				rSB	A *		hSBA	* *
coccal	Vaccine	Time point		≥8	GMT		≥8	GMT
group	group		N	(95% CI)	(95% CI)	N	(95% CI)	(95% CI)
		(Post-	7.4	100%	4020	74	100%	15544
		booster)(3,4)	74	(95.1; 100)	(3319; 4869)	71	(94.9; 100)	(11735; 20588)
		Month 1 ⁽¹⁾	74	100%	750	38 ⁽⁵⁾	39.5%	13.1
		MONUTE IN 7	74	(95.1; 100)	(555; 1014)	30\'/	(24.0; 56.6)	(5.4; 32.0)
		Year 5 ⁽²⁾	13	100%	128	n/a ⁽⁶⁾		
				(75.3; 100)	(56.4; 291)			
	ACWY-PS	Year 6 ⁽³⁾	24	79.2%	98.7	24	100%	235
	vaccine			(57.8; 92.9)	(42.2; 231)		(85.8; 100)	(122; 451)
		Year 10 ⁽³⁾	17	76.5%	96.2	17	100%	99.1
		(Pre-booster)		(50.1; 93.2)	(28.9; 320)		(80.5; 100)	(35.8; 274)
		(Post-	17	100%	15101	17	94.1	44794
		booster)(3,4)		(80.5; 100)	(7099; 32122)		(71.3; 99.9)	(10112; 198440)
		Month 1 ⁽¹⁾	225	100%	11777	107 ⁽⁵⁾	95.3%	134
		Wieriar		(98.4; 100)	(10666; 13004)	107	(89.4; 98.5)	(101; 178)
		Year 5 ⁽²⁾	98	78.6%	209	n/a ⁽⁶⁾		
		Tour o		(69.1; 86.2)	(128; 340)	Ti/G		
	Nimenrix [™]	Year 6 ⁽³⁾	98	73.5%	265	92	81.5%	62.5
	Millicilia	Tour o	00	(63.6; 81.9)	(155; 454)	02	(72.1; 88.9)	(42.0; 93.1)
		Year 10 ⁽³⁾	74	68.9%	206	59	61.0%	17.5
W-135		(Pre-booster)	74	(57.1; 79.2)	(109; 392)	33	(47.4; 73.5)	(10.5; 29.2)
W-133		(Post-	74	100%	27944	74	100%	6965
		booster) ^(3,4)	74	(95.1; 100)	(22214; 35153)	74	(95.1; 100)	(5274; 9198)
		Month 1 ⁽¹⁾	75	100%	2186	35 ⁽⁵⁾	34.3%	5.8
		IVIOHUI 1\'	75	(95.2; 100)	(1723; 2774)	30\	(19.1; 52.2)	(3.3; 9.9)
	ACWY-PS	Year 5 ⁽²⁾	12	0%	4.0	, (6)		
	vaccine	Teal of	13	(0.0; 24.7)	(4.0; 4.0)	n/a ⁽⁶⁾		-
		Year 6 ⁽³⁾	24	12.5%	7.6	23	30.4%	7.0
		Teal U.	4	(2.7; 32.4)	(3.7; 15.6)	23	(13.2; 52.9)	(2.9; 16.9)

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Table 14: rSBA and hSBA titres following a single dose of Nimenrix[™] (or ACWY-PS) in children aged 2-10 years, persistence up to 10 years, and post-booster administered 10 years following initial vaccination (Studies MenACWY-TT-027/032/100)

Meningo-				rSB	A*		hSBA	**
coccal	Vaccine	Time point		≥8	GMT		≥8	GMT
group	group		N	(95% CI)	(95% CI)	N	(95% CI)	(95% CI)
		Year 10 ⁽³⁾	17	23.5%	15.4	15	26.7%	4.1
		(Pre-booster)	17	(6.8; 49.9)	(4.2; 56.4)	15	(7.8; 55.1)	(2.0; 8.5)
		(Post-	17	94.1%	10463	15	100%	200
		booster)(3,4)	17	(71.3; 99.9)	(3254; 33646)	15	(78.2; 100)	(101; 395)
		Month 1 ⁽¹⁾	225	100%	6641	94 ⁽⁵⁾	83.0%	93.7
		Month 107	225	(98.4; 100)	(6044; 7297)	94\-'	(73.8; 89.9)	(62.1; 141)
		Year 5 ⁽²⁾	00	78.6%	143	n/a ⁽⁶⁾		
		Year 5\'/	98	(69.1; 86.2)	(88.0; 233)	n/a\		-
	Nimenrix™	Year 6 ⁽³⁾	98	71.4%	136	89	65.2%	40.3
	Nimema	real 0\7	90	(61.4; 80.1)	(82.6; 225)	09	(54.3; 75.0)	(23.9; 68.1)
		Year 10 ⁽³⁾	74	67.6%	98.5	65	72.3%	35.7
		(Pre-booster)		(55.7; 78.0)	(54.3; 179)	00	(59.8; 82.7)	(21.0; 60.6)
		(Post-	7.4	100%	7530	74	100%	11127
Y		booster)(3,4)	74	(95.1; 100)	(5828; 9729)	74	(95.1; 100)	(8909; 13898)
Y		Month 1 ⁽¹⁾	75	100%	1410	32 ⁽⁵⁾	43.8%	12.5
		MOHIT 117	75	(95.2; 100)	(1086; 1831)	32.7	(26.4; 62.3)	(5.6; 27.7)
		Year 5 ⁽²⁾	13	7.7%	5.5	n/a ⁽⁶⁾		
		Teal 5	13	(0.2; 36.0)	(2.7; 11.1)	11/a		
	ACWY-PS	Year 6 ⁽³⁾	24	20.8%	11.6	24	25.0%	7.3
	vaccine	Teal o	24	(7.1; 42.2)	(4.7; 28.7)	24	(9.8; 46.7)	(2.7; 19.8)
		Year 10 ⁽³⁾	17	17.6%	10.2	14	35.7%	7.8
		(Pre-booster)	''	(3.8; 43.4)	(3.5; 30.2)	14	(12.8; 64.9)	(2.5; 24.4)
		(Post-	17	100%	6959	17	100%	454
		booster)(3,4)	17	(80.5; 100)	(3637; 13317)	17	(80.5; 100)	(215; 960)

The analysis of immunogenicity was conducted on the ATP cohort for each time point.

- (1) Study MenACWY-TT-027
- (2) Study MenACWY-TT-032
- (3) Study MenACWY-TT-100

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- (4) Blood sampling was performed 1 month after a booster dose at Year 10.
- (5) Includes children aged 6 to <11 years. hSBA analysis was not performed for children aged 2 to <6 years (at time of vaccination).
- (6) Per the protocol for Study MenACWY-TT-032, hSBA was not measured for this age group at Year 5.
- *rSBA analysis performed at GSK laboratories for 1 month post primary vaccination samples and at PHE laboratories in UK for subsequent sampling time points.
- **hSBA analysis performed at GSK laboratories and at Neomed in Canada for time points in Study MenACWY-TT-100.

Immunogenicity in adolescents aged 11-17 years and adults aged ≥18 years

In two clinical studies, conducted in adolescents aged 11-17 years (Study MenACWY-TT-036) and in adults aged 18-55 years (Study MenACWY-TT-035), either one dose of **Nimenrix**[™] or one dose of the ACWY-PS vaccine was administered.

In both adolescents and adults, **Nimenrix**[™] was demonstrated to be immunologically non-inferior to the ACWY-PS vaccine in terms of vaccine response. rSBA titres to the four meningococcal groups elicited by **Nimenrix**[™] were either similar to or higher than those elicited by the ACWY-PS vaccine as shown in Table 15.

Table 15: rSBA*titres following a single dose of Nimenrix[™] (or ACWY-PS) in adolescents aged 11-17 years and adults aged 18-55 years (Studies MenACWY-TT-035/036)

Manin			Study MenACV	VY-TT-036		Study MenAC	WY-TT-035				
Meningo-	Vaccine		(11-17 yea	ars) ⁽¹⁾		(18-55 years) ⁽¹⁾					
coccal	group		VR GMT			VR	GMT				
group		N	(95% CI)	(95% CI)	N	(95% CI)	(95% CI)				
	Nimenrix [™]	553	85.4%	5928	743	80.1%	3625				
		333	(82.1; 88.2)	(5557; 6324)	743	(77.0; 82.9)	(3372; 3897)				
A	ACWY-PS	191	77.5%	2947	252	69.8%	2127				
	vaccine	191	(70.9; 83.2)	(2612; 3326)	232	(63.8; 75.4)	(1909; 2370)				
	Nimenrix [™]	642	97.4%	13110	849	91.5%	8866				
		042	(95.8; 98.5)	(11939; 14395)	049	(89.4; 93.3)	(8011; 9812)				
С	ACWY-PS	211	96.7%	8222	288	92.0%	7371				
	vaccine	211	(93.3; 98.7)	(6807; 9930)	200	(88.3; 94.9)	(6297; 8628)				
W-135	Nimenrix [™]	639	96.4%	8247	860	90.2%	5136				

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Table 15: rSBA*titres following a single dose of Nimenrix[™] (or ACWY-PS) in adolescents aged 11-17 years and adults aged 18-55 years (Studies MenACWY-TT-035/036)

Meningo-	Vaccine	Study MenACWY-TT-036 (11-17 years) ⁽¹⁾			Study MenACWY-TT-035 (18-55 years) ⁽¹⁾			
coccal group	group	N	VR (95% CI)	GMT (95% CI)	N	VR (95% CI)	GMT (95% CI)	
			(94.6; 97.7)	(7639; 8903)		(88.1; 92.1)	(4699; 5614)	
	ACWY-PS	216	87.5%	2633	283	85.5%	2461	
	vaccine	210	(82.3; 91.6)	(2299; 3014)	200	(80.9; 89.4)	(2081; 2911)	
	Nimenrix™	657	93.8%	14086	862	87.0%	7711	
		007	(91.6; 95.5)	(13168; 15069)	002	(84.6; 89.2)	(7100; 8374)	
Y	ACWY-PS	219	78.5%	5066	288	78.8%	4314	
	vaccine	213	(72.5; 83.8)	(4463; 5751)	200	(73.6; 83.4)	(3782; 4921)	

The analysis of immunogenicity was conducted on the ATP cohorts.

(1) Blood sampling performed 1 month post vaccination

VR: vaccine response defined as the proportion of subjects with:

- rSBA titres ≥32 for initially seronegative subjects (i.e., pre-vaccination rSBA titre <8)
- at least a 4-fold increase in rSBA titres from pre- to post vaccination for initially seropositive subjects (i.e., pre-vaccination rSBA titre ≥8)

rSBA titres were determined over a period of 10 years in subjects initially vaccinated with one dose of **Nimenrix**[™] or ACWY-PS at 11 to 17 years of age in Study MenACWY-TT-036. Persistence of rSBA titres was evaluated in two extension studies: MenACWY-TT-043 (up to 5 years) and MenACWY-TT-101 (at 10 years). Study MenACWY-TT-101 also evaluated the response to a single booster dose of **Nimenrix**[™] administered 10 years following the initial vaccination with **Nimenrix**[™] or ACWY-PS. Results are shown in Table 16.

^{*}rSBA analysis performed at GSK laboratories

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Table 16: rSBA* titres following a single dose of Nimenrix[™] (or ACWY-PS) in adolescents aged 11-17 years, persistence up to 10 years, and post-booster administered 10 years following initial vaccination (Studies MenACWY-TT-036/043/101)

Meningo-		Nimenrix™				ACWY-PS vaccine			
coccal	Time point		≥8	GMT		≥8	GMT		
group		N	(95% CI)	(95% CI)	N	(95% CI)	(95% CI)		
		074	100%	5929	004	99.6%	2947		
	Month 1 ⁽¹⁾	674	(99.5; 100)	(5557; 6324)	224	(97.5; 100)	(2612; 3326)		
	Year 3 ⁽²⁾	449	92.9%	448	150	82.7%	206		
	rear 5\	449	(90.1; 95.1)	(381; 527)	150	(75.6; 88.4)	(147; 288)		
A	Year 5 ⁽²⁾	236	97.5%	644	86	93.0%	296		
_ ^	rear 5\	230	(94.5; 99.1)	(531; 781)	00	(85.4; 97.4)	(202; 433)		
	Year 10 ⁽³⁾	162	85.2%	248	51	80.4%	143		
	(Pre-booster)	102	(78.8; 90.3)	(181; 340)	51	(66.9; 90.2)	(80.5; 253)		
	(Post-	162	100%	3760	51	100%	2956		
	booster)(3,4)	102	(97.7; 100)	(3268; 4326)	31	(93.0; 100)	(2041; 4282)		
	Month 1 ⁽¹⁾	673	100%	13110	224	100%	8222		
			(99.5; 100)	(11939; 14395)	224	(98.4; 100)	(6808; 9930)		
	Year 3 ⁽²⁾	449	91.1%	371	150	86.0%	390		
			(88.1; 93.6)	(309; 446)	150	(79.4; 91.1)	(262; 580)		
С	Year 5 ⁽²⁾	236	88.6%	249	85	87.1%	366		
			(83.8; 92.3)	(194; 318)	65	(78.0; 93.4)	(224; 599)		
	Year 10 ⁽³⁾	162	90.1%	244	51	82.4%	177		
	(Pre-booster)	102	(84.5; 94.2)	(182; 329)	31	(69.1; 91.6)	(86.1; 365)		
	(Post-	162	100%	8698	51	100%	3879		
	booster)(3,4)	102	(97.7; 100)	(7391; 10235)	31	(93.0; 100)	(2715; 5544)		
	Month 1 ⁽¹⁾	678	99.9%	8247	224	100%	2633		
	MOHUTTY	070	(99.2; 100)	(7639; 8903)	224	(98.4; 100)	(2299; 3014)		
	Year 3 ⁽²⁾	449	82.0%	338	150	30.0%	16.0		
W-135	real 5	449	(78.1; 85.4)	(268; 426)	150	(22.8; 38.0)	(10.9; 23.6)		
VV-135	Year 5 ⁽²⁾	236	86.0%	437	98	34.9%	19.7		
	Teal 3'	230	(80.9; 90.2)	(324; 588)	86	(24.9; 45.9)	(11.8; 32.9)		
	Year 10 ⁽³⁾	162	71.6%	146	51	43.1%	16.4		
	(Pre-booster)	102	(64.0; 78.4)	(97.6; 217)	31	(29.3; 57.8)	(9.2; 29.4)		

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Meningo-		Nimenrix™			ACWY-PS vaccine			
coccal group	Time point	N	≥8 (95% CI)	GMT (95% CI)	N	≥8 (95% CI)	GMT (95% CI)	
	(Post-booster) ^(3,4)	162	100% (97.7; 100)	11243 (9367; 13496)	51	100% (93.0; 100)	3674 (2354; 5734)	
	Month 1 ⁽¹⁾	677	100% (99.5; 100)	14087 (13168; 15069)	224	100% (98.4; 100)	5066 (4463; 5751)	
	Year 3 ⁽²⁾	449	93.1% (90.3; 95.3)	740 (620; 884)	150	58.0% (49.7; 66.0)	69.6 (44.6; 109)	
Y	Year 5 ⁽²⁾	236	96.6% (93.4; 98.5)	3.6% 1000 86	86	66.3% (55.3; 76.1)	125 (71.2; 219)	
	Year 10 ⁽³⁾ (Pre-booster)	162	90.7% (85.2; 94.7)	447 (333; 599)	51	49.0% (34.8; 63.4)	32.9 (17.1; 63.3)	
	(Post-booster) ^(3,4)	162	100% (97.7; 100)	7585 (6748; 8525)	51	98.0% (89.6; 100)	3296 (1999; 5434)	

The analysis of immunogenicity was conducted on the ATP cohort for each time point.

- (1) Study MenACWY-TT-036
- (2) Study MenACWY-TT-043
- (3) Study MenACWY-TT-101
- (4) Blood sampling was performed 1 month after a booster dose at Year 10.

*rSBA analysis performed at GSK laboratories for 1 month post primary vaccination samples and at PHE laboratories in UK for the subsequent sampling time points.

In Study MenACWY-TT-059, hSBA persistence was evaluated up to 5 years after vaccination in adolescents and adults aged 11-25 years initially vaccinated in Study MenACWY-TT-052.

For all meningococcal groups, the persistence of hSBA titres elicited by **Nimenrix**[™] was similar to or higher than those induced by the licensed quadrivalent meningococcal diphtheria toxoid (DT) conjugate (ACWY-DT) vaccine as shown in Table 17.

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Table 17: hSBA* titres following a single dose of Nimenrix[™] (or ACWY-DT) in adolescents and adults aged 11-25 years and persistence up to 5 years following vaccination (Studies MenACWY-TT-052/059)

(Studies Menacy					
Meningo-coccal group	Vaccine group	Time-point	N	≥8 (95% CI)	GMT (95% CI)
		Month 1 ⁽¹⁾	356	82.0% (77.6; 85.9)	58.7 (48.6; 70.9)
	Nimenrix™	Year 1 ⁽²⁾	350	29.1% (24.4; 34.2)	5.4 (4.5; 6.4)
_		Year 5 ⁽²⁾	141	48.9% (40.4; 57.5)	8.9 (6.8; 11.8)
Α		Month 1 ⁽¹⁾	107	73.8% (64.4; 81.9)	42.5 (28.5; 63.3)
	ACWY-DT	Year 1 ⁽²⁾	111	31.5% (23.0; 41.0)	6.0 (4.3; 8.5)
		Year 5 ⁽²⁾	45	44.4% (29.6; 60.0)	7.9 (4.8; 13.2)
		Month 1 ⁽¹⁾	359	96.1% (93.5; 97.9)	532 (424; 668)
	Nimenrix™	Year 1 ⁽²⁾	336	94.9% (92.0; 97.0)	172 (142; 207)
		Year 5 ⁽²⁾	140	92.9% (87.3; 96.5)	94.6 (65.9; 136)
С		Month 1 ⁽¹⁾	113	99.1% (95.2; 100)	317 (217; 462)
	ACWY-DT	Year 1 ⁽²⁾	105	73.3% (63.8; 81.5)	46.7 (30.2; 72.1)
		Year 5 ⁽²⁾	44	79.5% (64.7; 90.2)	30.6 (17.3; 54.4)
		Month 1 ⁽¹⁾	334	91.0% (87.4; 93.9)	117 (96.8; 141)
	Nimenrix [™]	Year 1 ⁽²⁾	327	98.5% (96.5; 99.5)	197 (173; 225)
W 405		Year 5 ⁽²⁾	138	87.0% (80.2; 92.1)	103 (76.3; 140)
W-135		Month 1 ⁽¹⁾	96	75.0% (65.1; 83.3)	70.4 (43.7; 113)
	ACWY-DT	Year 1 ⁽²⁾	107	75.7% (66.5; 83.5)	48.9 (32.5; 73.8)
		Year 5 ⁽²⁾	44	84.1% (69.9; 93.4)	70.4 (37.2; 133)
		Month 1 ⁽¹⁾	364	95.1% (92.3; 97.0)	246 (208; 291)
Y	Nimenrix™	Year 1 ⁽²⁾	356	97.8% (95.6; 99.0)	272 (237; 311)
		Year 5 ⁽²⁾	142	94.4% (89.2; 97.5)	225 (174; 290)
		Month 1 ⁽¹⁾	111	81.1% (72.5; 87.9)	103 (67.5; 159)
	ACWY-DT	Year 1 ⁽²⁾	112	86.6% (78.9; 92.3)	101 (69.6; 146)
		Year 5 ⁽²⁾	44	90.9% (78.3; 97.5)	129 (77.4; 216)

The analysis of immunogenicity was conducted on the ATP cohort for persistence adapted for each time-point.

- (1) Study MenACWY-TT-052
- (2) Study MenACWY-TT-059

^{*} hSBA analysis performed at GSK laboratories

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rSBA titres were determined over a period of 10 years in subjects initially vaccinated with one dose of **Nimenrix**[™] or ACWY-PS at 11 to 55 years of age in Study MenACWY-TT-015.

Persistence of rSBA titres was evaluated in two extension studies: MenACWY-TT-020 (up to 5 years) and MenACWY-TT-099 (up to 10 years). Study MenACWY-TT-099 also evaluated the response to a single booster dose of **Nimenrix**[™] administered 10 years following the initial vaccination with **Nimenrix**[™] or ACWY-PS. Results are shown in Table 18.

Table 18: rSBA* titres following a single dose of Nimenrix[™] (or ACWY-PS) in adolescents and adults aged 11-55 years, persistence up to 10 years, and post-booster administered 10 years following initial vaccination (Studies MenACWY-TT-015/020/099)

Meningo-			Nimenri	x TM	ACWY-PS vaccine			
coccal	Time point	N	≥8 GMT			≥8	GMT	
group	-		(95% CI)	(95% CI)	N	(95% CI)	(95% CI)	
	(1)		100%	4945		100%	2190	
	Month 1 ⁽¹⁾	323	(98.9; 100)	(4452; 5493)	112	(96.8; 100)	(1858; 2582)	
	. (2)		95.3%	365		76.5%	104	
	Year 4 ⁽²⁾	43	(84.2; 99.4)	(226; 590)	17	(50.1; 93.2)	(31.0; 351)	
	- (2)		84.3%	190	40	57.9%	37.0	
Α	Year 5 ⁽²⁾	51	(71.4; 93.0)	(108; 335)	19	(33.5; 79.7)	(12.6; 109)	
	Year 10 ⁽³⁾	455	78.1%	154	5 0	71.2%	75.1	
	(Pre-booster)	155	(70.7; 84.3)	(108; 219)	52	(56.9; 82.9)	(41.4; 136)	
	(Post-	455	100%	4060		100%	3585	
	booster)(3,4)	155	(97.6; 100)	(3384; 4870)	52	(93.2; 100)	(2751; 4672)	
	Month 1 ⁽¹⁾	341	99.7%	10074	444	100%	6546	
			(98.4; 100)	(8700, 11665)	114	(96.8; 100)	(5048; 8488)	
	1(2)	40	76.7%	126	47	41.2%	16.7	
	Year 4 ⁽²⁾	43	(61.4; 88.2)	(61.6; 258)	17	(18.4; 67.1)	(5.7; 48.7)	
	V =(2)	5 4	72.5%	78.5	40	38.9%	17.3	
С	Year 5 ⁽²⁾	51	(58.3; 84.1)	(41.8; 147)	18	(17.3; 64.3)	(6.0; 49.7)	
	Year 10 ⁽³⁾	454	90.9%	193	F0	88.5%	212	
	(Pre-booster)	154	(85.2; 94.9)	(141; 264)	52	(76.6; 95.6)	(110; 412)	
	(Post-	155	100%	13824	5 0	98.1%	3444	
	booster)(3,4)	155	(97.6; 100)	(10840; 17629)	52	(89.7; 100)	(1999; 5936)	

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Table 18: rSBA* titres following a single dose of Nimenrix[™] (or ACWY-PS) in adolescents and adults aged 11-55 years, persistence up to 10 years, and post-booster administered 10 years following initial vaccination (Studies MenACWY-TT-015/020/099)

Meningo-			Nimenri	x TM		ACWY-PS	vaccine
coccal	Time point		≥8 GMT			≥8	GMT
group		N	(95% CI)	(95% CI)	N	(95% CI)	(95% CI)
	(1)		99.7%	8577		100%	2970
	Month 1 ⁽¹⁾	340	(98.4; 100)	(7615; 9660)	114	(96.8; 100)	(2439; 3615)
	M 4(2)	40	90.7%	240	47	17.6%	8.3
	Year 4 ⁽²⁾	43	(77.9; 97.4)	(128; 450)	17	(3.8; 43.4)	(3.6; 19.5)
)o.	N =(2)		86.3%	282	40	31.6%	15.4
W-135	5 Year 5 ⁽²⁾	51	(73.7; 94.3)	(146; 543)	19	(12.6; 56.6)	(5.7; 41.9)
	Year 10 ⁽³⁾	454	71.4%	166	5 0	21.2%	10.9
	(Pre-booster)	154	(63.6; 78.4)	(107; 258)	52	(11.1; 34.7)	(6.1; 19.3)
	(Post-	455	100%	23431	50	98.1%	5793
	booster)(3,4)	155	(97.6; 100)	(17351; 31641)	52	(89.7; 100)	(3586; 9357)
	(1)	340	100%	10315	444	100%	4574
	Month 1 ⁽¹⁾		(98.9; 100)	(9317; 11420)	114	(96.8; 100)	(3864; 5414)
	M 4(2)	40	86.0%	443	47	47.1%	30.7
	Year 4 ⁽²⁾	43	(72.1; 94.7)	(230; 853)	17	(23.0; 72.2)	(9.0; 105)
	V = (2)	5 4	92.2%	770	40	63.2%	74.1
Y	Year 5 ⁽²⁾	51	(81.1; 97.8)	(439; 1351)	19	(38.4; 83.7)	(21.9; 250)
	Year 10 ⁽³⁾	454	86.4%	364	F0	61.5%	56.0
	(Pre-booster)	154	(79.9; 91.4)	(255; 519)	52	(47.0; 74.7)	(28.8; 109)
	(Post-	155	100%	8958	5 0	100%	5138
	booster)(3,4)	155	(97.6; 100)	(7602; 10558)	52	(93.2; 100)	(3528; 7482)

The analysis of immunogenicity was conducted on the ATP cohorts for 1 month and 5 years post vaccination and the booster ATP cohort.

- (1) Study MenACWY-TT-015
- (2) Study MenACWY-TT-020
- (3) Study MenACWY-TT-099
- (4) Blood sampling was performed 1 month after a booster dose at Year 10.

^{*}rSBA analysis performed at GSK laboratories for 1 month post primary vaccination samples and at PHE laboratories in UK for the subsequent sampling time points.

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In a descriptive study conducted in 194 adults aged 56 years and older (Study MenACWY-TT-085), **Nimenrix**TM was immunogenic, with a vaccine response rate \geq 63.4% and with \geq 97.4% of subjects with rSBA titres \geq 8 against all four meningococcal groups. Moreover, at least 93.2% of subjects achieved the more conservative threshold of protection of rSBA titres \geq 128.

Booster response for subjects previously vaccinated with a conjugate meningococcal vaccine against *Neisseria meningitidis*

Nimenrix[™] booster vaccination in subjects previously primed with a monovalent (MenC-CRM) or a quadrivalent conjugate meningococcal vaccine (MenACWY-TT) was studied in subjects from 12 months of age onwards who received a booster vaccination. Robust anamnestic responses to the antigen(s) in the priming vaccine were observed (see Tables 8, 9, 14, 16, and 18).

Response to NimenrixTM in subjects previously vaccinated with a plain polysaccharide meningococcal vaccine against *Neisseria meningitidis*

In Study MenACWY-TT-021 conducted in subjects aged 4.5-34 years, the immunogenicity of **Nimenrix**[™] administered between 30 and 42 months after vaccination with a ACWY-PS vaccine was compared to the immunogenicity of **Nimenrix**[™] administered to age-matched subjects who had not been vaccinated with any meningococcal vaccine in the preceding 10 years. The rSBA GMTs were significantly lower in the subjects who had received a dose of ACWY-PS vaccine 30-42 months prior to **Nimenrix**[™]. The clinical relevance of this observation is unknown since all subjects achieved rSBA titres ≥8 for all four meningococcal groups. Results are shown in Table 19.

Table 19: rSBA* titres 1 month after Nimenrix[™] vaccination in subjects according to their meningococcal vaccine history (Study MenACWY-TT-021)

Mening o-coccal group	-	cts vaccinated 30 previously with AC		Subjects who had not received a meningococcal vaccine in the preceding			
	N	≥8 (95% CI)	GMT (95% CI)	N	≥8 (95% CI)	GMT (95% CI)	
Α	146	100% (97.5; 100)	6869 (6045; 7805)	69	100% (94.8; 100)	13015 (10722; 15798)	
С	169	100%	1946	75	100%	5495	

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Mening o-coccal		cts vaccinated 30 to		Subjects who had not received a meningococcal vaccine in the preceding			
group	N	≥8 GMT		≥8	GMT		
	IN .	(95% CI)	(95% CI)	N	(95% CI)	(95% CI)	
		(97.8; 100)	(1583; 2391)		(95.2; 100)	(4266; 7076)	
		100% 4636		100%	9078		
W-135	169	(97.8; 100)	(3942; 5451)	75	(95.2; 100)	(7088; 11627)	
		100%	7800		100%	13895	
Y	169	(97.8; 100)	(6683; 9104)	75	(95.2; 100)	(11186; 17261)	

The analysis of immunogenicity was conducted on the ATP cohort.

Response to Nimenrix[™] in subjects at increased risk for meningococcal infections

Study MenACWY-TT-084 evaluated the immunogenicity of one and two doses of **Nimenrix**TM given 2 months apart in 43 at-risk subjects aged 2-17 years (at increased risk for meningococcal disease, i.e., asplenic subjects, and hyposplenic subjects) compared to 43 healthy age-matched subjects.

One month after the first vaccine dose, vaccine response rates (rSBA titre \geq 1:32 or a \geq 4-fold increase in rSBA titre from baseline) for groups A, C, W-135, and Y, respectively, were 100%, 92.5%, 100% and 97.5% in the at-risk group and were 97.5%, 97.5%, 97.5%, and 100% for healthy subjects. After the second vaccine dose, vaccine response rates in both at-risk and healthy subjects were 100% for each of the four meningococcal groups.

Impact of a single dose of Nimenrix[™]

The Netherlands introduced **Nimenrix**[™] into the national immunization program in 2018 as a single dose at 14 months of age. A catch-up campaign for individuals 14-18 years of age initiated in 2018 and in 2020 a single dose of **Nimenrix**[™] at 14 years of age became routine, resulting in a toddler and adolescent national immunization program. Within two years, the incidence of meningococcal disease caused by groups C, W, and Y was significantly reduced by 100% (95% CI: 14, 100) in individuals 14-18 years of age, 85% (95% CI: 32, 97) in all vaccine eligible ages (direct effect), and 50% (95% CI: 28, 65) in non-vaccine eligible ages (indirect effect).

^{*} rSBA analysis performed at GSK laboratories

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5.2 Pharmacokinetic properties

Not applicable.

5.3 Pre-clinical safety data

Non-clinical data reveal no special hazard for humans based on local tolerance, acute toxicity, repeated dose toxicity, developmental/reproductive toxicity and fertility studies.

6. PHARMACEUTICAL PARTICULARS

6.1 List of excipients

Powder: sucrose, trometamol.

Solvent: sodium chloride, water for injections.

6.2 Incompatibilities

In the absence of compatibility studies, this medicinal product must not be mixed with other medicinal products.

6.3 Shelf life

The expiry date is indicated on the label and packaging.

6.4 Special precautions for storage

- Store in a refrigerator (2°C − 8°C)
- The solvent may also be stored at ambient temperature (25°C)
- Do not freeze
- Protect from light

6.5 Nature and contents of container

Powder in a vial containing 1 dose (0.5 ml) and 0.5 ml of solvent in a pre-filled syringe.
 Pack sizes of 1 and 10 with or without needles.

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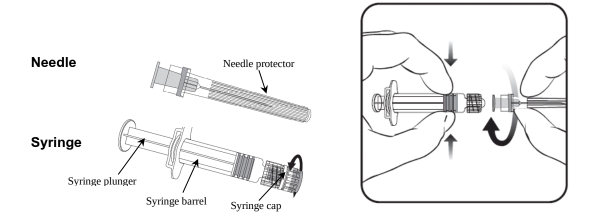
The powder is white. The solvent is clear and colorless.

6.6 Instructions for use and handling

Instructions for reconstitution of the vaccine with the solvent presented in pre-filled syringe

Nimenrix[™] must be reconstituted by adding the entire content of the pre-filled syringe of solvent to the vial containing the powder.

To attach the needle to the syringe, refer to the below picture. However, the syringe provided with **Nimenrix**[™] might be slightly different than the syringe described in the picture.



- 1. Holding the syringe <u>barrel</u> in one hand (avoid holding the syringe plunger), unscrew the syringe cap by twisting it anticlockwise.
- 2. To attach the needle to the syringe, twist the needle clockwise into the syringe until you feel it lock. (see picture).
- 3. Remove the needle protector, which on occasion can be a little stiff.

Add the solvent to the powder. After the addition of the solvent to the powder, the mixture should be well shaken until the powder is completely dissolved in the solvent.

The reconstituted vaccine is a clear colourless solution.

The reconstituted vaccine should be inspected visually for any foreign particulate matter and/or variation of physical aspect prior to administration. In the event of either being observed, discard the vaccine.

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After reconstitution, the vaccine should be used immediately.

A new needle should be used to administer the vaccine.

Any unused product or waste material should be disposed of in accordance with local requirements.

Not all presentations are available in every country.

Name and address of marketing authorization holder

Pfizer (Thailand) Limited

Bangkok, Thailand

Date of revision of package insert

09 February 2023

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