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

1. NAME OF THE MEDICINAL PRODUCT

*Nimenrix*TM powder and solvent for solution for injection in pre-filled syringe Meningococcal groups A, C, W-135 and Y conjugate vaccine

2. QUALITATIVE AND QUANTITATIVE COMPOSITION

After reconstitution, 1 dose (0.5 mL) contains:

Neisseria meningitidis group A polysaccharide ¹	5 micrograms
Neisseria meningitidis group C polysaccharide ¹	5 micrograms
Neisseria meningitidis group W-135 polysaccharide ¹	5 micrograms
Neisseria meningitidis group Y polysaccharide ¹	5 micrograms

¹conjugated to tetanus toxoid carrier protein

44 micrograms

3. PHARMACEUTICAL FORM

Powder and solvent for solution for injection.

The powder or cake is white.

The solvent is clear and colourless.

4. CLINICAL PARTICULARS

4.1 Therapeutic indications

*Nimenrix*TM is indicated for active immunisation of individuals from the age of 6 weeks against invasive meningococcal disease caused by *Neisseria meningitidis* groups A, C, W-135, and Y.

4.2 Posology and method of administration

Posology

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

Primary immunisation

Infants from 6 weeks to less than 6 months of age: two doses, each of 0.5 mL, should be administered with an interval of 2 months between doses.

Infants from 6 months of age, children, adolescents and adults: a single 0.5 mL dose should be administered.

An additional primary dose of $Nimenrix^{TM}$ may be considered appropriate for some individuals (see section 4.4).

Booster doses

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

After completion of the primary immunisation course in infants 6 weeks to less than 12 months of age, a booster dose should be given at 12 months of age with an interval of at least 2 months after the last *Nimenrix*TM vaccination (see section 5.1).

In previously vaccinated individuals 12 months of age and older, *Nimenrix*TM may be given as a booster dose if they have received primary vaccination with a conjugated or plain polysaccharide meningococcal vaccine (see sections 4.4 and 5.1).

Method of administration

Immunisation should be carried out by intramuscular injection only.

In infants, the recommended injection site is the anterolateral aspect of the thigh. In individuals from 1 year of age, the recommended injection site is the anterolateral aspect of the thigh or the deltoid muscle (see sections 4.4 and 4.5).

For instructions on reconstitution of the medicinal product before administration, see section 6.5.

4.3 Contraindications

Hypersensitivity to the active substances or to any of the excipients.

4.4 Special warnings and precautions for use

Traceability

In order to improve the traceability of biological medicinal products, the name and the batch number of the administered product should be clearly recorded.

*Nimenrix*TM 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.

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

Vaccination with *Nimenrix*TM 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 especially in adolescents as a psychogenic response to the needle injection. This can be accompanied by several neurological signs such as transient visual disturbance, paraesthesia and tonic-clonic limb movements during recovery. It is important that procedures are in place to avoid injury from faints.

Thrombocytopenia and coagulation disorders

*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.

Immunodeficiency

It may be expected that in patients receiving immunosuppressive treatment or patients with immunodeficiency, an adequate immune response may not be elicited.

Persons with familial complement deficiencies (for example, C5 or C3 deficiencies) and persons receiving treatments that inhibit 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*TM.

Protection against meningococcal disease

*Nimenrix*TM will only confer protection against *Neisseria meningitidis* groups A, C, W-135 and Y. The vaccine will not protect against any other *Neisseria meningitidis* groups.

A protective immune response may not be elicited in all vaccinees.

Effect of prior vaccination with plain polysaccharide meningococcal vaccine Subjects previously vaccinated with a plain polysaccharide meningococcal vaccine and vaccinated with *Nimenrix*TM 30 to 42 months later had lower Geometric Mean Titres (GMTs) measured with a serum bactericidal assay using rabbit complement (rSBA) than subjects who had not been vaccinated with any meningococcal vaccine in the preceding 10 years (see section 5.1). The clinical relevance of this observation is unknown.

Effect of pre-vaccination antibody to tetanus toxoid

The safety and immunogenicity of *Nimenrix*TM was evaluated when it was sequentially administered or co-administered with a vaccine containing, diphtheria and tetanus toxoids, acellular pertussis, inactivated polioviruses (1, 2 and 3), hepatitis B surface antigen and *Haemophilus influenzae* type b polyribosyl ribose phosphate conjugated to tetanus toxoid (DTaP-HBV-IPV/Hib) in the second year of life. The administration of *Nimenrix*TM one month after the DTaP-HBV-IPV/Hib vaccine resulted in lower rSBA GMTs against groups A, C and W-135 compared with co-administration (see section

4.5). The clinical relevance of this observation is unknown.

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 particular 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.

Immune responses in toddlers aged 12-14 months

Toddlers aged 12-14 months had similar rSBA titres to groups A, C, W-135 and Y at one month after one dose of *Nimenrix*TM or at one month after two doses of *Nimenrix*TM given two months apart.

A single dose was associated with lower hSBA titres to groups W-135 and Y compared with two doses given two months apart. Similar responses to groups A and C were observed after one or two doses (see section 5.1). The clinical relevance of this observation is unknown. If a toddler is expected to be at particular risk of invasive meningococcal disease due to exposure to groups W-135 and/or Y, consideration may be given to administering a second dose of *Nimenrix*TM after an interval of 2 months. Regarding waning of antibody against group A or group C after a first dose of *Nimenrix*TM in children aged 12-23 months, see under Persistence of serum bactericidal antibody titres.

Persistence of serum bactericidal antibody titres

Following administration of *Nimenrix*TM there is a waning of serum bactericidal antibody titres against group A when using 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 one year previously, consideration may be given to administering a booster dose.

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

Effect of *Nimenrix*TM on anti-tetanus antibody concentrations

Although an increase of the anti-tetanus toxoid (TT) antibody concentrations was observed following vaccination with *Nimenrix*TM, *Nimenrix*TM does not substitute for tetanus immunisation.

Giving *Nimenrix*TM with or one month before a TT-containing vaccine in the second year of life does not impair the response to TT or significantly affect safety. No data are available beyond the age of 2 years.

Sodium content

This vaccine contains less than 1 mmol sodium (23 mg) per dose, that is to say essentially 'sodium-free'.

4.5 Interaction with other medicinal products and other forms of interaction

In infants, *Nimenrix*TM can be given concomitantly with combined DTaP-HBV-IPV/Hib vaccines and with 10-valent pneumococcal conjugate vaccine.

From age 1 year and above, *Nimenrix*TM 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.

In the second year of life, *Nimenrix*TM can also be given concomitantly with combined diphtheria - tetanus - acellular pertussis (DTaP) vaccines, including combination DTaP vaccines with hepatitis B, inactivated poliovirus or *Haemophilus influenzae* type b (HBV, IPV or Hib), such as DTaP-HBV-IPV/Hib vaccine, and 13-valent pneumococcal conjugate vaccine.

In individuals aged 9 to 25 years, *Nimenrix*TM can be given concomitantly with human papillomavirus bivalent [Type 16 and 18] vaccine, recombinant (HPV2).

Whenever possible, *Nimenrix*TM and a TT containing vaccine, such as DTaP-HBV-IPV/Hib vaccine, should be co-administered or *Nimenrix*TM should be administered at least one month before the TT containing vaccine.

One month after co-administration with a 10-valent pneumococcal conjugate vaccine, 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 immune responses to the other nine pneumococcal serotypes.

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*TM or the tetanus or diphtheria antigens included in Tdap.

If *Nimenrix*TM is to be given at the same time as another injectable vaccine, the vaccines should always be administered at different injection sites.

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 $Nimenrix^{TM}$ in pregnant women.

Animal studies do not indicate direct or indirect harmful effects with respect to pregnancy, embryo/foetal development, parturition or post-natal development (see section 5.3).

*Nimenrix*TM should be used during pregnancy only when clearly needed, and the possible advantages outweigh the potential risks for the foetus.

Breast-feeding

It is unknown whether *Nimenrix*TM is excreted in human milk.

*Nimenrix*TM should only be used during breast-feeding when the possible advantages outweigh the potential risks.

Fertility

Animal studies do not indicate direct or indirect harmful effects with respect to fertility.

4.7 Effects on ability to drive and use machines

No studies on the effects of *Nimenrix*TM on the ability to drive and use machines have been performed. However, some of the effects mentioned under section 4.8 "Undesirable effects" may affect the ability to drive or use machines.

4.8 Undesirable effects

Summary of the safety profile

The safety of *Nimenrix*TM presented in the table below is based on two clinical study datasets as follows:

- A pooled analysis of data from 9,621 subjects administered a single dose of *Nimenrix*TM. This total included 3,079 toddlers (12 months to 23 months), 909 children between 2 and 5 years of age, 990 children between 6 and 10 years of age, 2,317 adolescents (11 to 17 years) and 2,326 adults (18 to 55 years).
- Data from a study in infants aged 6 to 12 weeks at the time of the first dose (Study MenACWY-TT-083), 1,052 subjects received at least one dose of a primary series of 2 or 3 doses of *Nimenrix*TM and 1,008 received a booster dose at approximately 12 months of age.

Safety data have also been evaluated in a separate study, in which a single dose of *Nimenrix*TM was administered to 274 individuals aged 56 years and older.

Local and general adverse reactions

In the 6-12 weeks and in the 12-14 months age groups who received 2 doses of *Nimenrix*TM given 2 months apart, the first and second doses were associated with similar local and systemic reactogenicity.

The local and general adverse reaction profile of a booster dose of *Nimenrix*TM given to subjects from 12 months through 30 years 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*TM, except for gastrointestinal symptoms (including diarrhoea, vomiting, and nausea), which were very common among subjects 6 years of age and older.

Tabulated list of adverse reactions

Adverse reactions reported are listed according to the following frequency categories:

Very common: $(\geq 1/10)$

Common: $(\ge 1/100 \text{ to } < 1/10)$ Uncommon: $(\ge 1/1,000 \text{ to } < 1/100)$ Rare: $(\ge 1/10,000 \text{ to } < 1/1,000)$

Very rare: (<1/10,000)

Not known (cannot be estimated from available data)

Table 1 shows the adverse reactions reported from the studies in subjects aged from 6 weeks up to 55 years of age and post-marketing experience. Adverse reactions reported in subjects aged >55 years were similar to those observed in younger adults.

Table 1: Tabulated summary of adverse reactions by system organ class

System Organ Class	Frequency	Adverse reactions
Blood and lymphatic system	Not known***	Lymphadenopathy
disorders		
Metabolism and nutrition	Very common	Appetite lost
disorders		
Psychiatric disorders	Very common	Irritability
	Uncommon	Insomnia
		Crying
Nervous system disorders	Very common	Drowsiness
		Headache
	Uncommon	Hypoaesthesia
		Dizziness
	Rare	Febrile convulsion
Gastrointestinal disorders	Common	Diarrhoea
		Vomiting
		Nausea*
Skin and subcutaneous tissue	Uncommon	Pruritus
disorders		Urticaria
		Rash**
Musculoskeletal and connective	Uncommon	Myalgia
tissue disorders		Pain in extremity
General disorders and	Very common	Fever
administration site conditions		Swelling at injection site
		Pain at injection site
		Redness at injection site

System Organ Class	Frequency	Adverse reactions		
		Fatigue		
	Common	Injection site haematoma*		
	Uncommon	Malaise		
		Injection site induration		
		Injection site pruritus		
		Injection site warmth		
		Injection site anaesthesia		
	Not known***	Extensive limb swelling at the injection		
		site, frequently associated with erythema,		
		sometimes involving the adjacent joint or		
		swelling of the entire injected limb		

^{*} Nausea and Injection site haematoma occurred at a frequency of Uncommon in infants

4.9 Overdose

No case of overdose has been reported.

5. PHARMACOLOGICAL PROPERTIES

5.1 Pharmacodynamic properties

Pharmacotherapeutic group: vaccines, meningococcal vaccines, ATC code: J07AH08

Mechanism of action

Anti-capsular meningococcal antibodies protect against meningococcal disease via complement mediated bactericidal activity. *Nimenrix*TM 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.

<u>Immunogenicity in infants</u>

In Study MenACWY-TT-083, the first dose was administered at 6 to 12 weeks of age, the second after an interval of 2 months, and a third (booster) dose administered at approximately 12 months of age. DTaP-HBV-IPV/Hib and a 10-valent pneumococcal vaccine were co-administered. *Nimenrix*TM elicited rSBA and hSBA titres against the four meningococcal groups as shown in Table 2. The response against group C was non-inferior to the one elicited by licensed MenC-CRM and MenC-TT vaccines in terms of percentages with rSBA titres ≥8 at 1 month after the second dose.

Data from this study support the extrapolation of the immunogenicity data and posology to infants from 12 weeks to less than 6 months of age.

Table 2: rSBA and hSBA titres following two doses of *Nimenrix*TM (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 12 months of age (Study MenACWY-TT-083)

^{**} Rash occurred at a frequency of Common in infants

^{***} ADR identified post-marketing

Meningo-	Manaina			rSBA*		hSBA**			
coccal	Vaccine	Time point	N	≥8	GMT	NI	≥8	GMT	
group	group		IN	(95% CI)	(95% CI)	N	(95% CI)	(95% CI)	
		Post dose 2 ⁽¹⁾	156	97.4%	203	202	96.5%	157	
	Nimenrix TM	Post dose 2(4)	456	(95.4; 98.6)	(182; 227)	202	(93.0; 98.6)	(131; 188)	
A	A Nimenrix TM	D = -4 14(1)	462	99.6%	1561	214	99.5%	1007	
	Post booster ⁽¹⁾	462	(98.4; 99.9)	(1412; 1725)	214	(97.4; 100)	(836; 1214)		
		Post dose 2 ⁽¹⁾	456	98.7%	612	218	98.6%	1308	
	Nim annivTM	Post dose 2(4)	430	(97.2; 99.5)	(540; 693)	218	(96.0; 99.7)	(1052; 1627)	
	Nimenrix TM	Post booster ⁽¹⁾	463	99.8%	1177	221	99.5%	4992	
		Post booster	403	(98.8; 100)	(1059; 1308)	221	(97.5; 100)	(4086; 6100)	
		Post dose 2 ⁽¹⁾	455	99.6%	958	202	100%	3188	
C	MenC-CRM	rost dose 200		(98.4; 99.9)	(850; 1079)	202	(98.2; 100)	(2646; 3841)	
	vaccine	Post booster ⁽¹⁾	446	98.4%	1051	216	100%	5438	
		rost booster	440	(96.8; 99.4)	(920; 1202)	210	(98.3; 100)	(4412; 6702)	
		Post dose 2 ⁽¹⁾	457	100%	1188	226	100%	2626	
	MenC-TT	rost dose 2.	437	(99.2; 100)	(1080; 1307)	220	(98.4; 100)	(2219; 3109)	
	vaccine	Post booster ⁽¹⁾	459	100%	1960	219	100%	5542	
		Tost booster	433	(99.2; 100)	(1776; 2163)	219	(98.3; 100)	(4765; 6446)	
		Post dose 2 ⁽¹⁾	455	99.1%	1605	217	100%	753	
W	Nimenrix TM	rost dose 2	433	(97.8; 99.8)	(1383; 1862)	217	(98.3; 100)	(644; 882)	
**	Timenta	Post booster ⁽¹⁾	462	99.8%	2777	218	100%	5123	
		Tost booster	402	(98.8; 100)	(2485; 3104)	210	(98.3; 100)	(4504; 5826)	
		Post dose 2 ⁽¹⁾	456	98.2%	483	214	97.7%	328	
Y	Nimenrix TM	1 ost dosc 2V	436	(96.6; 99.2)	(419; 558)	∠1 ⊤	(94.6; 99.2)	(276; 390)	
1	Tumentix	Post booster ⁽¹⁾	462	99.4%	881	217	100%	2954	
		1 OST DOOSTEL	702	(99.1; 99.9)	(787; 986)	217	(98.3; 100)	(2498; 3493)	

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

In Study MenACWY-TT-087, infants received either a single primary dose at 6 months followed by a booster dose at 15-18 months (DTaP-IPV/Hib and 10-valent pneumococcal conjugate vaccine was co-administered at both vaccination time points) or three primary doses at 2, 4, and 6 months followed by a booster dose at 15-18 months. 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 3.

Table 3: rSBA and hSBA titres following a single dose of *Nimenrix*TM in infants at 6 months of age and pre- and post-booster dose at 15-18 months of age (Study MenACWY-TT-087)

Meningo-			rSBA*		hSBA**				
coccal group	Time point	N	≥8 (95% CI)	GMT (95% CI)	N	≥8 (95% CI)	GMT (95% CI)		
	Post dose 1 ⁽¹⁾	163	98.8% (95.6; 99.9)	1333 (1035; 1716)	59	98.3% (90.9; 100)	271 (206; 355)		
A	Pre-booster	131	81.7% (74; 87.9)	125 (84.4; 186)	71	66.2% (54; 77)	20.8 (13.5; 32.2)		
	Post-booster ⁽¹⁾	139	99.3% (96.1; 100)	2762 (2310; 3303)	83	100% (95.7; 100)	1416 (1140; 1758)		
C	Post dose 1 ⁽¹⁾	163	99.4%	592	66	100%	523		

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

^{**} hSBA analysis performed at GSK laboratories

⁽¹⁾ blood sampling performed 21 to 48 days post vaccination

Meningo-			rSBA*			hSBA	**
coccal	Time point	N	≥8 (050/ CD)	GMT	N	≥8	GMT
group			(95% CI)	(95% CI)		(95% CI)	(95% CI)
			(96.6; 100)	(482; 726)		(94.6; 100)	(382; 717)
	Pre booster	131	65.6%	27.4	78	96.2%	151
	TTC 000StCI	131	(56.9; 73.7)	(20.6; 36.6)	70	(89.2; 99.2)	(109; 210)
	Post booster ⁽¹⁾	120	99.3%	2525	92	100%	13360
	rost booster	139	(96.1; 100)	(2102; 3033)	92	(96.1; 100)	(10953; 16296)
	D 4 1 1(1)	1.62	93.9%	1256	47	87.2%	137
	Post dose 1 ⁽¹⁾	163	(89; 97)	(917; 1720)	47	(74.3; 95.2)	(78.4; 238)
***	Pre booster	131	77.9%	63.3	52	100%	429
W			(69.8; 84.6)	(45.6; 87.9)	53	(93.3; 100)	(328; 559)
	D 41 4 (1)	120	100%	3145	50	100%	9016
	Post booster ⁽¹⁾	139	(97.4; 100)	(2637; 3750)	59	(93.9; 100)	(7045; 11537)
	D 4 1 1(1)	1.62	98.8%	1470	50	92.3%	195
	Post dose 1 ⁽¹⁾	163	(95.6; 99.9)	(1187; 1821)	52	(81.5; 97.9)	(118; 323)
37	D 1 /	121	88.5%	106	<i>C</i> 1	98.4%	389
Y	Pre booster	131	(81.8; 93.4)	(76.4; 148)	61	(91.2; 100)	(292; 518)
	D (1 (1)	120	100%	2749	60	100%	5978
	Post 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.

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. Results are shown in Table 3.

<u>Immunogenicity in toddlers aged 12-23 months</u>

In clinical studies MenACWY-TT-039 and MenACWY-TT-040, a single dose of *Nimenrix*TM 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 4.

Table 4: SBA* titres following a single dose of *Nimenrix*TM (or MenC-CRM) in toddlers aged 12-23 months (Studies MenACWY-TT-039/040)

Maninas				Study MenAC	CWY-T	T-039 ⁽¹⁾		Stı	Study MenACWY-TT-040 ⁽²⁾			
Meningo -coccal	Vaccine		rSBA	*		hSBA*	•	rSBA*				
group	group		≥8	GMT	N	≥8	GMT	N	≥8	GMT		
S. op	group	N	(95% CI)	(95% CI)	11	(95% CI)	(95% CI)	1	(95% CI)	(95% CI)		
A	Nimenrix TM	354	99.7%	2205	338	77.2%	19.0	183	98.4%	3170		
A	A Nimenrix ^{1 M}	334	(98.4; 100)	(2008; 2422)	336	(72.4; 81.6)	(16.4; 22.1)	103	(95.3; 99.7)	(2577; 3899)		
C	C Nimenrix TM	00.79/	99.7%	478	341	98.5%	196	183	97.3%	829		
C	Nimenrix	354	(98.4; 100)	(437; 522)	341	(96.6; 99.5)	(175; 219)		(93.7; 99.1)	(672; 1021)		

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

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

⁽¹⁾ blood sampling performed 1 month post vaccination

Maninga				Study MenAC	CWY-T	T-039 ⁽¹⁾		Stı	ıdy MenACWY	Y-TT-040 ⁽²⁾	
Meningo -coccal	Vaccine		rSBA	*		hSBA*	•	rSBA*			
group	group		≥8 (95% CI)	GMT (95% CI)	N	≥8 (95% CI)	GMT (95% CI)	N	≥8 (95% CI)	GMT (95% CI)	
	MenC- CRM vaccine	121	97.5% (92.9; 99.5)	212 (170; 265)	116	81.9% (73.7; 88.4)	40.3 (29.5; 55.1)	114	98.2% (93.8; 99.8)	691 (521; 918)	
W- 135	Nimenrix TM	354	100% (99.0; 100)	2682 (2453; 2932)	336	87.5% (83.5; 90.8)	48.9 (41.2; 58.0)	186	98.4% (95.4; 99.7)	4022 (3269; 4949)	
Y	Nimenrix TM	354	100% (99.0; 100)	2729 (2473; 3013)	329	79.3% (74.5; 83.6)	30.9 (25.8; 37.1)	185	97.3% (93.8; 99.1)	3168 (2522; 3979)	

The analysis of immunogenicity was conducted on the ATP cohorts.

Long-term immunogenicity in toddlers

Study MenACWY-TT-104 evaluated the immunogenicity after 1 month and the persistence of the response up to 5 years following 1 or 2 doses (given 2 months apart) of Nimenrix in toddlers aged 12 to 14 months. One month following one or two doses Nimenrix elicited rSBA titres against all four meningococcal groups that were similar in terms of the percentage of subjects with rSBA titre ≥ 8 and GMT. As a secondary endpoint hSBA titres were measured. One month post dose one or two Nimenrix elicited hSBA titres against groups W-135 and Y that were higher in terms of the percentage of subjects with hSBA titre ≥8 when two doses were given compared with one (see section 4.4). Nimenrix elicited hSBA titres against groups A and C that were similar in terms of the percentage of subjects with hSBA titre ≥8 when two doses were given compared with one. At Year 5 only a small difference in antibody persistence between one and two doses was observed, in terms of percentages of subjects with hSBA titres ≥8 against all groups. Antibody persistence was observed at Year 5 against groups C, W-135 and Y. After one and two doses the percentages of subjects with hSBA titres ≥8 for group C were 60.7% and 67.8%, group W-135 were 58.9% and 63.6% and group Y were 61.5% and 54.2%, respectively. For group A, 27.9% and 17.9% of subjects receiving one or two doses, respectively, had hSBA titres ≥8. Results are shown in Table 5.

⁽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

Table 5: rSBA and hSBA titres following one or two doses of $Nimenrix^{TM}$ with the first dose administered to toddlers aged 12-14 months and persistence up to 5

years (Study MenACWY-TT-104)

Mening-	rs (Study Me Nimenrix TM	Time		rSBA			hSBA**				
ococcal group	dose group	point ⁽¹⁾	N	≥8 (95% CI)	GMT (95% CI)	N	≥8 (95% CI)	GMT (95% CI)			
		Post dose 1	180	97.8% (94.4; 99.4)	1437 (1118; 1847)	74	95.9% (88.6, 99.2)	118 (86.8; 161)			
	1 dose	Year 1	167	63.5% (55.7; 70.8)	62.7 (42.6; 92.2)	70	35.1% (25.9; 49.5)	6.1 (4.1; 8.9)			
	1 dose	Year 3	147	46.9% (38.7; 55.3)	29.7 (19.8; 44.5)	55	36.4% (23.8; 50.4)	5.8 (3.8; 8.9)			
		Year 5	133	58.6% (49.8; 67.1)	46.8 (30.7; 71.5)	61	27.9% (17.1; 40.8)	4.4 (3.1; 6.2)			
A		Post dose 1	158	96.8% (92.8; 99.0)	1275 (970; 1675)	66	97.0% (89.5; 99.6)	133 (98.1; 180)			
		Post dose 2	150	98.0% (94.3; 99.6)	1176 (922; 1501)	66	97.0% (89.5; 99.6)	170 (126; 230)			
	2 doses	Year 1	143	70.6% (62.4; 77.9)	76.6 (50.7; 115.7)	62	35.5% (23.7; 48.7)	6.4 (4.2; 10.0)			
		Year 3	121	54.5% (45.2; 63.6)	28.5 (18.7; 43.6)	50	36.0% (22.9; 50.8)	5.4 (3.6; 8.0)			
		Year 5	117	65.8% (56.5; 74.3)	69.9 (44.7; 109.3)	56	17.9% (8.9; 30.4)	3.1 (2.4; 4.0)			
		Post dose 1	179	95.0% (90.7; 97.7)	452 (346; 592)	78	98.7% (93.1; 100)	152 (105; 220)			
	1 dose	Year 1	167	49.1% (41.3; 56.9)	16.2 (12.4; 21.1)	71	81.7% (70.7; 89.9)	35.2 (22.5; 55.2)			
	Tuose	Year 3	147	35.4% (27.7; 43.7)	9.8 (7.6; 12.7)	61	65.6% (52.3; 77.3)	23.6 (13.9; 40.2)			
C		Year 5	132	20.5% (13.9; 28.3)	6.6 (5.3; 8.2)	61	60.7% (47.3; 72.9)	18.1 (10.9; 30.0)			
		Post dose 1	157	95.5% (91.0; 98.2)	369 (281; 485)	70	95.7% (88.0; 99.1)	161 (110; 236)			
	2 doses	Post dose 2	150	98.7% (95.3; 99.8)	639 (522; 783)	69	100% (94.8; 100)	1753 (1278; 2404)			
	2 doses	Year 1	143	55.2% (46.7; 63.6)	21.2 (15.6; 28.9)	63	93.7% (84.5; 98.2)	73.4 (47.5; 113.4)			
		Year 3	121	33.9% (25.5; 43.0)	11.5 (8.4; 15.8)	56	67.9% (54.0; 79.7)	27.0 (15.6; 46.8)			

Mening-				rSBA			hSBA	
ococcal group	dose group	Time point ⁽¹⁾	N	≥8 (95% CI)	GMT (95% CI)	N	≥8 (95% CI)	GMT (95% CI)
		Year 5	116	28.4% (20.5; 37.6)	8.5 (6.4; 11.2)	59	67.8% (54.4; 79.4)	29.4 (16.3; 52.9)
		Post dose 1	180	95.0% (90.8; 97.7)	2120 (1601; 2808)	72	62.5% (50.3; 73.6)	27.5 (16.1; 46.8)
	1 dose	Year 1	167	65.3% (57.5; 72.5)	57.2 (39.9; 82.0)	72	95.8% (88.3; 99.1)	209.0 (149.9; 291.4)
	1 dose	Year 3	147	59.2% (50.8; 67.2)	42.5 (29.2; 61.8)	67	71.6% (59.3; 82.0)	30.5 (18.7; 49.6)
		Year 5	133	44.4% (35.8; 53.2)	25.0 (16.7; 37.6)	56	58.9% (45.0; 71.9)	20.8 (11.6; 37.1)
W-135		Post dose 1	158	94.9% (90.3; 97.8)	2030 (1511; 2728)	61	68.9% (55.7; 80.1)	26.2 (16.0; 43.0)
		Post dose 2	150	100% (97.6; 100)	3533 (2914; 4283)	70	97.1% (90.1; 99.7)	757 (550; 1041)
	2 doses	Year 1	143	77.6% (69.9; 84.2)	123 (82.7; 183)	65	98.5% (91.7; 100.0)	232.6 (168.3; 321.4)
		Year 3	121	72.7% (63.9; 80.4)	92.9 (59.9; 144)	54	87.0% (75.1; 94.6)	55.5 (35.3; 87.1)
		Year 5	117	50.4% (41.0; 59.8)	37.1 (23.3; 59.0)	44	63.6% (47.8; 77.6)	19.5 (10.7; 35.2)
		Post dose 1	180	92.8% (88.0; 96.1)	952 (705; 1285)	71	67.6% (55.5; 78.2)	41.2 (23.7; 71.5)
	1 dose	Year 1	167	73.1% (65.7; 79.6)	76.8 (54.2; 109.0)	62	91.9% (82.2; 97.3)	144 (97.2; 214.5)
	1 dose	Year 3	147	61.9% (53.5; 69.8)	58.0 (39.1; 86.0)	64	53.1% (40.2; 65.7)	17.3 (10.1; 29.6)
Y		Year 5	133	47.4% (38.7; 56.2)	36.5 (23.6; 56.2)	65	61.5% (48.6; 73.3)	24.3 (14.3; 41.1)
•		Post dose 1	157	93.6% (88.6; 96.9)	933 (692; 1258)	56	64.3% (50.4; 76.6)	31.9 (17.6; 57.9)
	2 doses	Post dose 2	150	99.3% (96.3; 100)	1134 (944; 1360)	64	95.3% (86.9; 99.0)	513 (339; 775)
	2 40363	Year 1	143	79.7% (72.2; 86.0)	112.3 (77.5; 162.8)	58	87.9% (76.7; 95.0)	143.9 (88.5; 233.8)
		Year 3	121	68.6% (59.5; 76.7)	75.1 (48.7; 115.9)	52	61.5% (47.0; 74.7)	24.1 (13.3; 43.8)

Mening-	Nimenrix TM	Time	rSBA*				hSBA**			
ococcal group	dose group	point ⁽¹⁾	N	≥8 (95% CI)	GMT (95% CI)	N	≥8 (95% CI)	GMT (95% CI)		
		Year 5	117	58.1% (48.6; 67.2)	55.8 (35.7; 87.5)	48	54.2% (39.2; 68.6)	16.8 (9.0; 31.3)		

The analysis of immunogenicity was conducted on the ATP cohort.

rSBA and hSBA titres were determined over a period of 10 years in children initially vaccinated with one dose of *Nimenrix*TM 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*TM administered 10 years following the initial vaccination with *Nimenrix*TM or MenC-CRM. Results are shown in Table 6 (see section 4.4).

Table 6: rSBA and hSBA titres following a single dose of *Nimenrix*TM (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 group	Time-point	N	≥8	GMT	N	≥8	GMT
group	group		11	(95% CI)	(95% CI)	1	(95% CI)	(95% CI)
		Month 1 ⁽¹⁾	222	100%	3707	217	91.2%	59.0
		Month 1	222	(98.4; 100)	(3327; 4129)	217	(86.7; 94.6)	(49.3; 70.6)
		Year 4 ⁽²⁾	45	64.4%	35.1	44	52.3%	8.8
		1 car 4 · /	43	(48.8; 78.1)	(19.4; 63.4)	44	(36.7; 67.5)	(5.4; 14.2)
A	Nimenrix TM	Year 5 ⁽²⁾	49	73.5%	37.4	45	35.6%	5.2
A	nimennix		49	(58.9; 85.1)	(22.1; 63.2)	43	(21.9: 51.2)	(3.4; 7.8)
		Year 10 ⁽³⁾	62	66.1%	28.9	59	25.4%	4.2
		(Pre-booster)	02	(53.0; 77.7)	(16.4; 51.0)	39	(15.0; 38.4)	(3.0; 5.9)
		(Post-	62	98.4%	5122	62	100%	1534
		booster)(3,4)	02	(91.3; 100)	(3726; 7043)	02	(94.2; 100)	(1112; 2117)
		Month 1 ⁽¹⁾	220	100%	879	221	99.1%	190
		Wionin 1	220	(98.3; 100)	(779; 991)	221	(96.8; 99.9)	(165; 219)
		Year 4 ⁽²⁾	45	97.8%	110	45	97.8%	370
			73	(88.2; 99.9)	(62.7; 192)	73	(88.2; 99.9)	(214; 640)
	Nimenrix TM		49	77.6%	48.9	48	91.7%	216
	vimenna		77	(63.4; 88.2)	(28.5; 84.0)		(80.0; 97.7)	(124; 379)
		Year 10 ⁽³⁾	62	82.3%	128	60	91.7%	349
		(Pre-booster)	02	(70.5; 90.8)	(71.1; 231)	00	(81.6; 97.2)	(197; 619)
		(Post-	62	100%	7164	59	100%	33960
C		booster)(3,4)	02	(94.2; 100)	(5478; 9368)	37	(93.9; 100)	(23890; 48274)
		Month 1 ⁽¹⁾	68	98.5%	415	68	72.1%	21.2
		Wionth 1	00	(92.1; 100)	(297; 580)	00	(59.9; 82.3)	(13.9; 32.3)
		Year 4 ⁽²⁾	10	80.0%	137	10	70.0%	91.9
	MenC-	1 cui ¬	10	(44.4; 97.5)	(22.6; 832)	10	(34.8; 93.3)	(9.8; 859)
	CRM vaccine	Year 5 ⁽²⁾	11	63.6%	26.5	11	90.9%	109
			11	(30.8; 89.1)	(6.5; 107)	11	(58.7; 99.8)	(21.2; 557)
		Year 10 ⁽³⁾	16	87.5%	86.7	15	93.3%	117
		(Pre-booster)	10	(61.7; 98.4)	(29.0; 259)	10	(68.1; 99.8)	(40.0; 344)
		(Post-	16	100%	5793	15	100%	42559
		booster)(3,4)	10	(79.4; 100)	(3631; 9242)	1.5	(78.2; 100)	(20106; 90086)

⁽¹⁾ blood sampling performed 21 to 48 days post vaccination

^{*}rSBA analysis performed at PHE laboratories

^{**}hSBA analysis performed at GSK laboratories

Meningo-	Vassins			rSBA	*		hSBA	**
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)	5395 (4870; 5976)	177	79.7% (73.0; 85.3)	38.8 (29.7; 50.6)
		Year 4 ⁽²⁾	45	60.0% (44.3; 74.3)	50.8 (24.0; 108)	45	84.4% (70.5; 93.5)	76.9 (44.0; 134)
W-135	Nimenrix TM	Year 5 ⁽²⁾	49	34.7% (21.7; 49.6)	18.2 (9.3; 35.3)	46	82.6% (68.6; 92.2)	59.7 (35.1; 101)
		Year 10 ⁽³⁾ (Pre-booster)	62	30.6% (19.6; 43.7)	15.8 (9.1; 27.6)	52	44.2% (30.5; 58.7)	7.7 (4.9; 12.2)
		(Post-booster) ^(3,4)	62	100% (94.2; 100)	25911 (19120; 35115)	62	100% (94.2; 100)	11925 (8716; 16316)
		Month 1 ⁽¹⁾	222	100% (98.4; 100)	2824 (2529; 3153)	201	66.7% (59.7; 73.1)	24.4 (18.6; 32.1)
		Year 4 ⁽²⁾	45	62.2% (46.5; 76.2)	44.9 (22.6; 89.3)	41	87.8% (73.8; 95.9)	74.6 (44.5; 125)
Y	Nimenrix TM	Year 5 ⁽²⁾	49	42.9% (28.8; 57.8)	20.6 (10.9; 39.2)	45	80.0% (65.4; 90.4)	70.6 (38.7; 129)
		Year 10 ⁽³⁾ (Pre-booster)	62	45.2% (32.5; 58.3)	27.4 (14.7; 51.0)	56	42.9% (29.7; 56.8)	9.1 (5.5; 15.1)
		(Post- booster) ^(3,4)	62	98.4% (91.3; 100)	7661 (5263; 11150)	61	100% (94.1; 100)	12154 (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. Subjects with a suboptimal response to meningococcal group C (defined as SBA titre below the pre-defined assay cut-off) were to receive an additional dose of MenC vaccine before Year 6. These subjects were excluded from the analysis at Years 4 and 5 but included in the analysis at Year 10.

- (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.

Persistence of booster response

Study MenACWY-TT-102 evaluated the persistence of SBA titres up to 6 years after a booster dose of Nimenrix 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 7 (see section 4.4).

Table 7: 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)

^{*} 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.

Meningo-	Vaccine			rSBA	*		hSBA	/ **
coccal	group	Time point	N	≥8 (050(CT)	GMT	N	≥8 (050(CI)	GMT
group	8 1			(95% CI) 99.7%	(95% CI) 2205		(95% CI) 77.2%	(95% CI) 19.0
		Month 1 ⁽¹⁾	354	(98.4; 100)	(2008; 2422)	338	(72.4; 81.6)	(16.4; 22.1)
		Year 4 ⁽²⁾ (Pre-Nimenrix booster)	212	74.5% (68.1; 80.2)	112 (80.3; 156)	187	28.9% (22.5; 35.9)	4.8 (3.9; 5.9)
A	Nimenrix	(Post-booster) ^(2,3)	214	100% (98.3; 100)	7173 (6389; 8054)	202	99.5% (97.3; 100)	1343 (1119; 1612)
		5 years after booster dose ⁽⁴⁾	137	89.8% (83.4; 94.3)	229 (163; 322)	135	53.3% (44.6; 62.0)	13.2 (9.6; 18.3)
		6 years after booster dose ⁽⁴⁾	134	92.5% (86.7; 96.4)	297 (214; 413)	130	58.5% (49.5; 67.0)	14.4 (10.5; 19.7)
		Month 1 ⁽¹⁾	354	99.7% (98.4; 100)	478 (437; 522)	341	98.5% (96.6; 99.5)	196 (175; 219)
	Nimenrix	Year 4 ⁽²⁾ (Pre-Nimenrix booster)	213	39.9% (33.3; 46.8)	12.1 (9.6; 15.2)	200	73.0% (66.3; 79.0)	31.2 (23.0; 42.2)
		(Post-booster) ^(2,3)	215	100% (98.3; 100)	4512 (3936; 5172)	209	100% (98.3; 100)	15831 (13626; 18394)
		5 years after booster dose ⁽⁴⁾	137	80.3% (72.6; 86.6)	66.0 (48.1; 90.5)	136	99.3% (96.0; 100)	337 (261; 435)
		6 years after booster dose ⁽⁴⁾	134	71.6% (63.2; 79.1)	39.6 (28.6; 54.6)	130	97.7% (93.4; 99.5)	259 (195; 345)
С		Month 1 ⁽¹⁾	121	97.5% (92.9; 99.5)	212 (170; 265)	116	81.9% (73.7; 88.4)	40.3 (29.5; 55.1)
	MenC- CRM vaccine	Year 4 ⁽²⁾ (Pre-MenC- CRM ₁₉₇ booster)	43	37.2% (23.0; 53.3)	14.3 (7.7; 26.5)	31	48.4% (30.2; 66.9)	11.9 (5.1; 27.6)
		(Post-booster) ^(2,3)	43	100% (91.8; 100)	3718 (2596; 5326)	33	100% (89.4; 100)	8646 (5887; 12699)
		5 years after booster dose ⁽⁴⁾	23	78.3% (56.3; 92.5)	47.3 (19.0; 118)	23	100% (85.2; 100)	241 (139; 420)
		6 years after booster dose ⁽⁴⁾	23	65.2% (42.7; 83.6)	33.0 (14.7; 74.2)	23	95.7% (78.1; 99.9)	169 (94.1; 305)
		Month 1 ⁽¹⁾	354	100% (99.0; 100)	2682 (2453; 2932)	336	87.5% (83.5; 90.8)	48.9 (41.2; 58.0)
		Year 4 ⁽²⁾ (Pre-Nimenrix booster)	213	48.8% (41.9; 55.7)	30.2 (21.9; 41.5)	158	81.6% (74.7; 87.3)	48.3 (36.5; 63.9)
W-135	Nimenrix	(Post-booster) ^(2,3)	215	100% (98.3; 100)	10950 (9531; 12579)	192	100% (98.1; 100)	14411 (12972; 16010)
		5 years after booster dose ⁽⁴⁾	137	88.3% (81.7; 93.2)	184 (130; 261)	136	100% (97.3; 100)	327 (276; 388)
		6 years after booster dose ⁽⁴⁾	134	85.8% (78.7; 91.2)	172 (118; 251)	133	98.5% (94.7; 99.8)	314 (255; 388)
		Month 1 ⁽¹⁾	354	100% (99.0; 100)	2729 (2473; 3013)	329	79.3% (74.5; 83.6)	30.9 (25.8; 37.1)
		Year 4 ⁽²⁾ (Pre-Nimenrix booster)	213	58.2% (51.3; 64.9)	37.3 (27.6; 50.4)	123	65.9% (56.8; 74.2)	30.2 (20.2; 45.0)
Y	Nimenrix	(Post-booster) ^(2,3)	215	100% (98.3; 100)	4585 (4129; 5093)	173	100% (97.9; 100)	6776 (5961; 7701)
		5 years after booster dose ⁽⁴⁾	137	92.7% (87.0; 96.4)	265 (191; 368)	137	97.8% (93.7; 99.5)	399 (321; 495)
		6 years after booster dose ⁽⁴⁾	134	94.0% (88.6; 97.4)	260 (189; 359)	131	97.7% (93.5; 99.5)	316 (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

Immunogenicity in children aged 2-10 years

In Study MenACWY-TT-081, a single dose of *Nimenrix*TM was demonstrated to be non-inferior to another licensed MenC-CRM vaccine 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]. The GMT was lower for the *Nimenrix*TM group [2795 (95% CI: 2393; 3263)] versus the MenC-CRM vaccine [5292 (95% CI: 3815; 7340)].

In Study MenACWY-TT-038, a single dose of *Nimenrix*TM 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 8.

Table 8: rSBA* titres following a single dose of *Nimenrix*TM (or ACWY-PS) in children aged 2-10 years (Study MenACWY-TT-038)

Meningo-		Nimenri	x ^{TM(1)}	ACWY-PS vaccine ⁽¹⁾				
coccal group	N	VR (95% CI)	GMT (95% CI)	N	VR (95% CI)	GMT (95% CI)		
A	594	89.1% (86.3; 91.5)	6343 (5998; 6708)	192	64.6% (57.4; 71.3)	2283 (2023; 2577)		
C	691	96.1% (94.4; 97.4)	4813 (4342; 5335)	234	89.7% (85.1; 93.3)	1317 (1043; 1663)		
W-135	691	97.4% (95.9; 98.4)	11543 (10873; 12255)	236	82.6% (77.2; 87.2)	2158 (1815; 2565)		
Y	723	92.7% (90.5; 94.5)	10825 (10233; 11452)	240	68.8% (62.5; 74.6)	2613 (2237; 3052)		

The analysis of immunogenicity was conducted on the ATP cohort.

- 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)

Persistence of SBA titres was evaluated in children initially vaccinated in Study MenACWY-TT-081 as shown in Table 9 (see section 4.4).

Table 9: rSBA and hSBA titres up to 44 months following Nimenrix (or MenC-CRM) in children aged 2-10 years at time of vaccination (Study MenACWY-TT-088)

Meningo-	Vaccine	Time-	rSBA*			hSBA**			
coccal group	group	point (months)	N	≥8 (95% CI)	GMT (95% CI)	N	≥8 (95% CI)	GMT (95% CI)	
A	Nimenrix 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)	
		44	189	85.7%	307	89	25.8%	4.8	

^{*} 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.

⁽¹⁾ Blood sampling performed 1 month post vaccination

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

^{*} rSBA analysis performed at GSK laboratories

Meningo-	Vaccina	Time-		rSBA	*		hSBA*	*
coccal group	Vaccine group	point (months)	N	≥8 (95% CI)	GMT (95% CI)	N	≥8 (95% CI)	GMT (95% CI)
				(79.9; 90.4)	(224; 423)		(17.1; 36.2)	(3.4; 6.7)
	A7. · TM	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 TM MenC-CRM vaccine	44	189	37.0% (30.1; 44.3)	14.5 (10.9; 19.2)	82	76.8% (66.2; 85.4)	36.4 (23.1; 57.2)
C		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)
		44	66	45.5% (33.1; 58.2)	31.0 (16.6; 58.0)	31	64.5% (45.4; 80.8)	38.8 (13.3; 113)
W-135	Nim aminTM	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 TM	44	189	68.3% (61.1; 74.8)	103 (72.5; 148)	87	80.5% (70.6; 88.2)	64.3 (42.7; 96.8)
V	NimannivTM	32	193	81.3% (75.1; 86.6)	227 (165; 314)	91	81.3% (71.8; 88.7)	79.2 (52.5; 119)
I	Nimenrix TM -	44	189	62.4% (55.1; 69.4)	78.9 (54.6; 114)	76	82.9% (72.5; 90.6)	127 (78.0; 206)

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

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 (Table 10) (see section 4.4).

Table 10: 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-			l month post-va			1 year persis			
_	Vaccine	(S	tudy MenACW	Y-TT-027)	(Study MenACWY-TT-028)				
coccal	group	NT	≥8	GMT	N.T	≥8	GMT		
group		N	(95% CI)	(95% CI)	N	(95% CI)	(95% CI)		
	Nimenrix TM	105	80.0%	53.4	104	16.3%	3.5		
	Nimenrix	103	(71.1; 87.2)	(37.3; 76.2)	104	(9.8; 24.9)	(2.7; 4.4)		
A	ACWY-PS	25	25.7%	4.1	25	5.7%	2.5		
vaccine	vaccine	35	(12.5;43.3)	(2.6; 6.5)	35	(0.7; 19.2)	(1.9; 3.3)		
	Nimenrix TM	101	89.1%	156	105	95.2%	129		
C	Nimenrix	101	(81.3; 94.4)	(99.3; 244)	105	(89.2; 98.4)	(95.4; 176)		
C	ACWY-PS	20	39.5%	13.1	21	32.3%	7.7		
	vaccine	38	(24.0; 56.6)	(5.4; 32.0)	31	(16.7; 51.4)	(3.5; 17.3)		
	Nimenrix TM	102	95.1%	133	102	100%	257		
W 125	Nimenrix	103	(89.0; 98.4)	(99.9; 178)	103	(96.5; 100)	(218; 302)		
W-135	ACWY-PS	25	34.3%	5.8	2.1	12.9%	3.4		
	vaccine	35	(19.1; 52.2)	(3.3; 9.9)	31	(3.6; 29.8)	(2.0; 5.8)		
	Nimenrix TM	89	83.1%	95.1	106	99.1%	265		
Y	Nimenrix	89	(73.7; 90.2)	(62.4; 145)	106	(94.9; 100)	(213; 330)		
Y	ACWY-PS	22	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

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

^{**} hSBA analysis performed at GSK laboratories

^{*} hSBA analysis performed at GSK laboratories

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 11 (see section 4.4).

Table 11: 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-	Vassins			rSBA	*		hSBA ³	**
coccal group	Vaccine group	Time point	N	≥8 (95% CI)	GMT (95% CI)	N	≥8 (95% CI)	GMT (95% CI)
		Month 1 ⁽¹⁾	225	100% (98.4; 100)	7301 (6586; 8093)	111(5)	81.1% (72.5; 87.9)	57.0 (40.3; 80.6)
		Year 5 ⁽²⁾	98	90.8% (83.3; 95.7)	141 (98.2; 203)	n/a ⁽⁶⁾		
	Nimenrix	Year 6 ⁽³⁾	98	79.6% (70.3; 87.1)	107 (66.0; 174)	90	41.1% (30.8; 52.0)	6.5 (4.8; 8.8)
		Year 10 ⁽³⁾ (Pre-booster)	73	89.0% (79.5; 95.1)	96.3 (57.1; 163)	62	33.9% (22.3; 47.0)	4.5 (3.3; 6.2)
		(Post-booster)(3,4)	74	95.9% (88.6; 99.2)	4626 (3041; 7039)	73	100% (95.1; 100)	1213 (994; 1481)
A		Month 1 ⁽¹⁾	75	100% (95.2; 100)	2033 (1667; 2480)	35(5)	25.7% (12.5; 43.3)	4.1 (2.6; 6.5)
	ACWY-	Year 5 ⁽²⁾	13	15.4% (1.9; 45.4)	4.7 (3.7; 6.0)	n/a ⁽⁶⁾		
	PS	Year 6 ⁽³⁾	24	12.5% (2.7; 32.4)	5.8 (3.5; 9.6)	21	33.3% (14.6; 57.0)	5.9 (3.0; 11.7)
	vaccine	Year 10 ⁽³⁾ (Pre-booster)	17	23.5% (6.8; 49.9)	8.0 (3.3; 19.3)	17	29.4% (10.3; 56.0)	6.2 (2.4; 15.7)
		(Post-booster) ^(3,4)	17	100% (80.5; 100)	6414 (3879; 10608)	17	100% (80.5; 100)	211 (131; 340)
		Month 1 ⁽¹⁾	225	100% (98.4; 100)	2435 (2106; 2816)	107(5)	89.7% (82.3; 94.8)	155 (101; 237)
		Year 5 ⁽²⁾	98	90.8% (83.3; 95.7)	79.7 (56.0; 113)	n/a ⁽⁶⁾		
	Nimenrix	Year 6 ⁽³⁾	98	82.7% (73.7; 89.6)	193 (121; 308)	97	93.8% (87.0; 97.7)	427 (261; 700)
		Year 10 ⁽³⁾ (Pre-booster)	74	85.1% (75.0; 92.3)	181 (106; 310)	73	91.8% (83.0; 96.9)	222 (129; 380)
		(Post-booster) ^(3,4)	74	100% (95.1; 100)	4020 (3319; 4869)	71	100% (94.9; 100)	15544 (11735; 20588)
С		Month 1 ⁽¹⁾	74	100% (95.1; 100)	750 (555; 1014)	38(5)	39.5% (24.0; 56.6)	13.1 (5.4; 32.0)
		Year 5 ⁽²⁾	13	100% (75.3; 100)	128 (56.4; 291)	n/a ⁽⁶⁾		
	ACWY- PS	Year 6 ⁽³⁾	24	79.2% (57.8; 92.9)	98.7 (42.2; 231)	24	100% (85.8; 100)	235 (122; 451)
	vaccine	Year 10 ⁽³⁾ (Pre-booster)	17	76.5% (50.1; 93.2)	96.2 (28.9; 320)	17	100% (80.5; 100)	99.1 (35.8; 274)
		(Post-booster) ^(3,4)	17	100% (80.5; 100)	15101 (7099; 32122)	17	94.1 (71.3; 99.9)	44794 (10112; 198440)

Meningo-	Vaccine			rSBA	\ *		hSBA ⁵	k **
coccal group	group	Time point	N	≥8 (95% CI)	GMT (95% CI)	N	≥8 (95% CI)	GMT (95% CI)
group		Month 1 ⁽¹⁾	225	100% (98.4; 100)	11777 (10666; 13004)	107 ⁽⁵⁾	95.3% (89.4; 98.5)	134 (101; 178)
		Year 5 ⁽²⁾	98	78.6% (69.1; 86.2)	209 (128; 340)	n/a ⁽⁶⁾		
Ni	Nimenrix	Year 6 ⁽³⁾	98	73.5% (63.6; 81.9)	265 (155; 454)	92	81.5% (72.1; 88.9)	62.5 (42.0; 93.1)
		Year 10 ⁽³⁾ (Pre-booster)	74	68.9% (57.1; 79.2)	206 (109; 392)	59	61.0% (47.4; 73.5)	17.5 (10.5; 29.2)
W 125		(Post-booster) ^(3,4)	74	100% (95.1; 100)	27944 (22214; 35153)	74	100% (95.1; 100)	6965 (5274; 9198)
W-135		Month 1 ⁽¹⁾	75	100% (95.2; 100)	2186 (1723; 2774)	35(5)	34.3% (19.1; 52.2)	5.8 (3.3, 9.9)
	ACWY-	Year 5 ⁽²⁾	13	0% (0.0; 24.7)	4.0 (4.0; 4.0)	n/a ⁽⁶⁾		
	PS vaccine	Year 6 ⁽³⁾	24	12.5% (2.7; 32.4)	7.6 (3.7; 15.6)	23	30.4% (13.2; 52.9)	7.0 (2.9; 16.9)
	vaccine	Year 10 ⁽³⁾ (Pre-booster)	17	23.5% (6.8; 49.9)	15.4 (4.2; 56.4)	15	26.7% (7.8; 55.1)	4.1 (2.0; 8.5)
		(Post-booster) ^(3,4)	17	94.1% (71.3; 99.9)	10463 (3254; 33646)	15	100% (78.2; 100)	200 (101; 395)
		Month 1 ⁽¹⁾	225	100% (98.4; 100)	6641 (6044; 7297)	94 ⁽⁵⁾	83.0% (73.8; 89.9)	93.7 (62.1; 141)
		Year 5 ⁽²⁾	98	78.6% (69.1; 86.2)	143 (88.0; 233)	n/a ⁽⁶⁾		
	Nimenrix	Year 6 ⁽³⁾	98	71.4% (61.4; 80.1)	136 (82.6; 225)	89	65.2% (54.3; 75.0)	40.3 (23.9; 68.1)
		Year 10 ⁽³⁾ (Pre-booster)	74	67.6% (55.7; 78.0)	98.5 (54.3; 179)	65	72.3% (59.8; 82.7)	35.7 (21.0; 60.6)
Y		(Post- booster) ^(3,4)	74	100% (95.1; 100)	7530 (5828; 9729)	74	100% (95.1; 100)	11127 (8909; 13898)
		Month 1 ⁽¹⁾	75	100% (95.2; 100)	1410 (1086; 1831)	32 ⁽⁵⁾	43.8% (26.4; 62.3)	12.5 (5.6; 27.7)
	ACWY-	Year 5 ⁽²⁾	13	7.7% (0.2; 36.0)	5.5 (2.7; 11.1)	n/a ⁽⁶⁾		
	PS vaccine	Year 6 ⁽³⁾	24	20.8% (7.1; 42.2)	11.6 (4.7; 28.7)	24	25.0% (9.8; 46.7)	7.3 (2.7; 19.8)
	, accine	Year 10 ⁽³⁾ (Pre-booster)	17	17.6% (3.8; 43.4)	10.2 (3.5; 30.2)	14	35.7% (12.8; 64.9)	7.8 (2.5; 24.4)
		(Post-booster) ^(3,4)	17	100% (80.5; 100)	6959 (3637; 13317)	17	100% (80.5; 100)	454 (215; 960)

The analysis of immunogenicity was conducted on the ATP cohort for each time point. Subjects with a suboptimal response to meningococcal group C (defined as SBA titre below the pre-defined assay cutoff) were to receive an additional dose of MenC vaccine before Year 6. These subjects were excluded from the analysis at Year 5 but included in the analyses at Years 6 and 10.

- (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.
- (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*TM or one dose of the ACWY-PS vaccine was administered.

*Nimenrix*TM was demonstrated to be immunologically non-inferior to the ACWY-PS vaccine in terms of vaccine response as shown in Table 12.

Table 12: rSBA* titres following a single dose of *Nimenrix*TM (or ACWY-PS) in adolescents aged 11-17 years and adults aged 18-55 years (Studies MenACWY-TT-035/036)

			Study MenAC	WY-TT-036	St	tudy MenACW	Y-TT-035		
Meningo- coccal	Vaccine	(11-17 years) ⁽¹⁾				(18-55 years) ⁽¹⁾			
group	group	N	VR (95% CI)	GMT (95% CI)	N	VR (95% CI)	GMT (95% CI)		
A	Nimenrix TM	553	85.4% (82.1; 88.2)	5928 (5557; 6324)	743	80.1% (77.0; 82.9)	3625 (3372; 3897)		
A	ACWY-PS vaccine	191	77.5% (70.9; 83.2)	2947 (2612; 3326)	252	69.8% (63.8; 75.4)	2127 (1909; 2370)		
C	Nimenrix TM	642	97.4% (95.8; 98.5)	13110 (11939; 14395)	849	91.5% (89.4; 93.3)	8866 (8011; 9812)		
C	ACWY-PS vaccine	211	96.7% (93.3; 98.7)	8222 (6807; 9930)	288	92.0% (88.3; 94.9)	7371 (6297; 8628)		
W-135	Nimenrix TM	639	96.4% (94.6; 97.7)	8247 (7639; 8903)	860	90.2% (88.1; 92.1)	5136 (4699; 5614)		
W-133	ACWY-PS vaccine	216	87.5% (82.3; 91.6)	2633 (2299; 3014)	283	85.5% (80.9; 89.4)	2461 (2081; 2911)		
Y	Nimenrix TM	657	93.8% (91.6; 95.5)	14086 (13168; 15069)	862	87.0% (84.6; 89.2)	7711 (7100; 8374)		
1	ACWY-PS vaccine	219	78.5% (72.5; 83.8)	5066 (4463; 5751)	288	78.8% (73.6; 83.4)	4314 (3782; 4921)		

The analysis of immunogenicity was conducted on the ATP cohorts.

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

- rSBA titres \ge 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 13.

Table 13: 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-

⁽¹⁾ Blood sampling performed 1 month post vaccination

^{*} rSBA analysis performed at GSK laboratories

036/043/101)

Meningo-	Time-		Nimenrix	$\mathcal{C}^{\mathrm{TM}}$		ACWY-PS v	accine
coccal	point	N	≥8	GMT	N	≥8	GMT
group	point	11	(95% CI)	(95% CI)	11	(95% CI)	(95% CI)
	Month 1 ⁽¹⁾	674	100%	5929	224	99.6%	2947
	Month 1	0/4	(99.5; 100)	(5557; 6324)	224	(97.5; 100)	(2612; 3326)
	Year 3 ⁽²⁾	449	92.9%	448	150	82.7%	206
	Teal 3	449	(90.1; 95.1)	(381; 527)	130	(75.6; 88.4)	(147; 288)
	Year 5 ⁽²⁾	226	97.5%	644	96	93.0%	296
A	Year 3(-)	236	(94.5; 99.1)	(531; 781)	86	(85.4; 97.4)	(202; 433)
	Year 10 ⁽³⁾		85.2%	248		80.4%	143
	(Pre-	162			51	(66.9; 90.2)	
	booster)		(78.8; 90.3)	(181; 340)		(66.9; 90.2)	(80.5; 253)
	(Post-	1.62	100%	3760	5.1	100%	2956
	booster)(3,4)	162	(97.7; 100)	(3268; 4326)	51	(93.0; 100)	(2041; 4282)
	M 4 1(1)	(72	100%	13110	224	100%	8222
	Month 1 ⁽¹⁾	673	(99.5; 100)	(11939; 14395)	224	(98.4; 100)	(6808; 9930)
	Year 3 ⁽²⁾	449	91.1%	371	150	86.0%	390
	rear 3(-)	449	(88.1; 93.6)	(309; 446)	130	(79.4; 91.1)	(262; 580)
	Year 5 ⁽²⁾	226	88.6%	249	0.5	87.1%	366
C	Year 3(2)	236	(83.8; 92.3)	(194; 318)	85	(78.0; 93.4)	(224; 599)
	Year 10 ⁽³⁾		90.1%	244		92.40/	177
	(Pre-	162		244	51	82.4%	177
	booster)		(84.5; 94.2)	(182; 329)		(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
	Month 1	0/8	(99.2; 100)	(7639; 8903)	224	(98.4; 100)	(2299; 3014)
	Year 3 ⁽²⁾	449	82.0%	338		30.0%	16.0
	Teal 3	449	(78.1; 85.4)	(268; 426)	130	(22.8; 38.0)	(10.9; 23.6)
	Year 5 ⁽²⁾	236	86.0%	437 86		34.9%	19.7
W-135		230	(80.9; 90.2)	(324; 588)	80	(24.9; 45.9)	(11.8; 32.9)
	Year 10 ⁽³⁾		71.6%	146		43.1%	16.4
	(Pre-	162	(64.0; 78.4)	(97.6; 217)	51	(29.3; 57.8)	(9.2; 29.4)
	booster)			,			(9.2, 29.4)
	(Post-	162	100%	11243	51	100%	3674
	booster)(3,4)	102	(97.7; 100)	(9367; 13496)	31	(93.0; 100)	(2354; 5734)
	Month 1 ⁽¹⁾	677	100%	14087	224	100%	5066
	IVIOIIIII I · /	0//	(99.5; 100)	(13168; 15069)	22 4	(98.4; 100)	(4463; 5751)
	Year 3 ⁽²⁾	449	93.1%	740	150	58.0%	69.6
	Tear 5	777	(90.3; 95.3)	(620; 884)	130	(49.7; 66.0)	(44.6; 109)
	Year 5 ⁽²⁾	236	96.6%	1000	86	66.3%	125
Y		230	(93.4; 98.5)	(824; 1214)	00	(55.3; 76.1)	(71.2; 219)
	Year 10 ⁽³⁾		90.7%	447		49.0%	32.9
	(Pre-	162	(85.2; 94.7)	(333; 599)	51	(34.8; 63.4)	(17.1; 63.3)
	booster)						
	(Post-	162	100%	7585	51	98.0%	3296
The on	booster)(3,4)	102	(97.7; 100)	(6748; 8525)	<i>J</i> 1	(89.6; 100)	(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.

hSBA persistence was evaluated up to 5 years after vaccination in adolescents and adults initially vaccinated in Study MenACWY-TT-052 as shown in Table 14 (see

^{*} 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.

section 4.4).

Table 14: hSBA* titres following a single dose of *Nimenrix*TM in adolescents and adults aged 11-25 years and persistence up to 5 years following vaccination (Studies MenACWY-TT-052/059)

(Studies Menac	W 1-11-052/03	,,,		
Meningococcal 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)
A	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)
	Month 1 ⁽¹⁾	359	96.1% (93.5; 97.9)	532 (424; 668)
\mathbf{C}	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)
	Month 1 ⁽¹⁾	334	91.0% (87.4; 93.9)	117 (96.8; 141)
W-135	Year 1 ⁽²⁾	327	98.5% (96.5; 99.5)	197 (173; 225)
	Year 5 ⁽²⁾	138	87.0% (80.2; 92.1)	103 (76.3; 140)
	Month 1 ⁽¹⁾	364	95.1% (92.3; 97.0)	246 (208; 291)
\mathbf{Y}	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)

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

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 15.

Table 15: 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)

Maninga			Nimeni	rix	ACWY-PS vaccine			
Meningo- coccal group	Time point	N	≥8 (95% CI)	GMT (95% CI)	N	≥8 (95% CI)	GMT (95% CI)	
	Month 1 ⁽¹⁾	323	100% (98.9; 100)	4945 (4452, 5493)	112	100% (96.8, 100)	2190 (1858, 2582)	
	Year 4 ⁽²⁾	43	95.3% (84.2; 99.4)	365 (226; 590)	17	76.5% (50.1; 93.2)	104 (31.0; 351)	
A	Year 5 ⁽²⁾	51	84.3% (71.4; 93.0)	190 (108; 335)	19	57.9% (33.5; 79.7)	37.0 (12.6; 109)	
	Year 10 ⁽³⁾ (Pre-booster)	155	78.1% (70.7; 84.3)	154 (108; 219)	52	71.2% (56.9; 82.9)	75.1 (41.4; 136)	
	(Post-booster) ^(3,4)	155	100% (97.6; 100)	4060 (3384; 4870)	52	100% (93.2; 100)	3585 (2751; 4672)	
	Month 1 ⁽¹⁾	341	99.7% (98.4; 100)	10074 (8700, 11665)	114	100% (96.8; 100)	6546 (5048; 8488)	
С	Year 4 ⁽²⁾	43	76.7% (61.4; 88.2)	126 (61.6; 258)	17	41.2% (18.4; 67.1)	16.7 (5.7; 48.7)	
	Year 5 ⁽²⁾	51	72.5% (58.3; 84.1)	78.5 (41.8; 147)	18	38.9% (17.3; 64.3)	17.3 (6.0; 49.7)	

^{*} hSBA analysis performed at GSK laboratories

Meningo- coccal group	Time point	Nimenrix			ACWY-PS vaccine		
		N	≥8 (95% CI)	GMT	N	≥8 (95% CI)	GMT
	Year 10 ⁽³⁾ (Pre-booster)	154	90.9% (85.2; 94.9)	(95% CI) 193 (141; 264)	52	88.5% (76.6; 95.6)	(95% CI) 212 (110; 412)
	(Post-booster) ^(3,4)	155	100% (97.6; 100)	13824 (10840; 17629)	52	98.1% (89.7; 100)	3444 (1999; 5936)
W-135	Month 1 ⁽¹⁾	340	99.7% (98.4; 100)	8577 (7615; 9660)	114	100% (96.8; 100)	2970 (2439; 3615)
	Year 4 ⁽²⁾	43	90.7% (77.9; 97.4)	240 (128; 450)	17	17.6% (3.8; 43.4)	8.3 (3.6; 19.5)
	Year 5 ⁽²⁾	51	86.3% (73.7; 94.3)	282 (146; 543)	19	31.6% (12.6; 56.6)	15.4 (5.7; 41.9)
	Year 10 ⁽³⁾ (Pre-booster)	154	71.4% (63.6; 78.4)	166 (107; 258)	52	21.2% (11.1; 34.7)	10.9 (6.1; 19.3)
	(Post-booster) ^(3,4)	155	100% (97.6; 100)	23431 (17351; 31641)	52	98.1% (89.7; 100)	5793 (3586; 9357)
Y	Month 1 ⁽¹⁾	340	100% (98.9; 100)	10315 (9317; 11420)	114	100% (96.8; 100)	4574 (3864; 5414)
	Year 4 ⁽²⁾	43	86.0% (72.1; 94.7)	443 (230; 853)	17	47.1% (23.0; 72.2)	30.7 (9.0; 105)
	Year 5 ⁽²⁾	51	92.2% (81.1; 97.8)	770 (439; 1351)	19	63.2% (38.4; 83.7)	74.1 (21.9; 250)
	Year 10 ⁽³⁾ (Pre-booster)	154	86.4% (79.9; 91.4)	364 (255; 519)	52	61.5% (47.0; 74.7)	56.0 (28.8; 109)
	(Post-booster) ^(3,4)	155	100% (97.6; 100)	8958 (7602; 10558)	52	100% (93.2; 100)	5138 (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.

In a separate study (MenACWY-TT-085), a single dose of *Nimenrix*TM was administered to 194 Lebanese adults aged 56 years and older (including 133 aged 56-65 years and 61 aged >65 years). The percentage of subjects with rSBA titres (measured at GSK's laboratories) \geq 128 before vaccination ranged from 45% (group C) to 62% (group Y). Overall, at 1 month post-vaccination the percentage of vaccines with rSBA titres \geq 128 ranged from 93% (group C) to 97% (group Y). In the subgroup aged >65 years the percentage of vaccines with rSBA titres \geq 128 at 1 month post-vaccination ranged from 90% (group A) to 97% (group Y).

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

NimenrixTM 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 6, 7, 11, 13, and 15).

^{*} 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.

Response to *Nimenrix*TM in subjects previously vaccinated with a plain polysaccharide vaccine against *Neisseria meningitidis*

In Study MenACWY-TT-021 conducted in subjects aged 4.5-34 years, the immunogenicity of *Nimenrix*TM administered between 30 and 42 months after vaccination with a ACWY-PS vaccine was compared to the immunogenicity of *Nimenrix*TM administered to age-matched subjects who had not been vaccinated with any meningococcal vaccine in the preceding 10 years. An immune response (rSBA titre ≥8) was observed against all four meningococcal groups in all subjects regardless of the meningococcal vaccine history. The rSBA GMTs were significantly lower in the subjects who had received a dose of ACWY-PS vaccine 30-42 months prior to *Nimenrix*TM, however 100% of subjects achieved rSBA titres ≥8 for all four meningococcal groups (A, C, W-135, Y) (see section 4.4).

Children (2-17 years) with anatomical or functional asplenia

Study MenACWY-TT-084 compared immune responses to two doses of *Nimenrix*TM given 2 months apart between 43 subjects aged 2-17 years with anatomic or functional asplenia subjects and 43 age-matched subjects with normal splenic function. One month after the first vaccine dose and 1 month after the second dose similar percentages of subjects in the two groups had rSBA titres ≥ 8 and ≥ 128 and hSBA titres ≥ 4 and ≥ 8 .

Impact of a single dose of *Nimenrix*TM

In 2018, the Netherlands added *Nimenrix*TM to the national immunisation programme as a single dose for toddlers at 14 months of age to replace the meningococcal C conjugate vaccine. A catch-up campaign with a single dose of *Nimenrix*TM for adolescents 14-18 years of age also initiated in 2018, and it became routine in 2020 leading to a toddler and adolescent national immunisation programme. 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). The impact of *Nimenrix*TM was primarily driven by a reduction in group W disease.

5.2 Pharmacokinetic properties

Not applicable.

5.3 Preclinical 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 Incompatibilities

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

other medicinal products.

6.2 Shelf life

Please refer to the outer carton for the expiry date.

After reconstitution

After reconstitution, the vaccine should be used promptly. Although delay is not recommended, stability has been demonstrated for 8 hours at 30°C after reconstitution. If not used within 8 hours, do not administer the vaccine.

6.3 Special precautions for storage

Store in a refrigerator $(2^{\circ}C - 8^{\circ}C)$.

Do not freeze.

Store in the original package in order to protect from light.

For storage conditions after reconstitution of the medicinal product, see section 6.2.

6.4 Nature and contents of container

Powder in a vial (type I glass) with a stopper (butyl rubber) and solvent in a pre-filled syringe with a stopper (butyl rubber).

Pack sizes of 1 with needles.

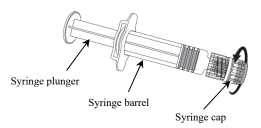
6.5 Special precautions for disposal and other handling

<u>Instructions</u> for reconstitution of the vaccine with the solvent presented in pre-filled syringe

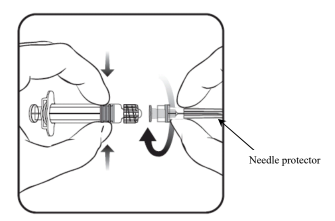
*Nimenrix*TM must be reconstituted by adding the entire content of the pre-filled syringe of solvent to the vial containing the powder.

To attach the needle to the syringe, refer to the below picture. However, the syringe provided with *Nimenrix*TM might be slightly different (without screw thread) than the syringe described in the picture. In that case, the needle should be attached without screwing.

1. Holding the syringe **barrel** 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.



4. 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.

After reconstitution, the vaccine should be used promptly.

A new needle should be used to administer the vaccine.

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

Pfizer Corporation Hong Kong Limited NOV 2022

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