This medicine protocol is a specific written instruction for the administration of Infanrix Hexa (6 in 1 vaccine), to Refugees and Applicants Seeking Protection in Ireland (hereafter referred to as vaccine recipients) and in the event of an outbreak as advised by the public health by registered nurses and registered midwives. This medicine protocol is valid for the 2023/2024 Health Service Executive (HSE) Primary Childhood Immunisation Programme (PCIP), Schools Immunisation Programme (SIP), National Immunisation Advisory Committee (NIAC) catch up Immunisation Programme and in the event of an outbreak as advised by public health.

This medicine protocol enables registered nurses and registered midwives employed in the voluntary and statutory services of the HSE including Mass vaccination clinics who have undertaken the required education and training programmes to administer Infanrix Hexa (6 in 1 vaccine), with reference to and guidance from Nursing & Midwifery Board of Ireland (NMBI), National Immunisation Working Group, National Immunisation Advisory Committee (NIAC), National Immunisation Office (NIO), HSE and in accordance with the Summary of Product Characteristics (SmPC) for Infanrix Hexa (6 in 1 vaccine) as detailed by the European Medicines Agency (EMA) at <u>www.ema.ie</u>

- An Bord Altranais (2007) Guidance to Nurses and Midwives on Medication Management. Dublin: An Bord Altranais
- National Immunisation Advisory Committee (2023) Anaphylaxis: Immediate Management in the Community available at:<u>https://rcpi.access.preservica.com/uncategorized/IO\_a36f9e4b-4c80-432d-8264-</u> 546089359925/
- National Immunisation Advisory Committee Immunisation Guidelines for Ireland: Royal College of Physicians of Ireland National Immunisation Advisory Committee available at: <u>https://www.rcpi.ie/Healthcare-</u> <u>Leadership/NIAC/Immunisation-Guidelines-for-Ireland</u>
- National Immunisation Office (2023/2024) Supporting Information for Staff: Schools Immunisation Programme available at:

https://www.hse.ie/eng/health/immunisation/hcpinfo/schoolproghcp/supportingdoc.pdf

- National Immunisation Office (2023) Clinical information to support healthcare staff to deliver catch-up vaccination for Refugees and Applicants Seeking Protection in Ireland and in the event of an outbreak available at: <a href="https://www.hse.ie/eng/health/immunisation/hcpinfo/botpipa/clinicalinfobotpipa.pdf">https://www.hse.ie/eng/health/immunisation/hcpinfo/botpipa/clinicalinfobotpipa.pdf</a>
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- Nursing and Midwifery Board of Ireland (2021) Code of Professional Conduct and Ethics for Registered Nurses and Registered Midwives. Dublin: Nursing and Midwifery Board of Ireland available at: <u>https://www.nmbi.ie/NMBI/media/NMBI/Code-of-Professional-Conduct-and-Ethics.pdf</u>
- Nursing and Midwifery Board of Ireland (2020) Guidance for Registered Nurses and Midwives on Medication Administration available at:

https://www.nmbi.ie/NMBI/media/NMBI/NMBI-Medication-Administration-2020.pdf?ext=.pdf

- Nursing and Midwifery Board of Ireland (2022) *Practice Standards for Midwives*. Dublin: Nursing and Midwifery Board of Ireland available at: <u>https://www.nmbi.ie/Standards-Guidance/MidwivesStandards</u>
- Nursing and Midwifery Board of Ireland (2015) Recording Clinical Practice. Guidance to Nurses and Midwives. Dublin: Nursing and Midwifery Board of Ireland available at: <u>https://www.nmbi.ie/Standards-Guidance/More-Standards-Guidance/More-Standards-Guidance/Recording-Clinical-Practice</u>
- Nursing and Midwifery Board of Ireland (2015) Scope of Nursing and Midwifery Practice Framework. Dublin: Nursing and Midwifery Board of Ireland available at: <u>https://www.nmbi.ie/StandardsGuidance/Scope-of-Practice/Nursing-Practice-Scope-Definition</u>

The Nursing and Midwifery Board of Ireland defines medicine protocols as "written directions that allow for the supply and administration of a named medicinal product by a nurse or midwife in identified clinical situations. A medicine protocol involves the authorisation of the nurse/midwife to supply and administer a medicine to groups of patients in a defined situation meeting specific criteria and who may not be individually identified before presentation for treatment. An individually named prescription is not required for the supply and administration of medication when a medicine protocol is in effect" (An Bord Altranais, 2007).



Document reference			
number	NIO 2023		
1.0 Critical Elements			
Name of Organisation	Health Service Providers/Mass vaccination clinics across the voluntary and statutory		
where medicine protocol			
applies	services of the fise. This inequality Protocol applies to:		
	<ul> <li>registered nurses and registered midwives employed in the voluntary and statutory</li> </ul>		
	services of the HSE including Mass vaccination clinics, congregated settings, temporary		
	clinics and mobile units		
Date the medicine	December 2023		
protocol comes into effect			
Date for review of	December 2024		
medicine protocol			
Document prepared by	HSE National Immunisation Office (NIO) in collaboration with the Office of the Nursing		
	and Midwifery Services Director (ONMSD) HSE.		
Names and Signatures of	Name: Dr. Eamonn O 'Moore Director of National Health Protection		
the employing authority			
who is authorising the			
implementation of the			
medicine protocol			
"On behalf of the			
authority employing			
professionals authorised to administer under this	5-5111		
medicine protocol, I have	DE 50'Moone		
read this medicine			
protocol and authorise its	Signature:		
implementation"			
	Name: <b>Dr Colm Henry</b> , Chief Clinical Officer, HSE		
	wenning		
	Signature:		
	Name: Dr Geraldine Shaw, Nursing and Midwifery Services Director, HSE		
	<u>An</u>		
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	1		

	Signature:	
2.0 Clinical Criteria		
Clinical condition for use of the medicine protocol	The clinical condition for which this medicine protocol has been developed is for the active immunisation against and prevention of Diphtheria, Tetanus, Pertussis, Hepatitis B, Poliomyelitis and Haemophilus type b disease.	
Circumstances in which the medicine protocol applies	To provide an Infanrix Hexa (6 in 1 vaccine) to Refugees and Applicants Seeking Protection in Ireland and in the event of an outbreak (hereafter referred to as vaccine recipients).	
Inclusion criteria for vaccine recipients receiving Infanrix Hexa (6 in 1 vaccine)	<b>For primary immunisation</b> : The primary course consists of 3 doses given at 2, 4 and 6 months.	
Infanrix Hexa (6 in 1 vaccine) under this medicine protocol	<b>For catch-up programme:</b> Children aged less than 10 years with no history of receipt of these antigens previously as per the NIAC catch up schedule.	
	For outbreak response: As advised by public health	
	Precautions	
	Acute severe febrile illness, defer until recovery.	
	• Type III (Arthus) hypersensitivity reaction to a previous dose (see adverse reactions in NIAC Chapter 6). Persons experiencing these reactions usually have very high serum diphtheria or tetanus antitoxin levels; they should not be given further routine or emergency booster doses of tetanus or diphtheria containing vaccines more frequently than every 10 years.	
	<b>Note:</b> COVID-19 vaccines (except for COVID-19 vaccine given to children aged 6 months	
	4 years where a 14-day interval is recommended) and other vaccines may be administered at the same time or at any interval.	
Exclusion criteria for vaccine recipients receiving Infanrix Hexa (6 in 1 vaccine) under this medicine protocol	<ul> <li>A known history of anaphylactic or hypersensitivity reaction to Infanrix Hexa (6 in 1 vaccine) or any of the vaccine's constituents.</li> </ul>	
Actions to be taken for those who are excluded	All recipients meeting exclusion criteria must be referred to the medical practitioner for an individual medical assessment.	
from the medicine protocol	<ul> <li>Document action in clinical notes</li> <li>Where Infanrix Hexa (6 in 1 vaccine) is prescribed following medical assessment, the nurse or midwife may administer Infanrix Hexa (6 in1 vaccine) within their scope of practice.</li> </ul>	
	<b>Note:</b> In determining their scope of practice, nurses and midwives must make judgements about their competency to carry out a role or activity (NMBI, 2015).	
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Description of circumstances and referral arrangements when further advice or consultation is required	<ul> <li>Discuss the vaccine recipien</li> <li>Previous adverse re</li> <li>Other clinical conce</li> </ul>		r in the event of:
Documentation required for the implementation of this medicine protocol	A consent form must be completed by the parent/legal guardian for all children who receive the Infanrix Hexa (6 in 1 vaccine), once understood and translation of consent undertaken with support of translator if required. Appropriate details including the batch number must be recorded on the consent form.         The following documents will be required at each vaccination session:         • Vaccination session report form         • Blank vaccine consent forms         • Vaccine Information Leaflets         • Patient held record cards/ Vaccine passport         • HPRA Adverse Reaction Reporting forms         • HSE Incident/Near Miss Report Forms         • Post vaccination information         It is the responsibility of each registered nurse or midwife to be familiar with the appropriate documentation to support the safe administration of Infanrix Hexa (6 in 1 vaccine) by registered nurses and midwives to Refugees and Applicants Seeking Protection in Ireland and in the event of an outbreak         • NIAC (2023) Anaphylaxis: Immediate Management in the Community available at: https://rcpi.access.preservica.com/uncategorized/IO a36f9e4b-4c80-432d-8264-546089359925/         Infanrix Hexa         Dose: 0.5ml		
	Route: Intramuscular inject Patients Age	Site	Needle length & Size
	Birth to <12 months	Vastus lateralis muscle	25 mm (Use a 16 mm needle in infants under 2.5 - 3 kg) 23-25 gauge
	12 to <36 months	Vastus lateralis or deltoid muscle (depending on muscle mass)	25 mm 23-25 gauge
	3 years and older	Deltoid muscle	25 mm 23-25 gauge
Link to Medicine Details of product	Link to Summary of Production <u>https://www.ema.europa.epar-product-information</u>	eu/en/documents/product-in	formation/infanrix-hexa-



information and other data including instructions for supply and administration is available from the Health Products Regulatory Authority at <u>www.ema.ie</u>	Link to Patient Information Leaflet: https://www.ema.europa.eu/en/documents/product-information/infanrix-hexa- epar-product-information_en.pdf	
Procedure for the reporting and documentation of errors and near misses involving the medicine	In the case of medicine errors that directly involve the vaccine recipient, i.e. wrong medicine/patient/dose/route being administered or another medicine error, the registered nurse or registered midwife must remain with the vaccine recipient and closely monitor them for any adverse reactions.Vital signs should be recorded and the vaccine recipient should be reviewed by the registered nurse/midwife and/or medical practitioner.The incident must be reported to the relevant line manager as soon as possible. The incident and all actions taken must be promptly recorded in the recipient's documentation/notes and the relevant National Incident Management Report Form completed: <a "="" a-z="" emitoolkit="" href="https://www.hse.ie/eng/about/who/nqpsd/qps-incident-&lt;/a&gt;&lt;br/&gt;management/nims/nirf-01-v12-person-interactive.pdfFor children, the child's parent and/or legal guardian must be informed of the incident.Any suspected adverse reactions associated with medicine errors should be reported to&lt;br/&gt;the HPRA as outlined below.Any errors and near misses not involving medication e.g. needle stick injuries, the&lt;br/&gt;incident and all actions taken must be promptly recorded on the relevant National&lt;br/&gt;Incident Management Report form and forwarded to the relevant line manager as per&lt;br/&gt;local policy. Refer to 'EMI Tool Kit' (&lt;a href=" https:="" www.hpsc.ie="">https://www.hpsc.ie/a-z/EMIToolkit/</a> )	
Procedure for reporting Adverse Drug Reactions to the HPRA Resources and equipment required for the	<ul> <li>The relevant nursing or midwifery staff should report to the HPRA any suspected adverse reactions, in accordance with criteria outlined by the HPRA. This reporting may be carried out online at <a href="https://www.hpra.ie">https://www.hpra.ie</a> or through the use of the yellow card system which is available in the downloadable format from the HPRA website, or on request from the HPRA.</li> <li>Infanrix Hexa vaccine</li> <li>Fridge/cool box with minimum/maximum temperature recording device to</li> </ul>	
administration of Infanrix Hexa (6in1) Vaccine	<ul> <li>monitor the cold chain temperature (between +2°C and +8°C)</li> <li>Disposable kidney dishes/coloured trays</li> <li>Gauze swabs/Plasters</li> <li>Sharps bins, and bags for disposal of healthcare risk and non-risk material</li> <li>Alcohol hand sanitiser</li> <li>Face masks (as per local guideline)</li> <li>Access to telephone</li> <li>Resuscitation equipment and drugs in accordance with the NIAC (2023) <i>Anaphylaxis: Immediate management in the Community</i> available at: https://rcpi.access.preservica.com/uncategorized/IO_a36f9e4b-4c80-432d- 8264-546089359925/</li> <li>Access to medical support</li> </ul>	



	Safe storage areas for medicines and equipment			
	<ul> <li>Current medicine protocol for Infanrix Hexa (6 in 1 vaccine).</li> </ul>			
Audit process to identify				
appropriate use of the	All documentation will be held for review and audit purposes as per local policy.			
medicine protocol or	All documentation will be held for review and audit purposes as per local policy.			
unexpected outcomes				
4.0 Information for vaccine	recipient/parent/legal guardian			
Advice to be given to the	Reiterate the information provided in the HSE patient information leaflet for the			
vaccine recipient before	vaccine in the appropriate language and translator support if required.			
treatment				
	For children, Patient Information Leaflet/Fact Sheet <b>must</b> be supplied with the consent			
	form to each parent/legal guardian prior to administration of the vaccine as above.			
Advice to be given to the	After Treatment			
Advice to be given to the vaccine recipient after	The vaccine recipient must be advised to remain seated in the post vaccination			
treatment	observation area for 15 minutes to allow monitoring of any immediate reaction			
treatment	including possible anaphylactic reaction and must be advised to report any side effects			
	to the registered nurse or registered midwife who is present.			
Details of any necessary	In the event of an adverse reaction the registered nurse or midwife must ensure that all			
follow-up, action and	procedures are adhered to as outlined in Section 3.			
referral arrangements				
5.0 Staff authorised to use t				
Professional qualifications,	Registered nurse or registered midwife must have completed all of the following:			
training, and competence				
required prior to using this				
medicine protocol	the NMBI			
	2) Immunisation Foundation Programme accessible on www.HSELanD.ie			
	3) Education programme for nurses and midwives on Primary Childhood Immunisation			
	Programme and any updates for nurses and midwives accessible on www.HSELanD.ie			
	4) Education programme for nurses and midwives on <i>Schools Immunisation Programme</i>			
	and any updates for nurses and midwives accessible on www.HSELanD.ie			
	5) NIO advertise an entropy of Cataly an entropy for Defense and Analismute			
	5) NIO education programme on <i>Catch up vaccination for Refugees and Applicants</i>			
	Seeking Protection in Ireland accessible on www.HSELanD.ie			
	6) An approved Basic Life Support for Health Care Providers Course within the last two			
	years (i.e. Irish Heart Foundation (IHF))			
	7) Initial National Anaphylaxis Education Programme for Health Care Professionals			
	accessible on <u>www.HSELanD.ie</u> followed by a two hour classroom based skills			
	workshop. Recertification is required every two years by completing the on-line			
	National Anaphylaxis Education Programme for Health Care Professionals accessible			
	on <u>www.HSELanD.ie</u>			

8) The registered nurse/midwife must complete the Competency Self-Assessment Form
to vaccinate* Refugees and Applicants Seeking Protection in Ireland *Health Service
Executive (HSE) Primary Childhood Immunisation Programme (PCIP), Schools
Immunisation Programmes (SIP) and the National Immunisation Advisory Committee
(NIAC) Catch Up Immunisation Programme available at <u>www.immunisation.ie</u>

## References

An Bord Altranais (2007) Guidance to Nurses and Midwives on Medication Management. Dublin: An Bord Altranais

Health Products Regulatory Authority available at www.hpra.ie

Infanrix Hexa Summary of Product Characteristics and Patient Information Leaflet, available at: https://www.ema.europa.eu/en/medicines/human/EPAR/infanrix-hexa#product-information-section

National Clinical Guideline No. 30 (2023) – Infection Prevention and Control (IPC) https://www.gov.ie/en/publication/a057e-infection-prevention-and-control-ipc/.

National Immunisation Advisory Committee (2023) *Anaphylaxis: Immediate Management in the Community* available at: <u>https://rcpi.access.preservica.com/uncategorized/IO\_a36f9e4b-4c80-432d-8264-546089359925/</u>

National Immunisation Advisory Committee *Immunisation Guidelines for Ireland*: Royal College of Physicians of Ireland National Immunisation Advisory Committee available at:<u>https://www.rcpi.ie/Healthcare-</u> Leadership/NIAC/Immunisation-Guidelines-for-Ireland

National Immunisation Office (2023/2024) *Supporting Information for Staff: Schools Immunisation Programme* available at: <u>https://www.hse.ie/eng/health/immunisation/hcpinfo/schoolproghcp/supportingdoc.pdf</u>

National Immunisation Office (2023) Clinical information to support healthcare staff to deliver catch-up vaccination for Refugees and Applicants Seeking Protection in Ireland and in the event of an outbreak available at:<u>https://www.hse.ie/eng/health/immunisation/hcpinfo/botpipa/clinicalinfobotpipa.pdf</u>

National Immunisation Office (2022) *Supporting Information for Vaccinations in General Practice* available at: <u>https://www.hse.ie/eng/health/immunisation/infomaterials/gpsupportingdocpci.pdf</u>

Nursing and Midwifery Board of Ireland (2021) *Code of Professional Conduct and Ethics for Registered Nurses and Registered Midwives*. Dublin: Nursing and Midwifery Board of Ireland available at: <u>https://www.nmbi.ie/Standards-Guidance/Code</u>

Nursing and Midwifery Board of Ireland (2020) *Guidance for Registered Nurses and Registered Midwives on Medication Administration*. Dublin: Nursing and Midwifery Board of Ireland available at: <a href="https://www.nmbi.ie/NMBI/media/NMBI/NMBI-Medication-Administration-2020">https://www.nmbi.ie/NMBI/media/NMBI/NMBI-Medication-Administration-2020</a>

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