



Patient Group Direction (PGD)

Administration of low-dose diphtheria, tetanus and inactivated poliomyelitis vaccine (Td/IPV) for the purpose of providing tetanus, diphtheria or polio vaccination for travel indications.

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Version history

Version	Date	Summary of changes
1.0	01/02/22	Version 1.0 new PGD

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Authorisation

PGD for administration of low-dose diphtheria, tetanus and inactivated poliomyelitis vaccine (Td/IPV) for the purpose of providing tetanus, diphtheria and polio vaccination for travel indications.

This Patient Group Direction (PGD) template has been produced by Public Health Scotland to assist NHS boards. NHS Borders has ensured that the final PGD is considered and approved in line with local clinical governance arrangements for PGDs.

The qualified health professionals who may administer low-dose diphtheria, tetanus and inactivated poliomyelitis vaccine (Td/IPV) under this PGD can only do so as named individuals. It is the responsibility of each professional to practice within the bounds of their own competence and in accordance with their own Code of Professional Conduct and to ensure familiarity with the manufacturer's product information/summary of product characteristics (SPC) for all vaccines administered in accordance with this PGD.

NHS board governance arrangements will indicate how records of staff authorised to operate this PGD will be maintained. Under PGD legislation there can be no delegation. Administration of the vaccine has to be by the same practitioner who has assessed the patient under the PGD.

This PGD has been produced for NHS Borders by:

Effective from: 01/02/2022

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Date approved: 21/02/2022		

Review date: 31/01/2024

Clinical situation

Category	Description
Indication	Active immunisation of individuals against tetanus, diphtheria or poliomyelitis.
Inclusion criteria	Valid consent has been given to receive the vaccine.
	Individuals aged 6 years and over who:
	intend to travel to or reside in countries where tetanus, diphtheria, or polio vaccination is currently recommended for travel in accordance with national recommendations issued by TRAVAX www.travax.nhs.uk/destinations/
	the risk of exposure should be determined after careful risk assessment of an individual's itinerary, duration of stay, planned activities and medical history.
	will be travelling to areas where proof of polio vaccination is required
Exclusion criteria	Individuals who:
	are aged less than 6 years
	are pregnant (as pertussis containing vaccine maybe more appropriate)
	have had a confirmed anaphylactic reaction to a previous dose of diphtheria, tetanus or poliomyelitis containing vaccine, including any conjugate vaccines where diphtheria or tetanus toxoid is used in the conjugate
	have had a confirmed anaphylactic reaction to any component of the vaccine, including formaldehyde, neomycin, streptomycin or polymyxin B (refer to relevant SPC)
	have a history of severe (i.e. anaphylactic reaction) to latex where the vaccine is not latex free.
	are suffering from acute severe febrile illness (the presence of a minor infection is not a contraindication for immunisation)
Cautions/need for further advice/ circumstances when further advice should be	The Green Book advises that there are very few individuals who cannot receive low-dose diphtheria, tetanus and inactivated poliomyelitis vaccine (Td/IPV). Where there is doubt, rather than withholding vaccination, appropriate advice should be sought from the lead clinician.
sought from a doctor	If under 10 years of age, ensure UK childhood immunisations are up to date as another tetanus/diphtheria/polio containing vaccine may be more appropriate. Refer to paediatric immunisation team.
	The presence of a neurological condition is not a contraindication to immunisation but if there is evidence of current neurological deterioration,

Category	Description
	deferral of vaccination may be considered, to avoid incorrect attribution of any change in the underlying condition. The risk of such deferral should be balanced against the risk of the preventable infection, and vaccination should be promptly given once the diagnosis and/or the expected course of the condition becomes clear.
	Individuals with immunosuppression can be given Td-IPV containing vaccines although these individuals may not make a full antibody response.
	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.
Action if excluded	Specialist advice must be sought on the vaccine and circumstances under which it could be given as immunisation using a patient specific direction may be indicated. The risk to the individual of not being immunised must be taken into account.
	Document the reason for exclusion and any action taken in accordance with local procedures.
	Inform or refer to the lead clinician in charge.
	In case of postponement due to acute severe febrile illness, advise when the individual can be vaccinated and ensure another appointment is arranged.
	Advise the individual/parent/carer of preventative measures to reduce exposure to polio through careful attention to food and water hygiene and hand washing.
	Advise the individual/parent/carer of preventative measures to reduce exposure to diphtheria by practising good respiratory and hand hygiene, especially in overcrowded or busy places.
	Advise the individual/parent/carer on wound cleansing and seeking medical help for tetanus prone injuries.
	Individuals who have had a confirmed anaphylactic reaction to a previous dose of a Td-IPV containing vaccine or any components of the vaccines should be referred to a clinician for specialist advice and appropriate management.
Action if patient declines	Advise the individual about the protective effects of the vaccine, the risks of infection and potential complications of disease. Advise how future immunisation may be accessed if they subsequently decide to receive the

Category	Description
	vaccine.
	Advise the individual/parent/carer of preventative measures to reduce exposure to polio through careful attention to food and water hygiene and hand washing.
	Advise the individual/parent/carer of preventative measures to reduce exposure to diphtheria by practising good respiratory and hand hygiene, especially in overcrowded or busy places.
	Advise the individual/parent/carer on wound cleansing and seeking medical help for tetanus prone injuries.
	Document advice given and decision reached. Inform or refer to the lead clinician in charge.

Description of treatment

Category	Description
Name of medicine	Adsorbed diphtheria, tetanus, and inactivated poliomyelitis vaccine (Td/IPV): Revaxis®
Form/strength	Suspension for injection in a pre-filled syringe.
Route of administration	Administer by intramuscular injection. The preferred site is the deltoid region of the upper arm.
	For individuals with a bleeding disorder, vaccines normally given by an intramuscular route should be given in accordance with the recommendations in the 'Green Book' Chapter 4
	The vaccine should be visually inspected for particulate matter and discoloration prior to administration. In the event of any foreign particulate matter and/or variation of physical aspect being observed, do not administer the vaccine.
Dosage	0.5ml
Frequency	Single 0.5ml dose
	Reinforcing doses:
	Booster doses are required where immunisation was received more than 10

Category	Description	
	years ago.	
	Polio: Boosting may be required more frequently in those are travelling to areas where proof of polio vaccination is required	
Duration of treatment	See dose and frequency of administration above.	
Maximum or minimum treatment period	See frequency of administration above.	
Quantity to supply/administer	Single 0.5ml dose per administration.	
▼ black triangle medicines	No	
Legal category	Prescription Only Medicine (POM)	
Is the use out with the SPC?	Vaccine should be stored according to the conditions detailed in the Storage section below. However, in the event of an inadvertent or unavoidable deviation of these conditions refer to national Vaccine Incident Guidance. Where vaccine is assessed in accordance with these guidelines as appropriate for continued use this would constitute off-label administration under this PGD.	
Storage requirements	Vaccine should be stored at a temperature of +2° to +8°C. Store in the original packaging to protect from light. Do not freeze.	
	NHS board guidance on Storage and Handling of vaccines should be observed.	
	In the event of an inadvertent or unavoidable deviation of these conditions, vaccine that has been stored outside the conditions stated above should be quarantined and risk assessed for suitability of continued off-label use or appropriate disposal.	
Additional information	Minor illnesses without fever or systemic upset are not valid reasons to postpone immunisation. If an individual is acutely unwell, immunisation may be postponed until they have fully recovered.	
	Adsorbed diphtheria (low dose), tetanus, and inactivated poliomyelitis vaccine (Td/IPV): can be given at the same time as other vaccines.	
	When administering at the same time as other vaccines, care should be taken to ensure that the appropriate route of injection is used for all the vaccinations.	

Category	Description
	The vaccines should be given at separate sites, preferably in different limbs. If given in the same limb, they should be given at least 2.5cm apart. The site at which each vaccine was given should be noted in the individual's records.

Adverse reactions

Category	Description
Warnings including possible adverse reactions and	For full details/information on possible side effects, refer to the marketing authorisation holder's SPC.
management of these	Local reactions following vaccination are very common such as pain, swelling or redness at the injection site. A small painless nodule may form at the injection site.
	Common adverse reactions include pyrexia, headache, vertigo, nausea and vomiting.
	As with all vaccines there is a very small possibility of anaphylaxis and facilities for its management must be available.
	In the event of severe adverse reaction individual should be advised to seek medical advice.
Reporting procedure for adverse reactions	Healthcare professionals and individuals/carers should report suspected adverse reactions to the Medicines and Healthcare products Regulatory Agency (MHRA) using the Yellow Card reporting scheme on http://yellowcard.mhra.gov.uk/
	Any adverse reaction to a vaccine should be documented in accordance with locally agreed procedures in the individual's record and the individual's GP should be informed.
Advice to patient	Written information to be given to individuals:
or carer including written information	Provide manufacturer's consumer information leaflet/patient information leaflet (PIL) provided with the vaccine.
	Individual advice / follow up treatment:
	Inform the individual/carer of possible side effects and their management.
	The individual should be advised to seek medical advice in the event of a severe adverse reaction.
	Inform the individual that they can report suspected adverse reactions

Category	Description	
	to the MHRA using the Yellow Card reporting scheme on: http://yellowcard.mhra.gov.uk .	
	Advise the individual/parent/carer of preventative measures to reduce exposure to polio through careful attention to food and water hygiene and hand washing.	
	Advise the individual/parent/carer of preventative measures to reduce exposure to diphtheria by practising good respiratory and hand hygiene, especially in overcrowded or busy places.	
	Advise the individual/parent/carer on wound cleansing and seeking medical help for tetanus prone injuries.	
Observation following vaccination	Following immunisation patients remain under observation in line with NHS board policy.	
Follow up	Not applicable	
Additional facilities	A protocol for the management of anaphylaxis and an anaphylaxis pack must always be available whenever vaccines are given. Immediate treatment should include early treatment with intramuscular adrenaline, with an early call for help and further IM adrenaline every 5 minutes. The health professionals overseeing the immunisation service must be trained to recognise an anaphylactic reaction and be familiar with techniques for resuscitation of a patient with anaphylaxis.	

Characteristics of staff authorised under the PGD

Category	Description
Professional qualifications	The following classes of registered healthcare practitioners are permitted to administer this vaccine:
	 nurses and midwives currently registered with the Nursing and Midwifery Council (NMC)
	pharmacists currently registered with the General Pharmaceutical Council (GPhC)
	 chiropodists/podiatrists, dieticians, occupational therapists, orthoptists, orthotists/prosthetists, paramedics, physiotherapists, radiographers and speech and language therapists currently registered with the Health and Care Professions Council (HCPC)

Category	Description
	 dental hygienists and dental therapists registered with the General Dental Council
	optometrists registered with the General Optical Council.
Specialist competencies or qualifications	Persons must only work under this PGD where they are competent to do so.
	All persons operating this PGD:
	 must be authorised by name by their employer as an approved person under the current terms of this PGD before working to it
	 must be familiar with the vaccine product and alert to changes in the manufacturers product information/summary of product information,
	must be competent to undertake immunisation and to discuss issues related to immunisation to assess patients for vaccination and obtain consent
	must be competent in the correct storage of vaccines and management of the cold chain if receiving, responsible for, or handling the vaccine
	 must be competent in the recognition and management of anaphylaxis or under the supervision of persons able to respond appropriately to immediate adverse reactions
	must have access to the PGD and associated online resources
	should fulfil any additional requirements defined by local policy
	Employer
	The employer is responsible for ensuring that persons have the required knowledge and skills to safely deliver the activity they are employed to provide under this PGD
	As a minimum, competence requirements stipulated in the PGD must be adhered to.
Continuing education and training	All practitioners operating under the PGD are responsible for ensuring they remain up to date with the use of vaccines included. If any training needs are identified these should be discussed with the individuals in the organisation responsible for authorising individuals to act under this PGD.

Audit trail

Name	Description
Record/ audit trail	Record:
	that valid informed consent was given
	 name of individual, address, date of birth and GP with whom the individual is registered
	name of person that undertook assessment of individual's clinical suitability for vaccine
	name of person that administered the vaccine
	name and brand of vaccine
	date of administration
	dose, form and route of administration of vaccine
	batch number
	where possible expiry date
	anatomical site of vaccination
	advice given, including advice given if excluded or declines immunisation
	details of any adverse drug reactions and actions taken
	administered under PGD
	Records should be kept in line with local procedures.
	Local policy should be followed to encourage information sharing with the individual's General Practice.
	All records should be clear, legible and contemporaneous and in an easily retrievable format.

Additional references

Name	Description
Additional references	Immunisation against Infectious Disease [Green Book]
	https://www.gov.uk/government/collections/immunisation-against-
	infectious-disease-the-green-book
	Immunisation Against Infectious Disease: The Green Book <u>chapter 30</u> <u>Chapter 15</u> and <u>chapter 26</u>
	Vaccination of individuals with uncertain or incomplete immunisation status.
	https://www.gov.uk/government/publications/vaccination-of-
	individuals-with-uncertain-or-incomplete-immunisation-status
	Current edition of British National Formulary.
	Marketing authorisation holder's Summary of Product Characteristics.
	All relevant Scottish Government Health Directorate advice including the relevant CMO letter(s).
	Professional Guidance on the Administration of Medicines in Healthcare Settings 2019
	https://www.rpharms.com/Portals/0/RPS document library/Open
	access/Professional standards/SSHM Admin/Admin Meds prof
	guidance.pdf/ver=2019-01-23-145026-567
	Professional Guidance on the Safe and Secure Handling of Medicines
	https://www.rpharms.com/recognition/setting-professional-
	standards/safe-and-secure-handling-of-medicines/professional-
	guidance-on-the-safe-and-secure-handling-of-medicines
	PHE guidance on the vaccination of individuals with uncertain or incomplete immunisation status:
	https://www.gov.uk/government/publications/vaccination-of-individuals-with-uncertain-or-incomplete-immunisation-status

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Practitioner

This PGD does not remove professional obligations and accountability. It is the responsibility of each professional to practice within the bounds of their own competence and in accordance with their Code of Professional Conduct and to ensure familiarity with the marketing authorisation holder's summary of product characteristics for all vaccines administered in accordance with this PGD.

By signing this Patient Group Direction you are indicating that you agree to its contents and that you will work within it. I agree to administer low-dose diphtheria, tetanus and inactivated poliomyelitis vaccine (Td/IPV) only in accordance with this PGD.

Name of professional	Signature	Date

Authorising Manager

Lead clinician for the service area

I confirm that the practitioners named above have declared themselves suitably trained and competent to work under this PGD. I give authorisation on behalf of **NHS Borders** for the above named health care professionals who have signed the PGD to work under it.

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Signature
Date

Authorised staff should be provided with an individual copy of the clinical content of the PGD and a photocopy of the document showing their authorisation. This authorisation sheet should be retained to serve as a record of those practitioners authorised to work under this PGD.