**Publications number:** **GOV-10648**

**Patient Group Direction (PGD) for the supply of oseltamivir for pre and post exposure of non-H7N9 avian influenza**

For the supply of oseltamivir for the pre and post exposure prophylaxis of non-H7N9 avian influenza, for adults and children aged one year and older, by registered healthcare practitioners identified in [Section 3,](#Section3) subject to any [limitations to authorisation](#limitations) detailed in [Section 2](#Section2).

Reference: 20211203 Oseltamivir non-H7N9 avian influenza PGD

Version no: 05.00

Valid from: 3 December 2021

Review date: 3 December 2023

Expiry date: 2 December 2024

**The UK Health Security Agency (UKHSA) has developed this PGD for local authorisation**

Those using this PGD must ensure it is organisationally authorised and signed in Section 2 by an appropriate authorising person, relating to the class of person by whom the product is to be supplied, in accordance with Human Medicines Regulations 2012 (HMR2012)[[1]](#footnote-1). **The PGD is not legal or valid without signed authorisation in accordance with** [**HMR2012 Schedule 16 Part 2**](http://www.legislation.gov.uk/uksi/2012/1916/schedule/16/part/2/made)**.**

Authorising organisations must not alter, amend or add to the clinical content of this document (sections 4, 5 and 6); such action will invalidate the clinical sign-off with which it is provided.

As operation of this PGD is the responsibility of commissioners and service providers, the authorising organisation can decide which staff groups, in keeping with relevant legislation, can work to the PGD. Sections 2, 3 and 7 must be completed and amended within the designated editable fields provided.

The final authorised copy of this PGD should be kept by the authorising organisation completing Section 2 for 25 years after the PGD expires. Provider organisations adopting authorised versions of this PGD should also retain copies for 25 years after the PGD expires.

**Individual practitioners must be authorised by name, under the current version of this PGD before working according to it.**

Practitioners and organisations must check they are using the current version of the PGD. Amendments may become necessary prior to the published expiry date. Current versions of UKHSA avian influenza PGDs for authorisation can be found from:

<https://www.gov.uk/government/publications/avian-influenza-pre-and-post-exposure-prophylaxis-pgd-template>

Any concerns regarding the content of this PGD should be addressed to: [respiratory.lead@phe.gov.uk](mailto:respiratory.lead@phe.gov.uk)

Enquiries relating to the availability of organisationally authorised PGDs and subsequent versions of this PGD should be directed to: insert local contact details

# **Change history**

|  |  |  |
| --- | --- | --- |
| **Version number** | **Change details** | **Date** |
| 01.00 | Original PGD template | 24 February 2017 |
| 02.00 | * update to off-label use * update to information for individuals with swallowing difficulties * amendment to age range for doses for children * addition of maximum duration of treatment * additional supply and labelling requirements * additional patient information * updates to references * minor typographical changes | 22 January 2018 |
| 03.00 | * addition of doses in renal failure * expansion of definition of immunosuppression * expansion of action to be taken if the patient is excluded * expansion of drug interaction with LAIV | 7 January 2021 |
| 04.00 | * removal of H5N8 as an exclusion * addition of ‘other materials’ to inclusion criteria | 4 March 2021 |
| 05.00 | * amendment of inclusion and exclusion criteria from 7 days or more to 8 days or more * additional information for doses in chronic kidney disease * references to PHE changed to UKHSA * minor rewording of standard text for consistency with other UKHSA PGDs; updated references | 3 December 2021 |

1. **PGD development**

This PGD has been developed by the following on behalf of the UKHSA:

|  |  |  |  |
| --- | --- | --- | --- |
| **Developed by:** | **Name** | **Signature** | **Date** |
| Pharmacist (Lead author) | Jacqueline Lamberty  Lead Pharmacist Medicines Governance, UKHSA |  | 3 December 2021 |
| Doctor | Dr John Astbury  Consultant in Health Protection  Head of Health Protection,  North West (Cumbria and Lancashire) Health Protection Team, UKHSA |  | 3 December 2021 |
| Registered nurse | Shelagh Snape  Senior Health Protection Practitioner  North West (Cumbria and Lancashire) Health Protection Team, UKHSA |  | 3 December 2021 |

This PGD has been peer reviewed by the Avian Influenza PGD Expert panel in accordance with the UKHSA PGD Policy. It has been agreed by the UKHSA Medicines Governance Group and ratified by the UKHSA Clinical Quality and Oversight Board.

**Expert panel**

|  |  |
| --- | --- |
| **Name** | **Designation** |
| Dr Meera Chand (Chair) | Co-director of Clinical and Emerging Infections (Interim), Clinical & Public Health Group, UKHSA |
| Dr Nicholas Aigbogun | Consultant in Communicable Disease Control, Yorkshire and Humber Health Protection Team, UKHSA |
| Dr Colin Brown | Director (Interim): Clinical and Emerging Infections & Deputy Director (Interim): HCAI, Fungal, AMR, AMU, & Sepsis Division, Clinical & Public Health Group, UKHSA |
| Dr Gavin Dabrera | Consultant in Acute Respiratory infections, Clinical and Public Health, UKHSA |
| Dr Emily Dobell | Consultant Epidemiologist, UKHSA |
| Adam John Grainger | Senior Medicines Performance Pharmacist, NHS Midlands and Lancashire CSU |
| Mark McGivern | Consultant in Health Protection, North West Health Protection Team, UKHSA |
| Dr Conall Watson | Consultant Epidemiologist – influenza and seasonal respiratory viruses, Immunisation & Vaccine-Preventable Diseases Division, UKHSA |

1. **Organisational authorisations**

The PGD is not legally valid until it has had the relevant organisational authorisation.

It is the responsibility of the organisation that has legal authority to authorise the PGD, to ensure all legal and governance requirements are met. The authorising body accepts governance responsibility for the appropriate use of the PGD.

Insert authorising body name authorises this PGD for use by the services or providers listed below:

|  |
| --- |
| Authorised for use by the following organisations and/or services |
| For instance, NHSEI services |
| Limitations to authorisation |
| For instance, any local limitations the authorising organisation feels they need to apply in line with the way services are commissioned locally. This organisation does not authorise the use of this PGD by …. |

|  |  |  |  |
| --- | --- | --- | --- |
| Organisational approval (legal requirement) | | | |
| Role | Name | Sign | Date |
| For instance, NHSEI Governance Lead, Medical Director |  |  |  |

|  |  |  |  |
| --- | --- | --- | --- |
| Additional signatories according to locally agreed policy | | | |
| Role | Name | Sign | Date |
|  |  |  |  |
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Section 7 provides a practitioner authorisation sheet. Individual practitioners must be authorised by name to work to this PGD. Alternative practitioner authorisation sheets may be used where appropriate in accordance with local policy, but this should be an individual agreement, or a multiple practitioner authorisation sheet as included at the end of this PGD.

#### Characteristics of staff

|  |  |
| --- | --- |
| **Qualifications and professional registration** | To be completed by the organisation authorising the PGD for instance, registered professional with one of the following bodies:   * nurses currently registered with the Nursing and Midwifery Council (NMC) * pharmacists currently registered with the General Pharmaceutical Council (GPhC)   The practitioners above must also fulfil the [Additional requirements](#addrequirements) detailed below.  Check [Section 2 Limitations to authorisation](#limitations) to confirm whether all practitioners listed above have organisational authorisation to work under this PGD. |
| **Additional requirements** | Additionally, practitioners:   * must be authorised by name as an approved practitioner under the current terms of this PGD before working to it * must have undertaken appropriate training for working under PGDs for supply or administration of medicines * must have undertaken training appropriate to this PGD * must be competent in the use of PGDs (see [NICE Competency framework](https://www.nice.org.uk/guidance/mpg2/resources) for health professionals using PGDs) * must be familiar with the product and alert to changes in the Summary of Product Characteristics * must be competent to assess the individual and discuss treatment options * must have access to the PGD and associated online resources * should fulfil any additional requirements defined by local policy * insert any additional requirements   **The individual practitioner must be authorised by name, under the current version of this PGD before working according to it.** |
| **Continued training requirements** | insert any continued training requirements |

1. **Clinical condition or situation to which this PGD applies**

|  |  |
| --- | --- |
| **Clinical condition or situation to which this PGD applies** | Pre and post exposure prophylaxis of non-H7N9 avian influenza as advised by the UKHSA. |
| **Criteria for inclusion[[2]](#footnote-2)** | Adults and children (one year of age or older) who have or will have:   * handled or been in close contact with live, sick, dying or dead birds infected or potentially infected with non-H7N9 avian influenza or * handled or been in close contact with faecal matter or contaminated litter/other materials from birds infected or potentially infected with non-H7N9 avian influenza * swabbed, culled or removed carcasses of birds infected or potentially infected with non-H7N9 avian influenza or * had a significant exposure as advised by the local UKHSA Health Protection Team   **unless:**   * 8 days or more have elapsed since the last exposure |
| **Criteria for exclusion**  Continued overleaf  **Criteria for exclusion**  (continued) | Individuals:   * with exposure to suspected or confirmed H7N9 avian influenza * whose last exposure was 8 days or more previously * who are aged under one year * with a body weight less than 10 kg * who have a known allergy or hypersensitivity to oseltamivir or to any of the excipients * with severe renal disease requiring haemodialysis * who are immunocompromised[[3]](#footnote-3) due to disease or treatment for instance: * severe primary immunodeficiency * current or recent (within 6 months) chemotherapy or radiotherapy for malignancy * solid organ transplant recipients on immunosuppressive therapy * bone marrow transplant recipients currently receiving immunosuppressive treatment, or within 12 months of receiving immunosuppression * individuals with current graft-versus-host disease * individuals currently receiving high dose systemic corticosteroids (equivalent to ≥40 mg prednisolone per day for >1 week in an adult, or ≥ 2mg/kg/day for ≥1 week in a child), and for at least 3 months after treatment has stopped * HIV infected individuals with severe immunosuppression (CD4<200/μl or <15% of total lymphocytes in an adult or child over 5; CD4< 500/μl or <15% of total lymphocytes in a child aged 1 to 5; expert clinical opinion in a child aged under 1) * individuals currently or recently (within 6 months) on other types of highly immunosuppressive therapy or where the individual’s specialist regards them as severely immunosuppressed. * who are taking other drugs with clinically significant drug interactions for instance, chlorpropamide, methotrexate, phenylbutazone |
| **Action to be taken if the patient is excluded** | For suspected or confirmed exposure to H7N9 avian influenza: use the [Oseltamivir H7N9 avian influenza PGD](https://www.gov.uk/government/publications/avian-influenza-pre-and-post-exposure-prophylaxis-pgd-template)  Where exposure was 8 days or more previously: inform the individual prophylaxis is not indicated beyond 7 days following exposure.  For individuals aged under one year, or with a body weight of less than 10kg, or with a known allergy or hypersensitivity to oseltamivir or to any of the excipients, or those who require haemodialysis: refer to a medical practitioner. A Patient Specific Direction (PSD) would be required for any alternative dosage or treatment recommended.  For individuals who specify a history of immunosuppression due to disease or treatment, discuss with a Consultant in Health Protection or a Consultant Virologist / Microbiologist for advice. Depending on the nature of the immunosuppression, discussion may be needed on a case by case basis between the Health Protection Team and specialists such as Consultant Virologists, Microbiologists or Epidemiologists. Some individuals might need a different dose, some might need an alternative medicine or, for some, complete cessation of all exposures, if possible, may be advised. A PSD would be required for any alternative dosage or treatment recommended as a result of this discussion.  Some individuals excluded under this PGD may be suitable for pre or post exposure prophylaxis if prescribed. Refer to a medical practitioner without delay. |
| **Action to be taken if the patient or carer declines prophylaxis** | Advise the individual or carer of the possible consequences of refusing treatment and of alternative sources of treatment.  Advise about the protective effects of the treatment, risks of infection, risk of spreading the disease to others and disease complications.  Document refusal and advice given.  Inform the relevant local Health Protection team and, if appropriate, refer toa medical practitioner for an alternative treatment. |
| **Cautions** | Refer individuals to a medical practitioner if:   * they are exhibiting sudden onset of symptoms of confusion, chest pain, breathing difficulties or any other symptoms giving cause for concern * they have long term conditions such as chronic respiratory or cardiovascular disease exhibiting rapidly worsening symptoms |

1. **Description of treatment**

|  |  |
| --- | --- |
| **Name, strength & formulation of drug** | Oseltamivir 75mg, 45mg and 30mg capsules |
| **Legal category** | POM - Prescription only medicine |
| **Black triangle▼** | No |
| **Off-label use** | Yes.  Oseltamivir is not licensed for avian influenza. [National UK guidance](https://assets.publishing.service.gov.uk/government/uploads/system/uploads/attachment_data/file/968566/Avian_influenza_guidance_and_algorithms_for_managing_incidents_in_birds.pdf) recommends chemoprophylaxis with oseltamivir as per the inclusion criteria.  Consider, as part of the consent process, informing the individual or their carer the product is offered in accordance with national guidance, but this is outside the product licence. |
| **Route / method of administration** | Oral.  The individual should start the medication as soon as possible.  The capsules should be swallowed whole with water.  For individuals with swallowing difficulties, the capsules can be opened and the contents mixed with a small amount of sweetened food, such as chocolate or cherry syrup, and dessert toppings such as caramel or fudge sauce or sugared water, just before administration (see [Patient Information Leaflet](https://www.medicines.org.uk/emc/)). |
| **Dose and frequency of administration**  Continued overleaf  **Dose and frequency of administration**  (continued) | **Adults and children aged 13 years and older:** see table below  The capsules should preferably be taken in the morning with breakfast, for the duration of treatment. Taking with food can reduce nausea or vomiting.   |  |  | | --- | --- | | **Renal function[[4]](#footnote-4)** | **Dose** | | No known chronic renal impairment | One 75mg capsule once a day | | Moderate impairment  (CrCl 31-60 mL/min) | One 30mg capsule once a day | | Severe impairment  (CrCl 11-30mL/min) | One 30mg capsule every 48 hours | | Established renal failure  (CrCl ≤10mL/min) | One 30mg capsule once, repeated every 7 days | | Haemodialysis | Refer to a medical practitioner; do not supply under this PGD | | Peritoneal dialysis | One 30mg capsule once, repeated every 7 days |   The doses given above are for individuals with stable chronic kidney disease. If there is a history of renal failure, supply as per the latest documented creatinine clearance (CrCl) results.  Estimated glomerular filtration rate (eGFR) may be more readily available. If eGFR is the only value available, do not delay chemoprophylaxis and supply a dose according to eGFR (substituting eGFR for the CrCL figure in the table above). Some individuals may receive a larger oseltamivir dose as a result, but this is unlikely to be harmful as clinical experience reveals a wide margin of safety.  For children with renal dysfunction aged less than 13 years, adjust the oseltamivir dose as per the [Oseltamivir chapter in the British National Formulary (BNF) for children.](https://bnfc.nice.org.uk/drug/oseltamivir.html#renalImpairment)  If CrCl or eGFR results are not known, refer to a medical practitioner. If a decision to supply is made, a Patient Specific Direction (PSD) will be required.  **For adults with a body weight less than 40 kg and children aged from 1 year to 12 years of age**: refer to the table below   |  |  | | --- | --- | | **Body Weight** | **Dose, preferably in the morning with breakfast** | | 10 kg to 15 kg | 30 mg once daily | | > 15 kg to 23 kg | 45 mg once daily | | > 23 kg to 40 kg | 60 mg once daily | | > 40 kg | 75 mg once daily |   If the child has a body weight less than 10 kg, they are excluded from this PGD. Refer them to a medical practitioner.  If the body weight cannot be determined and the child appears to be of average weight for their age, use the table below:   |  |  | | --- | --- | | **Age** | **Dose, preferably in the morning with breakfast** | | 1 to 3 years | 30 mg once daily | | 4 to 6 years | 45 mg once daily | | 7 to 12 years | 60 mg once daily | | Over 12 years | 75 mg once daily |   No dose adjustment is needed in obese individuals |
| **Duration of prophylaxis** | Individuals need to receive prophylaxis to cover the total exposure period and for 10 days following the last known exposure.  Once a worker has ended their exposure, any remaining doses should be properly disposed of by returning them to a community pharmacy for destruction.  The maximum period of treatment that an individual can receive for a single incident through this PGD is 42 days. |
| **Quantity to be supplied**  Continued overleaf  **Quantity to be supplied**  (continued) | Adults: sufficient to cover the duration of prophylaxis as above.  For adults with a body weight less than 40 kg and children aged from 1 year to 12 years of age: refer to the table overleaf.   |  |  |  | | --- | --- | --- | | **Body Weight** | **Age** | **Quantity of capsules to be supplied**  **for each 10 days of prophylaxis** | | 10 kg to 15 kg | 1 to 3 years | 10 x 30 mg | | > 15 kg to 23 kg | 3 to 6 years | 10 x 45 mg | | > 23 kg to 40 kg | 7 to 12 years | 20 x 30 mg | | > 40 kg | Over 12 years | 10 x 75 mg |   Renal impairment:   |  |  | | --- | --- | | **Quantity of capsules to be supplied** | **Quantity of capsules to be supplied for each 10 days of prophylaxis** | | Moderate impairment | 10 x 30mg | | Severe impairment | 5 x 30mg | | Established renal failure | 2 x 30mg | | Peritoneal dialysis | 2 x 30mg |   When supplying under PGD, this should be from the manufacturer’s original pack or over-labelled pre-packs so that the individual’s name, date and additional instructions can be written on the label at the time of supply. As split packs cannot be supplied, an over-supply might be required. |
| **Storage** | Medicines must be stored securely according to national guidelines and in accordance with the product’s SPC. Do not store above 25oC. |
| **Disposal** | Any unused product or waste material should be disposed of in accordance with local arrangements. |
| **Drug interactions** | Individuals taking the following medicines are excluded from this PGD (see [exclusion criteria](#drugsexclusion)):   * chlorpropamide * methotrexate * phenylbutazone   [The Green Book](https://www.gov.uk/government/publications/influenza-the-green-book-chapter-19) states administration of influenza antiviral agents within two weeks of administration of a live attenuated influenza vaccine (LAIV) may adversely affect the effectiveness of the vaccine. Therefore, oseltamivir and LAIV should not be administered concomitantly. LAIV should be delayed until 48 hours following the cessation of treatment with oseltamivir.  If LAIV has been given in the past two weeks, the individual may need to be revaccinated with another appropriate influenza vaccine and medical advice should be obtained. |
| **Identification & management of adverse reactions**  (continued overleaf)  **Identification & management of adverse reactions**  (continued) | Very common (≥ 1/10) and common (≥ 1/100 to < 1/10) adverse reactions include nausea, vomiting, headache, abdominal pain and dyspepsia.  These reactions may only occur on a single occasion, on either the first or second day of treatment, and resolve spontaneously within 1-2 days. However, if symptoms persist individuals should consult a healthcare professional.  Individuals should be advised not to discontinue treatment without consulting a doctor or pharmacist.  Other commonly reported adverse reactions include bronchitis, dizziness (including vertigo), fatigue, insomnia, herpes simplex, nasopharyngitis, upper respiratory tract infections, sinusitis, cough, sore throat, pyrexia, rhinorrhoea, pain including limb pain.  A detailed list of adverse reactions is available in the [SPC](http://www.medicines.org.uk/) |
| **Reporting procedure of adverse reactions** | Any adverse reaction to the product should be documented in the medical records.  Alert a doctor in the event of serious adverse reaction.  Healthcare professionals and individuals/parents/carers are encouraged to report all suspected adverse reactions in children and severe adverse reactions in adults to the Medicines and Healthcare products Regulatory Agency (MHRA) using the [Yellow Card](http://yellowcard.mhra.gov.uk) reporting scheme or search for MHRA Yellow Card in the Google Play or Apple App Store. |
| **Written information to be given to patient or carer** | Supply the marketing authorisation holder's patient information leaflet (PIL).  Each individual should be given a copy of the Information for contact of avian influenza available from [Managing the human health implications of avian influenza - guidance for health protection teams (publishing.service.gov.uk)](https://assets.publishing.service.gov.uk/government/uploads/system/uploads/attachment_data/file/968566/Avian_influenza_guidance_and_algorithms_for_managing_incidents_in_birds.pdf) |
| **Patient advice /follow up treatment** | Advise the individual or their carer:   * taking the medication with a small amount of food can reduce nausea or vomiting * the capsules can be opened and taken with a small amount of sweetened food as explained in the PIL * of any possible side effects and their management * to seek medical advice in the event of a severe adverse reaction * to seek advice if common side effects do not spontaneously resolve 48 hours after they first appear, but to continue taking the medicine * to take the medication for the specified number of days * to read the PIL leaflet before taking the medication * consider explaining the PIL does not mention avian influenza because the manufacturer has not sought a product license for this indication, but national guidance recommends the use of this medicine in these circumstances and it is deemed best practice * to seek medical advice if they experience influenza symptoms within 10 days of last exposure to source of non-H7N9 infection * if an over-supply has been required, to take any remaining capsules to a community pharmacy for destruction |
| **Additional information**  Continued overleaf  **Additional information**  (continued) | Pregnancy: oseltamivir is considered safe for use in pregnancy. Recent studies suggest there is no evidence of harm in pregnant women treated with oseltamivir, however published data is limited.  Breastfeeding: oseltamivir is considered acceptable for use in breastfeeding mothers. The benefits of breastfeeding are considered to outweigh any, albeit unidentified, risks. Use of oseltamivir is not a reason to discontinue or put limitations on breastfeeding.  Oseltamivir and its active metabolite are excreted into human breast milk in very small amounts. Limited data suggest clinical sequelae from maternal use would not be expected in a breastfed infant.  The UK Drugs in Lactation Advisory Service (UKDILAS) advises, as a precaution, infants should be monitored for vomiting or diarrhoea. This guidance applies to infants born full term and healthy. If an infant is unwell, premature, or the mother is taking multiple medicines, then an individual risk assessment will need to be made. |
| **Records** | Record:   * whether valid informed consent was given or a decision to supply was made in the individual’s best interests in accordance with the [Mental Capacity Act 2005](https://www.legislation.gov.uk/ukpga/2005/9/contents) * name of individual, address, date of birth and GP with whom the individual is registered (or record where an individual is not registered with a GP) * name of the healthcare practitioner who supplied the product * name and brand/manufacturer of the product * date of supply * dose, form and route of administration of the product * quantity supplied * batch number and expiry date * advice given, including advice given if the individual is excluded or declines treatment * details of any adverse drug reactions and actions taken * record the product was supplied via PGD   Records should be signed and dated (or password-controlled record on e-records).  All records should be clear, legible and contemporaneous  A record of all individuals receiving treatment under this PGD should also be kept for audit purposes in accordance with local policy. |

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#### Key references

|  |  |
| --- | --- |
| **Key references** | * [Summary of Product Characteristics](http://www.medicines.org.uk/) accessed 1 December 2021 * [Patient Information Leaflet](http://www.medicines.org.uk/) accessed 1 December 2021 * [Managing the human health implications of avian influenza in poultry and wild birds](https://www.gov.uk/government/publications/avian-influenza-guidance-and-algorithms-for-managing-incidents-in-birds) Guidance for health protection teams Version 53.0 March 2021 * [Guidance: Investigation and initial clinical management of possible human cases of avian influenza with potential to cause severe human disease](https://www.gov.uk/government/publications/avian-influenza-guidance-and-algorithms-for-managing-human-cases/investigation-and-initial-clinical-management-of-possible-human-cases-of-avian-influenza-with-potential-to-cause-severe-human-disease) Updated 17 November 2021 * [HSE guidance: Avoiding the risk of infection when working with poultry that is suspected of having H5 or H7 notifiable avian influenza](http://www.hse.gov.uk/biosafety/diseases/aisuspected.pdf) accessed 1 December 2021 * [Influenza: treatment and prophylaxis using anti-viral agents](https://www.gov.uk/government/publications/influenza-treatment-and-prophylaxis-using-anti-viral-agents) updated November 2021 * [Influenza: the green book, chapter 19](https://www.gov.uk/government/publications/influenza-the-green-book-chapter-19) last updated 29 October 2020 * [Oseltamivir or zanamivir - can they be used in breastfeeding mothers for the treatment or prophylaxis of influenza.](https://www.sps.nhs.uk/articles/oseltamivir-or-zanamivir-can-mothers-breastfeed-after-treatment-for-influenza-2/) 7 August 2020 * [British National Formulary (BNF) and British National Formulary for children (BNFc)](https://bnf.nice.org.uk/) accessed 1 December 2021 * [NICE Medicines Practice Guideline 2 (MPG2): Patient Group Directions](https://www.nice.org.uk/guidance/mpg2) last updated 27 March 2017 * [NICE MPG2 Patient group directions: competency framework for health professionals using patient group directions](https://www.nice.org.uk/guidance/mpg2/resources) last updated 27 March 2017 * [Health Technical Memorandum 07-01: Safe Management of Healthcare Waste.](https://www.england.nhs.uk/estates/health-technical-memoranda/) Department of Health 20March 2013 |

1. **Individual practitioner authorisation sheet**

By signing this PGD you are indicating you agree to the contents and you will work within it

PGDs do not remove inherent professional obligations or accountability

It is the responsibility of each professional to practice only within the bounds of their own competence

**Practitioner**

**I confirm I have read and understood the content of this PGD and I am willing and competent to work to it within my professional code of conduct**

Signed……………………………….………………………….…..Date……….….…………..............

Name (Print)…………….…………..………….………………………………………….…….............

Designation……………………………………………………………….…..………………................

**Authorising manager**

Manager to give authorisation on behalf of **Insert name of organisation** for the named healthcare professional who has signed the PGD

Signed…………………………………….………………………. Date………………………..........

Name (Print)………………………..…………………………………….……………..………..........

Designation………………………………………………………………..…………….…….............

**Note to authorising manager**

By signing above, you are confirming you have assessed the staff member as competent to work under this PGD and they have the organisational approval to do so

You must give this signed PGD to each authorised practitioner as it shows their authorisation to use the PGD

1. This includes any relevant amendments to legislation (such as [2013 No.235](http://www.legislation.gov.uk/uksi/2013/235/contents/made), [2015 No.178](http://www.legislation.gov.uk/nisr/2015/178/contents/made), [2015 No.323](http://www.legislation.gov.uk/uksi/2015/323/contents/made) and [the Human Medicines (Coronavirus and Influenza) (Amendment) Regulations 2020](https://www.legislation.gov.uk/uksi/2020/1125/contents/made) [↑](#footnote-ref-1)
2. Criteria for post exposure antiviral prophylaxis can be discussed with the local Health Protection Team. [↑](#footnote-ref-2)
3. [UKHSA Guidance on use of antiviral agents for the treatment and prophylaxis of seasonal influenza Version 11, November 2021](https://assets.publishing.service.gov.uk/government/uploads/system/uploads/attachment_data/file/1037465/ukhsa-guidance-antivirals-influenza-11v4.pdf) [↑](#footnote-ref-3)
4. [UKHSA Guidance on use of antiviral agents for the treatment and prophylaxis of seasonal influenza Version 11, November 2021](https://assets.publishing.service.gov.uk/government/uploads/system/uploads/attachment_data/file/1037465/ukhsa-guidance-antivirals-influenza-11v4.pdf) [↑](#footnote-ref-4)