**COVID-19 mRNA Vaccine BNT162b2 Pfizer/BioNTech Vaccine**

**National Protocol**

Reference no: COVID-19 mRNA Vaccine BNT162b2 Pfizer/BioNTech Vaccine Protocol

Version no: v01

Valid from: 21January

Review date: 30 November 2021

Expiry date: 31 December 2021

**1. About the National Protocol**

This protocol is for the supply and administration of COVID-19 mRNA Vaccine BNT162b2 Pfizer/BioNTech to individuals in accordance with the national COVID-19 vaccination programme. This protocol only allows administration during or in anticipation of COVID-19 pandemic where the disease represents a serious risk or potentially serious risk to human health.

This protocol is for the supply and administration of COVID-19 mRNA Vaccine BNT162b2 Pfizer/BioNTech by appropriately trained persons in accordance with [regulation 247A](https://www.legislation.gov.uk/uksi/2020/1125/regulation/14/made) of the [Human Medicines Regulation 2012](https://www.legislation.gov.uk/uksi/2012/1916/contents), as inserted by [The Human Medicines (Coronavirus and Influenza) (Amendment) Regulations 2020](https://www.legislation.gov.uk/uksi/2020/1125/contents/made)

The Scottish Government has developed this protocol which has been approved by the Scottish Ministers to facilitate the delivery of the national COVID-19 vaccination programme by Health Boards in Scotland and any organisation a Health Board makes arrangements with to deliver such services on its behalf, referred to as “the provider”. Please note that in the context of this protocol, “the provider” means either the Health Board or an organisation delivering services on behalf of a Health Board.

This protocol may be followed wholly from patient assessment through to post-vaccination by a single person. Alternatively, obtaining consent and patient assessment may be undertaken by a registered healthcare practitioner with the process of administration undertaken by a non-registered practitioner under clinical supervision.

Where multiple person models are used the provider must ensure that all elements of the protocol are complied with in the provision of vaccination to each patient.

The provider is responsible for ensuring that persons are trained and competent to safely deliver the activity they are authorised to provide under this protocol. As a minimum, competence requirements stipulated in the protocol under ‘Characteristics of staff’ must be adhered to.

The provider must identify a clinical supervisor who has overall responsibility for provision of vaccination under the protocol at all times. The clinical supervisor must be a registered healthcare professional trained and competent in all aspects of the protocol and provide clinical supervision for the overall provision of clinical care provided under the protocol.

The clinical supervisor must be identifiable to service users. Whenever the protocol is used, the name of the clinical supervisor taking responsibility and all of the people working under different activity stages of the protocol must be recorded for the session using the schedule in Annex C or maintaining an equivalent electronic record. The clinical supervisor has ultimate responsibility for safe care being provided under the terms of the protocol. Persons working under the protocol may be supported by additional registered healthcare professionals, but the clinical supervisor retains responsibility.

Persons working to the protocol must understand who the clinical supervisor for their practice is at any time and can only work under their authority. The clinical supervisor may withdraw this authority for all persons or individual persons at any time and has authority to stop and start service provision under the protocol as necessary. All members of staff have a responsibility to, and should, report immediately to the clinical supervisor any concerns they have about working under the protocol in general or about a specific individual, process, issue or event.

Individual practitioners must be designated by name to work to this protocol. Individuals working in accordance with this protocol must ensure they meet the staff characteristics for the activity they are undertaking, make a declaration of competence and be authorised in writing by the provider. This can be done by completing Annex B of this protocol or maintaining an equivalent electronic record.

It is a Health Board’s responsibility to adhere to this protocol. Where the Health Board is not the provider, it is the Health Board’s responsibility to ensure that the provider adheres to this protocol. The final authorised copy of this protocol should be kept, by Health Boards for 8 years after the protocol expires. Providers adopting authorised versions of this protocol should also retain copies, along with the details of those authorised to work under it, for 8 years after the protocol expires.

Providers must check that they are using the current version of this protocol. Amendments may become necessary prior to the published expiry date. Current versions of protocols authorised by the Scottish Ministers in accordance with regulation 247A of the Human Medicines Regulations 2012 can be found on the Scottish Government website: (tbc)

Any concerns regarding the content of this protocol should be addressed to: [VaccinationsDelivery@gov.scot](mailto:VaccinationsDelivery@gov.scot)

**2. Approval and Clinical Authorisation**

This protocol is not legally valid, in accordance with [regulation 247A](https://www.legislation.gov.uk/uksi/2020/1125/regulation/14/made) of [Human Medicines Regulation 2012](https://www.legislation.gov.uk/uksi/2012/1916/contents), as inserted by [The Human Medicines (Coronavirus and Influenza) (Amendment) Regulations 2020](https://www.legislation.gov.uk/uksi/2020/1125/contents/made), until approved by the Scottish Ministers.

On 21 January 2021 the Scottish Ministers, approved this protocol in accordance with [regulation 247A](https://www.legislation.gov.uk/uksi/2020/1125/regulation/14/made) of Human Medicines Regulation 2012. Approval of clinical information in Annex A is via the Scottish Government Chief Medical Officer, Chief Pharmaceutical Officer and Chief Nursing Officer for the delivery of the national COVID-19 vaccination programme, with defined limitations to authorisation that may be updated from time to time as may be required.

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| Authorised for use by the following organisations and/or services |
| All Health Boards in Scotland, and organisations Health Boards make arrangements with to deliver services on their behalf. |
| Limitations to authorisation |
| This authorisation applies to the supply and administration of the vaccine(s) only under the conditions set out in the authorisation for supply or license set out by the Medicines and Healthcare products Regulatory Agency |

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| **Clinical authorisation** | | | |
| **Role** | **Name** | **Sign** | **Date** |
| CMO | **Gregor Smith** |  | **19/01/21** |
| CNO | **Fiona Mcqueen** |  | **19/01/21** |
| CPO | **Alison Strath** |  | **19/01/21** |

It is Health Boards’ responsibility to ensure they and any organisations they make arrangements with to deliver services on their behalf operate the specified vaccination services in accordance with the protocol. Any provider administering COVID-19 mRNA Vaccine BNT162b2 Pfizer/BioNTech vaccine under protocol must work strictly within the terms of this protocol.

The national COVID-19 vaccination programme may also be provided under patient group direction, under written instruction for supply and administration in the course of an occupational health scheme, or on a patient specific basis, by or on the directions of an appropriate prescriber. Supply and administration in these instances are not related to this protocol.

**3. Change history**

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| **Version number** | **Change details** | **Date** |
| V01.00 | New protocol for COVID-19 mRNA Vaccine BNT162b2 Pfizer/BioNTech vaccine. | 21 January 2021 |

**4. Characteristics of staff**

The Provider is responsible for the designation and authorisation of persons within the classes set out below permitted to administer medicinal products under this protocol. In doing so the provider must establish that those persons

1. demonstrate appropriate knowledge and skills to work under the National Protocol for the supply/administration of COVID-19 vaccine.
2. have met the requirements of the NES Proficiency document -COVID-19 vaccine administration for registered staff or the NES Proficiency document –COVID-19 vaccine administration - Healthcare support workers as appropriate <https://learn.nes.nhs.scot/37676/immunisation/covid-19-vaccines>

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| Classes of persons permitted to administer medicinal products under this protocol |
| This protocol may be adhered to wholly from assessment through to post-vaccination by a single appropriately specified registered healthcare professional. Alternatively, multiple persons may undertake specific activity stages in the vaccination pathway in accordance with this protocol.  Activity stages of the vaccination pathway under this protocol   |  |  |  | | --- | --- | --- | | Stage 1 | 1. Assessment of the individual presenting for vaccination 2. Provide information and obtain informed consent 3. Provide advice to the individual | Registered Healthcare Professionals Only | | Stage 2 | * Vaccine Preparation | Registered or non-registered persons | | Stage 3 | * Vaccine Administration | Registered or non-registered persons | | Stage 4 | * Record Keeping | Registered or non-registered persons |   Providers are responsible for assessing the competency of, designating and recording the names of all those persons permitted to supply and administer under this protocol.  The following specified registered health professionals are permitted to administer under the protocol subject to the requirements set out below:   * 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, operating department practitioners, orthoptists, orthotists/prosthetists, paramedics, physiotherapists, radiographers and speech and language therapists currently registered with the Health and Care Professions Council (HCPC). * Dental hygienists and dental therapists registered with the General Dental Council. * Optometrists registered with the General Optical Council. * Doctors currently registered with General Medical Council. * Dentists currently registered with General Dental Council.   The following professionals (who are in the main non-registered) are permitted to administer under the protocol with appropriate supervision as set out below, subject to the requirements set out below   * Healthcare support workers. * Pharmacy technicians, provisionally registered pharmacists, pre-registration pharmacists and other pharmacy support practitioners. * Retired clinical practitioners such as doctors, dentists, pharmacists, nurses, optometrists, chiropodists/podiatrists, dieticians, occupational therapists, orthoptists, orthotists/prosthetists, paramedics, pharmacy technicians, physiotherapists, radiographers, speech and language therapists, dental hygienists and dental therapists not currently registered. * Student doctors, dentists, pharmacists, nurses, midwives, optometrists, chiropodists/podiatrists, dieticians, occupational therapists, orthoptists, orthotists/prosthetists, paramedics, physiotherapists, radiographers, speech and language therapists, dental hygienists and dental therapists not currently registered. * Healthcare Scientists. * Dental nurses. * Physician’s assistants.   All those working under this protocol must have undertaken training, be assessed as competent and receive supervision appropriate to the stage of activity they are undertaking. Where multiple person models are used, the provider must ensure that all elements of the protocol are complied with in the provision of vaccination to each individual. The provider is responsible for ensuring that persons are trained and competent to safely deliver the activity they are employed to provide under this protocol. As a minimum, competence requirements stipulated in the protocol must be adhered to.  All persons must be designated by name by the provider as an approved person under the current terms of this protocol before working to it, and listed on the practitioner authorisation sheet in Annex B. All staff listed on the sheet will be covered by NHS indemnity extended by the Health Board who is responsible for the COVID 19 vaccination programme in that locality. Protocols do not remove inherent obligations or accountability. All practitioners operating under this protocol must work within their terms of employment at all times; registered healthcare professionals should also abide by their professional code of conduct.  There are three underpinning principles to which every person undertaking activities under the remit of this protocol must adhere  1. Training   * They must have undertaken training appropriate to this protocol and relevant to their role, as required by local policy and health board standard operating procedures and in line with the training recommendations for COVID-19 vaccinators. * They must have met the requirements set out in the NES Proficiency document -COVID-19 vaccine administration for registered staff or the NES Proficiency document –COVID-19 vaccine administration- Healthcare support workers   2. Competency   * Those providing clinical supervision to those administering the vaccine must be competent to assess individuals for suitability for vaccination, identify any contraindications or precautions, discuss issues related to vaccination and obtain informed consent from the individuals being vaccinated. * All persons must either be an appropriate prescriber or one of above noted registered professionals. Those that are not registered professionals, and those returning to immunisation after a prolonged interval (more than 12 months), should be assessed and signed off as meeting the requirements of the relevant NES Proficiency document -COVID-19 vaccine administration. They should be observed administering the vaccine until both they, and their supervisor or trainer, feel confident that they have the necessary knowledge and skills to administer vaccines safely and competently. * Experienced vaccinators should use the relevant NES Proficiency document to self-assess that they are able to meet all the competencies listed and confirm that they have the knowledge and skills necessary to administer COVID-19 vaccine. * They must have completed local IPC training and comply with the vaccination guidance with the National COVID-19 IPC guidelines available:  [National Infection Prevention and Control Manual: Scottish COVID-19 Infection Prevention and Control Addendum for Acute Settings](http://www.nipcm.hps.scot.nhs.uk/scottish-covid-19-infection-prevention-and-control-addendum-for-acute-settings/)   In addition and where indicated as relevant to the role-   * They must be familiar with the vaccine product and alert to any changes in the manufacturers summary of product characteristics (SPC), should it become licensed, or the Regulation 174 Information for UK Healthcare Professionals and familiar with the national recommendations for the use of this vaccine. * They must be familiar with, and alert to changes in relevant chapters of Immunisation Against Infectious Disease: the Green Book [COVID-19: the green book, chapter 14a - GOV.UK (www.gov.uk)](https://www.gov.uk/government/publications/covid-19-the-green-book-chapter-14a). * They must be familiar with, and alert to changes in the relevant provider’s standard operating procedures (SOPs) and provider’s arrangements for the national COVID-19 vaccination programme * They must be competent in the correct handling and storage of vaccines and management of the cold chain if receiving, responsible for, or handling the vaccine. * They must be competent in the recognition and management of anaphylaxis, have completed basic life support training and be able to respond appropriately to immediate adverse reactions. * They must have access to the provider’s protocols and relevant COVID-19 vaccination programme online resources. * They must be competent in intramuscular injection technique if they are administering the vaccine, this should include a practical element. * For those preparing the vaccine, they must be competent in the handling of the vaccine product, procedure for dilution of the vaccine and use of the correct technique for drawing up the correct dose. * For those in record keeping roles, they must understand the importance of making sure vaccine information is recorded on the vaccination management app. * They should fulfil any additional requirements defined by local policies developed in accordance with any national guidance.   3. Supervision   * A period of supervised practice to allow observation of, and development of skills in vaccine administration and application of knowledge to practice is essential. Supervision for new immunisers and support for all immunisers is critical to the safe and successful delivery of the COVID-19 immunisation programme. * Non-registered persons must be supervised and supported by a registered healthcare professional at all times. * The clinical supervisor must be a registered healthcare professional trained and competent in all aspects of the protocol and provide clinical supervision for the overall provision of clinical care provided under the protocol. |

**5. Clinical condition or situation to which this Protocol applies**

COVID-19 mRNA Vaccine BNT162b2 Pfizer/BioNTech is indicated for active immunisation against COVID-19 disease caused by SARS-CoV-2 virus in accordance with Scottish Government COVID-19 immunisation programme and recommendations given in Chapter 14a of the Immunisation Against Infectious Disease: the ‘Green Book’ [COVID-19: the green book, chapter 14a - GOV.UK (www.gov.uk)](https://www.gov.uk/government/publications/covid-19-the-green-book-chapter-14a) and Scottish Government CMO letters relating to COVID-19 vaccination.

**ANNEX A: Clinical Information**

This Annex provides information about the clinical situation or condition and treatment in relation to the National Protocol.

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

| Category | Description |
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| **Indication** | COVID-19 mRNA Vaccine BNT162b2 is indicated for active immunisation against COVID-19 disease caused by SARS-CoV-2 virus in accordance with Scottish Government COVID-19 immunisation programme and recommendations given in Chapter 14a of the Immunisation Against Infectious Disease: the ‘Green Book’, JCVI statement on priority groups for COVID-19 vaccination from 30th December 2020 and subsequent correspondence/publications from Scottish Government. |
| **Inclusion criteria** | National policy must be followed in relation to the priority groups eligible for vaccination at a particular point in time.  COVID-19 vaccine should be offered to the following individuals:   * Residents in a care home for older adults and their carers * All those 80 years of age and over * Frontline health and social care workers (as included in COVID-19 –SARS-Cov-2 chapter of Green Book, JCVI statement and Scottish Government CMO letters) * All those 75 years of age and over * All those 70 years of age and over * Clinically extremely vulnerable (CEV) individuals (not including all pregnant women and those under 16 years) as defined by Scottish Government at <https://www.gov.scot/publications/covid-shielding/pages/highest-risk-classification/> * All those 65 years of age and over * Individuals aged 16 years to 64 years with underlying health conditions which puts them at higher risk of serious disease and mortality included in Table 3 COVID-19 –SARS-Cov-2 chapter 14a of Green Book\* * All those 60 years of age and over * All those 55 years of age and over * All those 50 years of age and over * Further guidance for those under age 50 years not included in the above groups will follow in phase 2.   \*This also includes those who are in receipt of a carer’s allowance, or those who are the main carer of an elderly or disabled person whose welfare may be at risk if the carer falls ill.  The list above is not exhaustive, and clinician should apply clinical judgment to take into account the risk of COVID-19 exacerbating any underlying disease that a patient may have, as well as the risk of serious illness from COVID-19 itself. COVID-19 vaccine should be offered in such cases even if the individual is not in the clinical risk groups specified above, this may be provided under a Patient Specific Direction (PSD). |
| **Exclusion criteria** | The vaccine should not be given to:   * Those who have had a confirmed anaphylactic reaction to a previous dose of this COVID-19 vaccine * Those who have had a confirmed anaphylactic reaction to any components of this vaccine * Those with a history of immediate-onset anaphylaxis to multiple classes of drugs or unexplained anaphylaxis. * Those in whom no valid consent has been received * Those who are under 16 years of age * Women who are known to be pregnant (routine questioning about last menstrual period and/or pregnancy testing is not required before offering the vaccine) * Those with confirmed COVID-19 infection to avoid confusing the differential diagnosis. As clinical deterioration can occur up to two weeks after infection, ideally vaccination should be deferred until around four weeks after onset of symptoms or from the first PCR positive specimen in those who are asymptomatic. * Those with evidence of current deterioration of COVID-19 symptoms, deferral of vaccination may be considered to avoid incorrect attribution of any change in the person’s underlying condition to the vaccine. * Those who are participating in a clinical trial of COVID-19 vaccines * Those with acute febrile illness – consider postponing immunisation until individual has fully recovered. * Those with evolving neurological condition – consider postponing immunisation until individual has stabilised. |
| **Cautions/need for further advice/ circumstances when further advice should be sought from a doctor** | The COVID-19 chapter of the Green Book advises that there are very few individuals who cannot receive COVID vaccine. Where there is doubt, rather than withholding vaccination, appropriate advice should be sought from the relevant specialist, or from the local immunisation or health protection team.  Individuals with a bleeding disorder may develop a haematoma at the injection site (see Route of Administration).  The MHRA has advised that individuals with a history of anaphylaxis to food, an identified drug or vaccine, or an insect sting can receive any COVID-19 vaccine, as long as they are not known to be allergic to any component of the vaccine.  Because of the absence of data on co-administration with COVID-19 vaccines, it should not be routine to offer appointments to give this vaccine at the same time as other vaccines. Based on current information about the first COVID-19 vaccines being deployed, scheduling should ideally be separated by an interval of at least 7 days to avoid incorrect attribution of potential adverse events.  As COVID-19 mRNA Vaccine BNT162b2 is considered inactivated, where individuals in an eligible cohort present having received another inactivated or live vaccine, COVID-19 vaccination should still be considered. The same applies for other live and inactivated vaccines where COVID-19 vaccination has been received first. In many cases, vaccination should proceed to avoid any further delay in protection and to avoid the risk of the patient not returning for a later appointment. In such circumstances, patients should be informed about the likely timing of potential adverse events relating to each vaccine.  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.  JCVI advise there is no known risk associated with giving these types of vaccines during pregnancy. These vaccines cannot replicate, so they cannot cause infection in either the woman or the unborn child.  Although the available data do not indicate any safety concern or harm to pregnancy, there is insufficient evidence to recommend routine use of COVID-19 vaccines during pregnancy.  JCVI advises that, for women who are offered vaccination with the Pfizer-BioNTech or AstraZeneca COVID-19 vaccines, vaccination in pregnancy should be considered where the risk of exposure to Severe Acute Respiratory Syndrome coronavirus 2 (SARS-CoV2) infection is high and cannot be avoided, or where the woman has underlying conditions that put them at very high risk of serious complications of COVID-19. In these circumstances, clinicians should discuss the risks and benefits of vaccination with the woman, who should be told about the absence of safety data for the vaccine in pregnant women.  There is no known risk associated with giving non-live vaccines whilst breastfeeding. JCVI advises that breastfeeding women may be offered vaccination with the Pfizer-BioNTech or AstraZeneca COVID-19 vaccines.  The developmental and health benefits of breastfeeding should be considered along with the woman’s clinical need for immunisation against COVID-19, and the woman should be informed about the absence of safety data for the vaccine in breastfeeding women. |
| **Action if excluded** | Specialist advice should be sought on the vaccine and circumstances under which it could be given as vaccination using a patient specific direction may be indicated.  Individuals who are participating in a clinical trial of COVID-19 vaccines who present for vaccination should be referred back to the investigators.  In case of postponement due to acute illness or evolving neurological condition, advise when the individual can be vaccinated and ensure another appointment is arranged.  In case of postponement due to COVID-19 symptoms or positive COVID test in the last four weeks advise when the individual can be vaccinated and how future vaccination may be accessed.  Document the reason for exclusion and any action taken in accordance with local procedures. |
| **Action if patient declines** | Advise the individual/carer about the protective effects of the vaccine, the risks of infection and potential complications if not immunised.  Advise how future immunisation may be accessed if they subsequently decide to receive the COVID-19 vaccine  Document patient’s declined consent and advice given. |

**2. Description of treatment**

| Category | Description |
| --- | --- |
| **Name of medicine** | COVID-19 mRNA Vaccine BNT162b2 concentrate for solution for injection |
| **Form/strength** | COVID-19 mRNA Vaccine BNT162b2 30micrograms/0.3ml dose concentrate for solution for injection multidose vials  COVID-19 mRNA Vaccine BNT162b2 is a multidose vial and must be diluted with 1.8mL of 0.9% sodium chloride before use. 1 vial contains 5 doses of 30 micrograms of BNT162b2 RNA (embedded in lipid nanoparticles). |
| **Route of** **administration** | After dilution, the vial contains 5 doses of 0.3 ml. Withdraw the required 0.3 mL dose of diluted vaccine using a sterile needle and syringe and administer. Vial volume was optimized to reliably obtain 5 doses regardless of syringe type used as most syringe and needle combinations require withdrawal of excess volume in order to ensure the full 0.3 mL dose of vaccine can be administered.  When low dead-volume syringes and/or needles are used, the amount remaining in the vial after 5 doses have been extracted may be sufficient for an additional (sixth) dose. Care should be taken to ensure a full 0.3 mL will be administered to the subject and that all doses from a single prepared vial are administered within 6 hours of the time of dilution. Where a full 0.3 mL dose cannot be extracted the contents should be discarded. Any unused vaccine should be discarded 6 hours after dilution.  COVID-19 mRNA Vaccine BNT162b2 must be administered by intramuscular (IM) injection preferably into the deltoid area of the upper arm. Where administration into the deltoid is not possible the anterolateral thigh can be considered.  Inspect visually prior to administration and ensure appearance is consistent with the description in the manufacturer’s product literature or summary of product characteristics.  Individuals with bleeding disorders may be vaccinated intramuscularly if, in the opinion of a doctor familiar with individual’s bleeding risk, vaccines or similar small volume intramuscular injections can be administered with reasonable safety by this route. If the individual receives medication/ treatment to reduce bleeding, for example treatment for haemophilia, intramuscular vaccination can be scheduled shortly after such medication/treatment is administered. Individuals on stable anticoagulation therapy, including individuals on warfarin who are up-to-date with their scheduled INR testing and whose latest INR is below the upper level of the therapeutic range, can receive intramuscular vaccination. A fine needle (23 or 25 gauge) should be used for the vaccination, followed by firm pressure applied to the site without rubbing for at least 2 minutes. The individual/parent/carer should be informed about the risk of haematoma from the injection.  The site at which each vaccine was given should be noted in the individual’s records. |
| **Dosage** | The dose of COVID-19 mRNA Vaccine BNT162b2 is 30 micrograms contained in 0.3mL of the diluted vaccine. |
| **Frequency** | COVID-19 mRNA Vaccine BNT162b2 course consists of two separate doses of 0.3ml each, a minimum of 21 days apart.  Operationally, it is recommended in the COVID-19 chapter of Green Book that the second dose of both vaccines should be routinely scheduled between four and 12 weeks after the first dose. This will allow more people to benefit from the protection provided from the first dose during the roll out phase. Longer term protection will then be provided by the second dose.  If an interval longer than the recommended interval is left between doses, the second dose should still be given (preferably using the same vaccine as was given for the first dose if possible). The course does not need to be restarted.  JCVI advises that the second vaccine dose should be with the same vaccine as for the first dose. Switching between vaccines or missing the second dose is not advised as this may affect the duration of protection.  There is no evidence on the interchangeability of the COVID-19 vaccines although studies are underway. Therefore, every effort should be made to determine which vaccine the individual received and to complete with the same vaccine. For individuals who started the schedule and who attend for vaccination at a site where the same vaccine is not available, or if the first product received is unknown, it is reasonable to offer a single dose of the locally available product. This option is preferred if the individual is likely to be at immediate high risk or is considered unlikely to attend again. In these circumstances, as both the vaccines are based on the spike protein, it is likely the second dose will help to boost the response to the first dose. For this reason, until additional information becomes available, further doses are not required. |
| **Duration of treatment** | See Dose and frequency of administration above.  Booster doses of COVID-19 vaccine are not yet recommended because the need for, and timing of, boosters has not yet been determined. |
| **Maximum or minimum treatment period** | See Frequency of administration above. |
| **Quantity to supply/administer** | Administer 30 micrograms in 0.3mL per administration. |
| **▼ black triangle medicines** | COVID-19 mRNA Vaccine BNT162b2 did not have a UK marketing authorisation at the time this protocol was written and is authorised for temporary supply in the UK in accordance with a Regulation 174 authorisation.  All adverse reactions occurring in individuals of any age after vaccination should be reported to the MHRA using the Coronavirus Yellow Card Scheme. Anyone can report a suspected adverse reaction to the MHRA using the Coronavirus Yellow Card reporting scheme  <https://coronavirus-yellowcard.mhra.gov.uk/> |
| **Legal category** | COVID-19 mRNA Vaccine BNT162b2 is provided temporary authorisation by the Medicines & Healthcare products Regulatory Agency (MHRA) for supply in the UK under regulation 174 and 174A, pending UK marketing authorisation.  The regulation 174 authorised product is categorised as a prescription only medicine (POM). |
| **Is the use out with the SPC?** | COVID-19 mRNA Vaccine BNT162b2 is supplied in the UK in accordance with regulation 174 and did not have a UK marketing authorisation at the time of writing this protocol.  As part of the consent process, inform the individual/carer that this vaccine does not have a UK marketing authorisation but has been authorised for temporary supply in the UK by the MHRA and that it is being offered in accordance with national guidance.  The vaccine manufacturer’s information for UK healthcare professionals states that the vaccine should be given as a series of two doses (0.3mL, each) 21 days apart. This is superseded by the JCVI advice that the second dose of both vaccines should be routinely scheduled between four and 12 weeks after the first dose. This will allow more people to benefit from the protection provided from the first dose during the roll out phase. Longer term protection will then be provided by the second dose. |
| **Storage requirements** | COVID-19 mRNA Vaccine BNT162b2 must be stored frozen at ultra-low temperature in accordance with manufacturer’s advice.  Once removed from the freezer BNT162b2 vaccine can be stored for 5 days in a fridge between +2 to +8°C prior to dilution.  NHS Board guidance on Storage and Handling of vaccines should be observed.  COVID-19 mRNA Vaccine BNT162b2 should be diluted as close to use as possible. However, reconstituted vaccine which is not required immediately must be used within 6 hours from the time of dilution and stored between +2°C to +25°C.  The vaccine vial has space to write the date and time of dilution; write this on the vial label.  During storage, minimise exposure to room light and avoid exposure to direct sunlight and ultraviolet light.  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 use or appropriate disposal.  The manufacturer may advise of updated storage requirements and product stability as new data becomes available, vaccine may be stored in accordance with updated recommendations from the manufacturer. |
| **Additional information** | Minor illnesses without fever or systemic upset are not valid reasons to postpone immunisation. If an individual is acutely unwell, immunisation should be postponed until they have fully recovered.  There is no evidence of any safety concerns from vaccinating individuals with a past history of COVID-19 infection, or with detectable COVID-19 antibody. Inclusion of antibody positive individuals in the Pfizer phase 3 analysis did not give any safety signals.  Vaccination of individuals who may be infected but asymptomatic or incubating COVID-19 infection is unlikely to have a detrimental effect on the illness. Vaccination should be deferred in those with confirmed infection to avoid confusing the differential diagnosis. As clinical deterioration can occur up to two weeks after infection, ideally vaccination should be deferred until around four weeks after onset of symptoms or from the first PCR positive specimen in those who are asymptomatic.  Having prolonged COVID-19 symptoms is not a contraindication to receiving COVID-19 vaccine but if the patient is seriously debilitated, still under active investigation, or has evidence of recent deterioration, deferral of vaccination may be considered to avoid incorrect attribution of any change in the person’s underlying condition to the vaccine. |

**3. Adverse reactions**

| Category | Description |
| --- | --- |
| **Warnings including possible adverse reactions and management of these** | Local reactions at the injection site are fairly common after COVID-19 mRNA Vaccine BNT162b2 primarily pain at the injection site, usually without redness and swelling. Systemic events reported were generally mild and short lived. In the final safety analysis of over 21,000 participants 16 years and older, the most common events were injection site pain (>80%), fatigue (>60%), and headache (>50%). Myalgia, arthralgia and chills were also common with fever in 10-20%. Most were classified as mild or moderate. Lymphadenopathy was reported in less than 1%.  In an earlier analysis of around 8,000 recipients, severe side effects, defined as those that interfere with daily activity, included fatigue in around 4% and headache in 2%. Older adults tend to report fewer adverse events following vaccination. This earlier analysis also showed no signal for enhanced disease in vaccine recipients with only 1 case of severe COVID in the 8 vaccine failures at that time.  A protocol for the management of anaphylaxis and an anaphylaxis pack must always be available whenever COVID-19 mRNA Vaccine BNT162b2 is given. Immediate treatment should include early treatment with 0.5mg intramuscular adrenaline (0.5ml of 1:1000 or 1mg/ml 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.  In the event of a severe adverse reaction individual should be advised to seek medical advice.  For full details/information on possible adverse reaction, refer to manufacturer’s product literature or summary of product characteristics. |
| **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 Coronavirus Yellow Card reporting scheme on:<https://coronavirus-yellowcard.mhra.gov.uk/>  As this vaccine is labelled with a black triangle, all adverse reactions occurring in individuals of any age after vaccination should be reported to the MHRA using the Coronavirus Yellow Card Scheme. Anyone can report a suspected adverse reaction to the MHRA using the Coronavirus Yellow Card reporting scheme <https://coronavirus-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.  Anaphylaxis is a very rare, recognised side effect of most vaccines and suspected cases should be reported via the Coronavirus Yellow Card Scheme. Chapter 8 of the Green Book gives detailed guidance on distinguishing between faints, panic attacks and the signs and symptoms of anaphylaxis. If a case of suspected anaphylaxis meets the clinical features described in Chapter 8, this should be reported via the Yellow Card Scheme as a case of ‘anaphylaxis’ (or if appropriate ‘anaphylactoid reaction’). Cases of less severe allergic reactions (i.e. not including the clinical features of anaphylaxis) should not be reported as anaphylaxis but as ‘allergic reaction’.  Programmatic Adverse Events should be recorded in line with local procedures and where appropriate escalated in accordance with the national framework. |
| **Advice to patient or carer including written information** | Written information to be given to individual   * Provide manufacturer’s consumer information leaflet/patient information leaflet (PIL) provided with the vaccine. * Provide copy of Public Health Scotland post- vaccination leaflet * Provide copy of Pregnant, planning a pregnancy or breastfeeding, a guide to COVID-19 vaccine to women of child bearing years   Individual advice / follow up treatment   * Inform the individual/carer of possible side effects and their management. * Vaccinated individuals should be advised that it is common to develop a fever after vaccination and that this normally happens within 48 hours after the vaccination and usually goes away within 48 hours. This is a common, expected reaction, and self-isolation and testing for COVID-19 are not required unless the individual has other COVID-19 symptoms; has been told by NHS Test and Protect they are a close contact of someone who has tested positive for COVID-19; they live with someone who has recently tested positive for COVID-19; or they live with someone who has symptoms of COVID-19. * Vaccinated individuals should be advised that if the fever started 48 hours after the vaccination or lasts longer than 48 hours, they should self-isolate and book a test. * Vaccinated individuals should be advised that feeling generally unwell, shivery, achy and tired were also symptoms commonly reported by vaccine recipients in the clinical trials. Generally, these symptoms were found to resolve within one to two days without treatment but paracetamol can be taken if necessary to relieve any of these symptoms. * As has always been recommended, any fever after vaccination should be monitored and if individuals are concerned about their health at any time, they should seek advice from their GP or NHS24 * 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 to the MHRA using the Yellow Card reporting scheme on: http://yellowcard.mhra.gov.uk. * Immunosuppressed individuals should be advised that they may not make a full immune response to the vaccine and they should continue to take appropriate measures to protect themselves against this infection. * When administration is postponed advise the individual how future vaccination may be accessed * When applicable, advise the individual/carer when to return for vaccination or when a subsequent vaccine dose is due. |
| **Observation following vaccination** | Vaccine recipients should be monitored for 15 minutes after vaccination, with a longer observation period when indicated after clinical assessment.  As syncope (fainting) can occur following vaccination, all vaccinees should either be driven by someone else or should not drive for 15 minutes after vaccination. |
| **Follow up** | Not applicable |
| **Additional facilities** | A protocol for the management of anaphylaxis and an anaphylaxis pack must always be available whenever COVID-19 mRNA Vaccine BNT162b2 is given. Immediate treatment should include early treatment with 0.5mg intramuscular adrenaline (0.5ml of 1:1000 or 1mg/ml 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. |

**4. Audit Trail/Records**

| 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 protocol   Records should kept line with local procedures. Ideally records should be kept within the NHS Scotland COVID-19 vaccine administration app.  Local policy should be followed to encourage information sharing with the individual’s General Practice.  All records should be clear, legible and contemporaneous. |

**5. References**

| Name | Description |
| --- | --- |
| **Additional references** | Immunisation against Infectious Disease [Green Book] <https://www.gov.uk/government/organisations/public-health-england/series/immunisation-against-infectious-disease-the-green-book>  Immunisation against Infectious Disease [Green Book] COVID-19 <https://www.gov.uk/government/publications/covid-19-the-green-book-chapter-14a>  JCVI: advice on priority groups for COVID-19 vaccine 30th December 2020  <https://www.gov.uk/government/publications/priority-groups-for-coronavirus-covid-19-vaccination-advice-from-the-jcvi-30-december-2020>  Manufacturer’s product information/ Summary of Product Characteristics <https://www.gov.uk/government/publications/regulatory-approval-of-pfizer-biontech-vaccine-for-covid-19>  Educational resources for registered professionals produced by National Education for Scotland  <https://learn.nes.nhs.scot/37676/immunisation/covid-19-vaccines>  All relevant Scottish Government advice including the relevant CMO letter(s) |

**ANNEX B: Practitioner authorisation sheet**

**COVID-19 mRNA Vaccine BNT162b2 Pfizer/BioNTech Vaccine Protocol v00.01 Valid from: DDDDDD Expiry: DDDDDD**

Before signing this Protocol, check that the document has had the necessary authorisations in section 1 and 2. Without these, this Protocol is not lawfully valid.

**Practitioner**

By signing this Protocol you are indicating that you agree to its contents and that you will work within it.

Protocols do not remove inherent professional obligations or accountability.

It is the responsibility of each practitioner to practise only within the bounds of their own competence and any appropriate professional code of conduct.

|  |  |  |  |
| --- | --- | --- | --- |
| I confirm that I have read and understood the content of this Protocol and that I am willing and competent to work to it within my professional code of conduct. | | | |
| Name | Designation | Signature | Date |
|  |  |  |  |
|  |  |  |  |
|  |  |  |  |
|  |  |  |  |
|  |  |  |  |

**Person authorising on behalf of Provider**

|  |  |  |  |
| --- | --- | --- | --- |
| I confirm that the practitioners named above have declared themselves suitably trained and competent to work under this Protocol. I give authorisation on behalf of **insert name of organisation**  for the above named health care professionals who have signed the Protocol to work under it. | | | |
| Name | Designation | Signature | Date |
|  |  |  |  |

**Note to person authorising on behalf of Provider**

Score through unused rows in the list of practitioners to prevent practitioner additions post managerial authorisation.

This authorisation sheet should be retained to serve as a record of those practitioners authorised to work under this Protocol.

**ANNEX C: Clinical Supervision sheet**

**COVID-19 mRNA Vaccine BNT162b2 Pfizer/BioNTech Vaccine Protocol v00.01 Valid from: DDDDDD Expiry: DDDDDD**

This sheet must record the name of the clinical supervisor taking responsibility and all of the people working under different activity stages of the protocol.

Activity stages of the vaccination pathway under this protocol:

|  |  |  |
| --- | --- | --- |
| Stage 1 | 1. Assessment of the individual presenting for vaccination 2. Provide information and obtain informed consent 3. Provide advice to the individual | Registered Healthcare Professionals Only |
| Stage 2 | * Vaccine Preparation | Registered or non-registered persons |
| Stage 3 | * Vaccine Administration | Registered or non-registered persons |
| Stage 4 | * Record Keeping | Registered or non-registered persons |

The clinical supervisor has ultimate responsibility for safe care being provided under the terms of the protocol. Persons working under the protocol may be supported by additional registered healthcare professionals, but the clinical supervisor retains responsibility.

Before signing this Protocol, check that the document has had the necessary authorisations. Without these, this Protocol is not lawfully valid.

**Clinical Supervisor**

|  |  |  |  |
| --- | --- | --- | --- |
| Name | Designation | Signature | Date |
|  |  |  |  |

**Practitioner(s) and Activity Stages**

|  |  |  |  |
| --- | --- | --- | --- |
| Name | Activity Stage(s) | Signature | Date |
|  |  |  |  |
|  |  |  |  |
|  |  |  |  |

**Note to Clinical Supervisor**

Score through unused rows in the list of practitioners to prevent practitioner additions post managerial authorisation.

This authorisation sheet should be retained to serve as a record of clinical supervision arrangements for those working under this Protocol.