IMPORTANT: It is important that you carry this card with you at all times.

Please ensure you show this card to all Healthcare Professionals (including nurses, pharmacists and

dentists), to any doctor involved in your treatment, and at any visits to the hospital.

Name of oncologist:___ Contact number:

After-hours contact number:

My name:

My contact number: Emergency contact:

> Important Information for **Health Care Providers**

(atezolizumab), which can cause immune-related adverse reactions that involve the lungs, liver, intestines, hormone glands, heart, and other organs, as well as infusionrelated reactions. Early diagnosis and appropriate management are essential to minimise any consequences

For suspected immune-related adverse reactions, ensure adequate evaluation to confirm aetiology or exclude other cause. Based on the severity of the adverse reaction, withhold Tecentriq® and administer corticosteroids. Specific guidelines for managing immune-related adverse reactions are provided in the Summary of Product Characteristics for atezolizumab, available at www.medicines.ie and the Physician Education Materials, available for download via the Medicines Authority of Malta website (http://www.

Please contact the patient's Oncologist (details on this

Assess patients for signs and symptoms of pneumonitis,

Please consult the Summary of Product Characteristics

Reporting of suspected adverse events

Reporting suspected adverse events or reactions after authorisation of the medicinal product is important. It allows continued monitoring of the benefit/ risk balance of the medicinal product. Healthcare professionals are asked to report any suspected adverse events or reactions (see details below). In the event of a suspected adverse event,

Roche Products (Ireland) Limited, 3004 Lake Drive,

Tel: 00 353 (0)1 4690700; Fax: 00 353 (0)1 4690793 Email: ireland.drug_surveillance_centre@roche.com Alternatively, suspected adverse reactions

Medicines Authority, Sir Temi Żammit Buildings, Malta Life Sciences Park, San Gwann SGN 3000, Malta Reporting forms and information can be found at www.medicinesauthority.gov.mt/adrportal

For additional copies of this risk minimisation material, refer to the Medicines Authority of Malta website (http://www.medicinesauthority.gov.mt/rmm) and download the required material or alternatively if you would like hard copies, please contact Roche Products (Ireland) Limited, 3004 Lake Drive, Citywest, Naas Road, Dublin 24 by mail, telephone (00 353 (0)1 4690700), fax (00 353 (0)1 4690791)

For further information about Tecentriq®, please contact Medical Information at Roche Products (Ireland) Limited by telephone (00 353 (0)1 4690700),

hepatitis, colitis, endocrinopathies (including hypophysitis, adrenal insufficiency, type 1 diabetes mellitus, hypothyroidism, hyperthyroidism), myocarditis, pancreatitis, nephritis and infusion-related reactions. Other immune-related adverse reactions reported in patients receiving atezolizumab include: neuropathies (Guillain-Barré syndrome, myasthenic syndrome/ myasthenia gravis), and meningoencephalitis.

for Tecentriq® available at www.medicines.ie.

▼ This medicinal product is subject to additional monitoring. This will allow quick identification of new

This patient is being treated with Tecentriq®

of immune-related adverse reactions.

medicinesauthority.gov.mt/rmm).

card) for more information.

safety information

or reactions

please report it to:

The Drug Surveillance Centre,

Citywest, Naas Road, Dublin 24.

should be reported to:

Further information

or email (Ireland.dra@roche.com).

fax (00 353 (0)1 4690791) or email (Ireland.druginfo@roche.com).

Roche

MT Version 3.0.0

safety information

Roche>

Emergency contact number: _

All rights reserved. Tecentrig®▼ (atezolizumab) ▼ This medicinal product is subject to additional monitoring. This will allow quick identification of new

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Patient Alert Card

Date of Malta Medicines Authority Approval: March 2019 MT Version 3.0.0

Symptoms may occur at any time during treatment or even after your treatment has ended. Call your doctor right away if you develop any of these new signs or symptoms listed on this card or if your symptoms should get worse. Also tell your doctor if you experience any other symptoms not listed on this card.

Tecentriq® can cause serious side effects in many parts of your body that need to be treated right away.

IMPORTANT:

Carry this card with you at all times, especially when you travel, whenever you go to the Accident and Emergency department, or when you see another doctor.

Do not try to treat your symptoms on your own.

IMPORTANT Reminders for Patients Like all medicines, Tecentriq® (atezolizumab) may cause side effects, although not everybody gets them. It is important to tell your doctor immediately if you develop any of the signs or symptoms listed on this card after starting treatment with atezolizumab. Before you start atezolizumab or during your treatment, you should also tell your doctor immediately if you:

Have an autoimmune disease (a condition

where the body attacks its own cells, examples include autoimmune thyroid disease, systemic lupus erythematosus (SLE), Sjogren's syndrome, multiple sclerosis, rheumatoid arthritis, vasculitis,

glomerulonephritis)

Have been told that your cancer has spread to your brain Have any history of inflammation of your lungs

(pneumonitis)

Have or have had chronic viral infection of the liver, including hepatitis B (HBV) or hepatitis C (HCV)

immune deficiency syndrome (AIDS)

· Have human immunodeficiency virus (HIV) or acquired Experienced serious side effects because of other

antibody therapies that help your immune system to fight cancer

Have been given medicines to stimulate your immune system such as interferons or interleukin-2 as these medicines may worsen the side effects of atezolizumab Have been given medicines to suppress your immune

system such as corticosteroids, since these medicines

may interfere with the effect of atezolizumab · Have been given a live, attenuated vaccine such as influenza intranasal vaccine, yellow fever vaccine

You should not start any other medicines during your treatment without talking to your doctor first.

If you develop any signs or symptoms listed on this card or if you notice any signs or symptoms not listed on this

card, please contact your doctor immediately. Getting medical treatment early may stop the problem from

becoming more serious.

If you have any further questions about your treatment or on the use of this medicine, please contact your doctor.

Reporting of side effects

Tel: 00 353 (0)1 4690700; Fax: 00 353 (0)1 4690793 Email: ireland.drug_surveillance_centre@roche.com

Medicines Authority, Sir Temi Żammit Buildings, Malta Life Sciences Park, San Gwann SGN 3000, Malta Reporting forms and information can be found at www.medicinesauthority.gov.mt/adrportal

Select important safety information

events may result in signs or symptoms such as:

chest pain.

stomach pain.

blood in stools, stomach pain.

increased sensitivity to cold or heat.

tingling in your hands and feet.

pelvis and swelling of the body.

breath, flushing.

decreased exercise tolerance, ankle swelling.

Serious side effects may include lung problems (pneumonitis), liver problems (hepatitis), intestinal problems (colitis), problems in hormone glands (for example thyroid problems or diabetes), heart, nervous system, and other organs. These

Lungs: new or worsening cough, shortness of breath,

Liver: yellowing of skin or the whites of eyes, severe nausea or vomiting, bleeding or bruising, dark urine,

Intestines: diarrhoea (watery, loose or soft stools),

Hormone glands: extreme tiredness, persistent headache, weight loss, weight gain, change in mood, hair loss, constipation, dizziness, feeling more hungry or thirsty than usual, need to urinate more often,

Heart: chest pain, shortness of breath, irregular heart beat,

 Brain: neck stiffness, headache, fever, chills, vomiting, eye sensitivity to light, confusion, sleepiness.

Nerves: severe muscle weakness and numbness,

Kidneys: changes in urine output and colour, pain in

Reactions associated with infusion (during or within 1 day of infusion): fever, chills, shortness of

Getting medical treatment immediately may stop the problems from becoming serious. Your doctor may decide to give you other medicines to prevent complications and reduce your symptoms, and may withhold the next dose or stop your treatment.

Pancreas: abdominal pain, nausea, vomiting.

If you get any side effects, talk to your doctor, pharmacist or nurse. This includes any possible side

effects not listed in the Tecentriq® Package Leaflet.

You can also report side effects directly (see details below). By reporting side effects you can help provide

Citywest, Naas Road, Dublin 24.

Alternatively, report to:

more information on the safety of this medicine. Please report it to:

The Drug Surveillance Centre, Roche Products (Ireland) Limited, 3004 Lake Drive,