ANNEX I SUMMARY OF PRODUCT CHARACTERISTICS

1. NAME OF THE VETERINARY MEDICINAL PRODUCT

Versican Plus Pi lyophilisate and solvent for suspension for injection for dogs

2. QUALITATIVE AND QUANTITATIVE COMPOSITION

Each dose of 1 ml contains:

Active substances:

	Minimum	Maximum
Lyophilisate (live attenuated):		
Canine parainfluenza Type 2 virus, strain CPiV-2 Bio 15	$10^{3.1} \text{ TCID}_{50}*$	$10^{5.1} \text{ TCID}_{50}*$

^{*} Tissue culture infectious dose 50%.

Excipients:

Qualitative composition of excipients and	
other constituents	
Lyophilisate:	
Trometamol	
Edetic Acid	
Sucrose	
Dextran 70	
Solvent:	
Water for injections (Aqua ad iniectabilia)	

The visual appearance is as follows:

Lyophilisate: spongy matter of white colour.

Solvent: clear colourless liquid.

3. CLINICAL INFORMATION

3.1 Target species

Dogs.

3.2 Indications for use for each target species

Active immunisation of dogs from 6 weeks of age:

 to prevent clinical signs (nasal and ocular discharge) and reduce viral excretion caused by canine parainfluenza virus.

Onset of immunity:

3 weeks after completion of the primary course.

Duration of immunity:

At least one year following the primary vaccination course.

3.3 Contraindications

None.

3.4 Special warnings

A good immune response is reliant on a fully competent immune system. Immunocompetence of the animal may be compromised by a variety of factors including poor health, nutritional status, genetic factors, concurrent medicinal therapy and stress.

Vaccinate healthy animals only.

3.5 Special precautions for use

Special precautions for safe use in the target species:

The live attenuated virus vaccine strain CPiV may be shed by vaccinated dogs following vaccination. However, due to the low pathogenicity of this strain, it is not necessary to keep vaccinated dogs separated from non-vaccinated dogs.

Special precautions to be taken by the person administering the veterinary medicinal product to animals:

In case of accidental self-injection, seek medical advice immediately and show the package leaflet or the label to the physician.

Special precautions for the protection of the environment:

Not applicable.

3.6 Adverse events

Dogs:

Common	injection site swelling ¹
(1 to 10 animals / 100 animals treated):	
Rare	hypersensitivity reaction ² (anaphylaxis, angioedema,
(1 to 10 animals / 10,000 animals	circulatory shock, collapse, diarrhoea, dyspnoea,
treated):	vomiting)
	anorexia, decreased activity
Very rare	hyperthermia, lethargy, malaise
(<1 animal / 10,000 animals treated,	
including isolated reports):	

¹A transient swelling (up to 5 cm) which can be painful, warm or reddened. Any such swelling will either have spontaneously resolved or be greatly diminished by 14 days after vaccination.

Reporting adverse events is important. It allows continuous safety monitoring of a veterinary medicinal product. Reports should be sent, preferably via a veterinarian, to either the marketing authorisation holder or the national competent authority via the national reporting system. See the package leaflet for respective contact details.

3.7 Use during pregnancy, lactation or lay

Pregnancy and lactation:

Can be used during the second and third stages of pregnancy. Safety of the product during the early stage of pregnancy and during lactation has not been investigated.

²If a hypersensitivity reaction occurs, appropriate treatment should be administered without delay. Such reactions may evolve to a more severe condition which may be life-threatening.

3.8 Interaction with other medicinal products and other forms of interaction

No information is available on the safety and efficacy of this vaccine when used with any other veterinary medicinal product other than Versiguard Rabies and Versican Plus L4. A decision to use this vaccine before or after any other veterinary medicinal product therefore needs to be made on a case by case basis.

Leptospira:

If protection against *Leptospira* is required, dogs can be vaccinated with two doses of Versican Plus Pi mixed with Versican Plus L4 3–4 weeks apart from 6 weeks of age:

The contents of a single vial of Versican Plus Pi should be reconstituted with the contents of a single vial of Versican Plus L4 (instead of the solvent). Once mixed, the contents of the vial should appear a whitish to yellowish colour with light opalescence. The mixed vaccines should be injected immediately via the subcutaneous route.

Rabies:

If protection against rabies is required:

First dose: Versican Plus Pi from 8–9 weeks of age.

Second dose: Versican Plus Pi mixed with Versiguard Rabies 3–4 weeks later, but not before 12 weeks of age.

The contents of a single vial of Versican Plus Pi should be reconstituted with the contents of a single vial of Versiguard Rabies (instead of the solvent). Once mixed, the contents of the vial should appear a pink/red or yellowish colour with light opalescence. The mixed vaccines should be injected immediately via the subcutaneous route.

The efficacy of the rabies fraction is proven after a single dose from 12 weeks of age in laboratory studies. However, in field studies 10% of seronegative dogs did not show seroconversion (>0.1 IU/ml) 3–4 weeks after single primary vaccination against rabies. Some animals may also not show titres > 0.5 IU/ml after the primary vaccination. Antibody titres drop over the course of the 3-year duration of immunity, although dogs are protected when challenged. In case of travelling to risk areas or outside the EU, veterinary surgeons may wish to give additional rabies vaccinations after 12 weeks of age to ensure that the vaccinated dogs have an antibody titre of \geq 0.5 IU/ml, which is generally regarded as sufficiently protective and that they meet the travel test requirements (antibody titres \geq 0.5 IU/ml).

Although the efficacy of the rabies fraction has been demonstrated following administration at 12 weeks, at the discretion of the veterinary surgeon, in case of need, dogs younger than 8 weeks can be vaccinated with Versican Plus Pi mixed with Versiguard Rabies as the safety of this association has been demonstrated in 6-week old dogs.

3.9 Administration routes and dosage

Subcutaneous use.

Dosage and route of administration:

Aseptically reconstitute the lyophilisate with the solvent. Shake well and administer immediately the entire contents (1 ml) of the reconstituted product.

Appearance of the reconstituted vaccine: clear whitish to yellowish colour with light opalescence.

Primary vaccination scheme:

Two doses of Versican Plus Pi 3-4 weeks apart from 6 weeks of age.

Re-vaccination scheme:

A single dose of Versican Plus Pi to be given annually.

3.10 Symptoms of overdose (and where applicable, emergency procedures and antidotes)

No adverse events other than those mentioned in section 3.6 were observed after administration of a 10-fold overdose of the vaccine. However, in a minority of animals pain was observed at the injection site immediately after administration of a 10-fold overdose of the vaccine.

3.11 Special restrictions for use and special conditions for use, including restrictions on the use of antimicrobial and antiparasitic veterinary medicinal products in order to limit the risk of development of resistance

Not applicable.

3.12 Withdrawal periods

Not applicable.

4. IMMUNOLOGICAL INFORMATION

4.1 ATCvet code: QI07AD08

The vaccine is intended for the active immunisation of healthy puppies and dogs against disease caused by canine parainfluenza virus.

5. PHARMACEUTICAL PARTICULARS

5.1 Major incompatibilities

Do not mix with any other veterinary medicinal product except those mentioned in section 3.8 above.

5.2 Shelf life

Shelf life of the veterinary medicinal product as packaged for sale: 2 years. Shelf life after reconstitution according to directions: use immediately.

5.3 Special precautions for storage

Store and transport refrigerated (2 $^{\circ}C$ – 8 $^{\circ}C$). Do not freeze.

Protect from light.

5.4 Nature and composition of immediate packaging

Type I glass vial containing 1 dose of lyophilisate closed with a bromobutyl rubber stopper and aluminium cap.

Type I glass vial containing 1 ml of solvent closed with a chlorobutyl rubber stopper and aluminium cap.

Pack sizes:

Plastic box containing 25 vials (1 dose) of lyophilisate and 25 vials (1 ml) of solvent.

Plastic box containing 50 vials (1 dose) of lyophilisate and 50 vials (1 ml) of solvent.

Not all pack sizes may be marketed.

5.5 Special precautions for the disposal of unused veterinary medicinal products or waste materials derived from the use of such products

Medicines should not be disposed of via wastewater or household waste.

Use take-back schemes for the disposal of any unused veterinary medicinal product or waste materials derived thereof in accordance with local requirements and with any national collection systems applicable to the veterinary medicinal product concerned.

6. NAME OF THE MARKETING AUTHORISATION HOLDER

Zoetis Belgium

7. MARKETING AUTHORISATION NUMBER(S)

EU/2/14/168/001 EU/2/14/168/002

8. DATE OF FIRST AUTHORISATION

Date of first authorisation: 04/07/2014.

9. DATE OF THE LAST REVISION OF THE SUMMARY OF THE PRODUCT CHARACTERISTICS

10. CLASSIFICATION OF VETERINARY MEDICINAL PRODUCTS

Veterinary medicinal product subject to prescription.

Detailed information on this veterinary medicinal product is available in the Union Product Database (https://medicines.health.europa.eu/veterinary).

	ANNEX II	
OTHER CONDITIONS AND REQUI	ANNEX II REMENTS OF THE MARKETING AUTI	HORISATION
OTHER CONDITIONS AND REQUIRENCE.		HORISATION
		HORISATION

ANNEX III LABELLING AND PACKAGE LEAFLET

A. LABELLING

PARTICULARS TO APPEAR ON THE OUTER PACKAGE			
BOX			
1. NAME OF THE VETERINARY MEDICI	NAL PRODUCT		
Versican Plus Pi lyophilisate and solvent for suspension for injection.			
2. STATEMENT OF ACTIVE SUBSTANCE	S		
Each dose of 1 ml contains:			
Active substances:			
Lyophilisate (live attenuated): Canine parainfluenza Type 2 virus	Minimum 10 ^{3.1} TCID ₅₀	Maximum 10 ^{5.1} TCID ₅₀	
Solvent: Water for injections (Aqua ad iniectabilia)			
3. PACKAGE SIZE			
25 x 1 dose 50 x 1 dose			
4. TARGET SPECIES			
Dogs.			
5. INDICATIONS			
6. ROUTES OF ADMINISTRATION			
Subcutaneous use.			
7. WITHDRAWAL PERIODS			
8. EXPIRY DATE			
Exp. {mm/yyyy} Once reconstituted use immediately.			

Store and transport refrigerated. Do not freeze. Protect from light.
10. THE WORDS "READ THE PACKAGE LEAFLET BEFORE USE"
Read the package leaflet before use.
11. THE WORDS "FOR ANIMAL TREATMENT ONLY"
For animal treatment only.
12. THE WORDS "KEEP OUT OF THE SIGHT AND REACH OF CHILDREN"
Keep out of the sight and reach of children.
13. NAME OF THE MARKETING AUTHORISATION HOLDER
Zoetis Belgium
14. MARKETING AUTHORISATION NUMBERS
EU/2/14/168/001 25x 1 dose EU/2/14/168/002 50 x 1 dose
15. BATCH NUMBER
Lot {number}

9.

SPECIAL STORAGE PRECAUTIONS

MINIMUM PARTICULARS TO APPEAR ON SMALL IMMEDIATE PACKAGING UNITS VIAL (1 DOSE LYOPHILISATE)

1. NAME OF THE VETERINARY MEDICINAL PRODUCT

Versican Plus Pi



2. QUANTITATIVE PARTICULARS OF THE ACTIVE SUBSTANCES

Pi

1 dose

3. BATCH NUMBER

Lot {number}

4. EXPIRY DATE

Exp. {mm/yyyy}

Once reconstituted use immediately.

MINIMUM PARTICULARS TO APPEAR ON SMALL IMMEDIATE PACKAGING UNITS VIAL (1 ML SOLVENT) 1. NAME OF THE VETERINARY MEDICINAL PRODUCT

Versican Plus Pi



2. QUANTITATIVE PARTICULARS OF THE ACTIVE SUBSTANCES

Aqua ad iniectabilia 1 ml

3. BATCH NUMBER

Lot {number}

4. EXPIRY DATE

Exp. {mm/yyyy}

B. PACKAGE LEAFLET

PACKAGE LEAFLET

1. Name of the veterinary medicinal product

Versican Plus Pi lyophilisate and solvent for suspension for injection for dogs

2. Composition

Each dose of 1 ml contains:

Active substances:

	Minimum	Maximum
<u>Lyophilisate (live attenuated):</u> Canine parainfluenza Type 2 virus, strain CPiV-2 Bio 15	10 ^{3.1} TCID ₅₀ *	10 ^{5.1} TCID ₅₀ *
Solvent: Water for injections (<i>Aqua ad iniectabilia</i>)		1 ml

^{*} Tissue culture infectious dose 50%.

The visual appearance is as follows:

Lyophilisate: spongy matter of white colour.

Solvent: clear colourless liquid.

3. Target species

Dogs.

4. Indications for use

Active immunisation of dogs from 6 weeks of age:

 to prevent clinical signs (nasal and ocular discharge) and reduce viral excretion caused by canine parainfluenza virus.

Onset of immunity:

3 weeks after completion of the primary course.

Duration of immunity:

At least one year following the primary vaccination course.

5. Contraindications

None.

6. Special warnings

Special warnings:

A good immune response is reliant on a fully competent immune system. Immunocompetence of the animal may be compromised by a variety of factors including poor health, nutritional status, genetic factors, concurrent medicinal therapy and stress.

Vaccinate healthy animals only.

Special precautions for safe use in the target species:

The live attenuated virus vaccine strain CPiV may be shed by vaccinated dogs following vaccination. However, due to the low pathogenicity of this strain, it is not necessary to keep vaccinated dogs separated from non-vaccinated dogs.

Special precautions to be taken by the person administering the veterinary medicinal product to animals:

In case of accidental self-injection, seek medical advice immediately and show the package leaflet or the label to the physician.

Special precautions for the protection of the environment:

Not applicable.

Pregnancy and lactation:

Can be used during the second and third stages of pregnancy. Safety of the product during the early stage of pregnancy and during lactation has not been investigated.

<u>Interaction</u> with other medicinal products and other forms of interaction:

No information is available on the safety and efficacy of this vaccine when used with any other veterinary medicinal product other than Versiguard Rabies and Versican Plus L4. A decision to use this vaccine before or after any other veterinary medicinal product therefore needs to be made on a case by case basis.

Leptospira:

If protection against *Leptospira* is required, dogs can be vaccinated with two doses of Versican Plus Pi mixed with Versican Plus L4 3–4 weeks apart from 6 weeks of age:

The contents of a single vial of Versican Plus Pi should be reconstituted with the contents of a single vial of Versican Plus L4 (instead of the solvent). Once mixed, the contents of the vial should appear a whitish to yellowish colour with light opalescence. The mixed vaccines should be injected immediately via the subcutaneous route.

Rabies:

If protection against rabies is required:

First dose: Versican Plus Pi from 8–9 weeks of age.

Second dose: Versican Plus Pi mixed with Versiguard Rabies 3–4 weeks later, but not before 12 weeks of age.

The contents of a single vial of Versican Plus Pi should be reconstituted with the contents of a single vial of Versiguard Rabies (instead of the solvent). Once mixed, the contents of the vial should appear a pink/red or yellowish colour with light opalescence. The mixed vaccines should be injected immediately via the subcutaneous route.

The efficacy of the rabies fraction is proven after a single dose from 12 weeks of age in laboratory studies. However, in field studies 10% of seronegative dogs did not show seroconversion (>0.1 IU/ml) 3–4 weeks after single primary vaccination against rabies.

Some animals may also not show titres > 0.5 IU/ml after the primary vaccination. Antibody titres drop over the course of the 3-year duration of immunity, although dogs are protected when challenged. In case of travelling to risk areas or outside the EU, veterinary surgeons may wish to give additional rabies vaccinations after 12 weeks of age to ensure that the vaccinated dogs have an antibody titre of ≥ 0.5 IU/ml, which is generally regarded as sufficiently protective and that they meet the travel test requirements (antibody titres ≥ 0.5 IU/ml).

Although the efficacy of the rabies fraction has been demonstrated following administration at 12 weeks, at the discretion of the veterinary surgeon, in case of need, dogs younger than 8 weeks can be vaccinated with Versican Plus Pi mixed with Versiguard Rabies as the safety of this association has been demonstrated in 6-week old dogs.

Overdose:

No adverse events other than those mentioned in section "Adverse events" were observed after administration of a 10-fold overdose of the vaccine. However, in a minority of animals pain was observed at the injection site immediately after administration of a 10-fold overdose of the vaccine.

Special restrictions for use and special conditions for use:

Not applicable.

Major incompatibilities:

Do not mix with any other veterinary medicinal product except those mentioned in section "Interaction with other medicinal products and other forms of interaction".

7. Adverse events

Dogs:

Common (1 to 10 animals / 100 animals treated):

injection site swelling¹

Rare (1 to 10 animals / 10,000 animals treated):

hypersensitivity reaction² (anaphylaxis, angioedema, circulatory shock, collapse, diarrhoea, dyspnoea, vomiting)

anorexia, decreased activity

Very rare (<1 animal / 10,000 animals treated, including isolated reports):

hyperthermia, lethargy, malaise

Reporting adverse events is important. It allows continuous safety monitoring of a product. If you notice any side effects, even those not already listed in this package leaflet, or you think that the medicine has not worked, please contact, in the first instance, your veterinarian. You can also report any adverse events to the marketing authorisation holder using the contact details at the end of this leaflet, or via your national reporting system: {national system details}.

8. Dosage for each species, routes and method of administration

Subcutaneous use.

Primary vaccination scheme:

Two doses of Versican Plus Pi 3-4 weeks apart from 6 weeks of age.

Re-vaccination scheme:

A single dose of Versican Plus Pi to be given annually.

9. Advice on correct administration

Aseptically reconstitute the lyophilisate with the solvent. Shake well and administer immediately the entire contents (1 ml) of the reconstituted product.

Appearance of the reconstituted vaccine: clear whitish to yellowish colour with light opalescence.

¹A transient swelling (up to 5 cm) which can be painful, warm or reddened. Any such swelling will either have spontaneously resolved or be greatly diminished by 14 days after vaccination.

²If a hypersensitivity reaction occurs, appropriate treatment should be administered without delay. Such reactions may evolve to a more severe condition which may be life-threatening.

10. Withdrawal periods

Not applicable.

11. Special storage precautions

Keep out of the sight and reach of children.

Store and transport refrigerated ($2 \, ^{\circ}\text{C} - 8 \, ^{\circ}\text{C}$).

Do not freeze.

Protect from light.

Do not use this veterinary medicinal product after the expiry date which is stated on the label after Exp.

The expiry date refers to the last day of that month.

Shelf life after reconstitution according to directions: use immediately.

12. Special precautions for disposal

Medicines should not be disposed of via wastewater or household waste.

Use take-back schemes for the disposal of any unused veterinary medicinal product or waste materials derived thereof in accordance with local requirements and with any applicable national collection systems. These measures should help to protect the environment.

Ask your veterinary surgeon or pharmacist how to dispose of medicines no longer required.

13. Classification of veterinary medicinal products

Veterinary medicinal product subject to prescription.

14. Marketing authorisation numbers and pack sizes

EU/2/14/168/001-002

Plastic box containing 25 vials (1 dose) of lyophilisate and 25 vials (1 ml) of solvent. Plastic box containing 50 vials (1 dose) of lyophilisate and 50 vials (1 ml) of solvent.

Not all pack sizes may be marketed.

15. Date on which the package leaflet was last revised

{MM/YYYY}

Detailed information on this veterinary medicinal product is available in the Union Product Database (https://medicines.health.europa.eu/veterinary).

16. Contact details

Marketing authorisation holder and contact details to report suspected adverse reactions:

Zoetis Belgium Rue Laid Burniat 1 1348 Louvain-La-Neuve Belgium

België/Belgique/Belgien

Tél/Tel: +32 (0) 800 99 189 pharmvig-belux@zoetis.com

Република България

Тел: +359 888 51 30 30 zoetisromania@zoetis.com

Česká republika

Tel: +420 257 101 111 infovet.cz@zoetis.com

Danmark

Tlf: +45 70 20 73 05 adr.scandinavia@zoetis.com

Deutschland

Tel: +49 30 2020 0049

tierarzneimittelsicherheit@zoetis.com

Eesti

Tel: +370 610 05088 zoetis.estonia@zoetis.com

Ελλάδα

Tηλ: +30 210 6791900 infogr@zoetis.com

España

Tel: +34 91 4191900 regulatory.spain@zoetis.com

France

Tél: +33 (0)800 73 00 65 contacteznous@zoetis.com

Hrvatska

Tel: +385 1 6441 462

pv.westernbalkans@zoetis.com

Ireland

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Ísland

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tierarzneimittelsicherheit@zoetis.com

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farmacovigilanza.italia@zoetis.com

Κύπρος

Tηλ: +30 210 6791900 infogr@zoetis.com

Latvija

Tel: +370 610 05088 zoetis.latvia@zoetis.com

<u>Manufacturer responsible for batch release</u>: Bioveta a.s.

Komenskeho 212/12 683 23 Ivanovice Na Hane Czechia

Suomi/Finland

Puh/Tel: +358 10 336 7000 laaketurva@zoetis.com

Sverige

Tel: +46 (0) 76 760 0677 adr.scandinavia@zoetis.com

United Kingdom (Northern Ireland)

Tel: +353 (0) 1 256 9800 pvsupportireland@zoetis.com

17. Other information

The vaccine is intended for the active immunisation of healthy puppies and dogs against disease caused by canine parainfluenza virus.