ANNEX I

SUMMARY OF PRODUCT CHARACTERISTICS

1. NAME OF THE VETERINARY MEDICINAL PRODUCT

Purevax RCPCh lyophilisate and solvent for suspension for injection

2. QUALITATIVE AND QUANTITATIVE COMPOSITION

Per dose of 1 ml or 0.5 ml:

Active substances:

Attenuated feline rhinotracheitis herpesvirus (FHV F2 strain)	$\geq 10^{4.9} \text{ CCID}_{50}^{1}$
Inactivated feline calicivirus (FCV 431 and G1 strains) antigens	\geq 2.0 ELISA U.
Attenuated Chlamydophila felis (905 strain)	$\geq 10^{3.0} \text{ EID}_{50}^2$
Attenuated feline panleucopenia virus (PLI IV)	$\geq 10^{3.5} \text{ CCID}_{50}^{1}$

 1 cell culture infective dose 50%

 2 egg infective dose 50%

Excipients:

Qualitative composition of excipients and other constituents	Quantitative composition if that information is essential for proper administration of the veterinary medicinal product
Lyophilisate:	
Sucrose	
Sorbitol	
Dextran 40	
Casein hydrolysate	
Collagen hydrolysate	
Dipotassium phosphate	
Potassium dihydrogen phosphate	
Potassium hydroxide	
Sodium chloride	
Disodium hydrogen orthophosphate	
Monopotassium phosphate anhydrous	
Water for injections	
Solvent:	
Water for injections	q.s. 1 ml or 0.5 ml

Lyophilisate: homogeneous beige pellet. Solvent: clear colourless liquid.

3. CLINICAL INFORMATION

3.1 Target species

Cats

3.2 Indications for use for each target species

Active immunisation of cats aged 8 weeks and older:

- against feline viral rhinotracheitis to reduce clinical signs,
- against calicivirus infection to reduce clinical signs,
- against Chlamydophila felis infection to reduce clinical signs,
- against feline panleucopenia to prevent mortality and clinical signs.

Onsets of immunity have been demonstrated 1 week after primary vaccination course for rhinotracheitis, calicivirus, *Chlamydophila felis* and panleucopenia components.

Duration of immunity:

- Rhinotracheitis, calicivirosis and panleucopenia components: 1 year after primary vaccination course and 3 years after the last re-vaccination
- Chlamydophila felis component: 1 year after the last re-vaccination.

3.3 Contraindications

None.

3.4 Special warnings

Vaccinate healthy animals only.

3.5 Special precautions for use

Special precautions for safe use in the target species: None.

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.

This vaccine should not be handled by persons who are immunodeficient or taking immunosuppressive medicinal products. If self-injection occurs, immediate medical advice should be sought and the doctor informed that self-injection with a living chlamydial vaccine has occurred.

Special precautions for the protection of the environment:

Not applicable.

3.6 Adverse events

Cats:

Common (1 to 10 animals / 100 animals treated):	Transient apathy, anorexia, and hyperthermia ¹ (observed during safety and field studies). Injection site reactions (slight pain at palpation, itching or limited oedema) ² (observed during safety and field studies)
Uncommon	Hypersensitivity reaction ³ (observed in field studies)
(1 to 10 animals / 1,000 animals treated):	

Very rare	Emesis ⁴ ; transient hyperthermia and lethargy,
(<1 animal / 10,000 animals treated, including isolated reports):	sometimes associated with lameness ₅ (based on post- marketing experience)

¹ lasting usually for 1 or 2 days

³ may require appropriate symptomatic treatment

⁴ mostly within 24 to 48 hours

⁵ observed 1 to 3 weeks following booster vaccination in adult cats

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 its local representative or the national competent authority via the national reporting system. See also the last section of the package leaflet for respective contact details.

3.7 Use during pregnancy, lactation or lay

Do not use during the whole pregnancy and lactation.

3.8 Interactions with other medicinal products and other forms of interaction

Safety and efficacy data are available which demonstrate that this vaccine can be mixed and administered with Boehringer Ingelheim non-adjuvanted vaccine against feline leukaemia and/or administered the same day but not mixed with Boehringer Ingelheim adjuvanted vaccine against rabies.

Safety and efficacy data are available which demonstrate that this vaccine can be mixed and administered with Boehringer Ingelheim non-adjuvanted vaccine against rabies.

No information is available on the safety and efficacy of this vaccine when used with any other veterinary medicinal product except the products mentioned above. 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.

3.9 Administration routes and dosage

Subcutaneous route.

Reconstitute gently the vaccine in order to obtain a uniform suspension with limited foam formation. Visual appearance after reconstitution: clear slightly yellow suspension.

After reconstitution of the lyophilisate with 0.5 ml or 1 ml of the solvent (depending on the presentation chosen), inject one dose of vaccine according to the following vaccination scheme:

Primary vaccination course:

- first injection: from 8 weeks of age,
- second injection: 3 to 4 weeks later.

Where high levels of maternal antibodies against rhinotracheitis, calicivirosis, panleucopenia or *Chlamydophila* components are expected to be present (e.g. in kittens of 9 to 12 weeks of age born from queens which were vaccinated before pregnancy and/or with known or suspected previous exposure to the pathogen(s)), the primary vaccination course should be delayed until 12 weeks of age.

Revaccination:

- the first revaccination must be carried out for all components one year after the primary vaccination course,

² disappearing within 1 or 2 weeks at most

- subsequent revaccinations must be carried out
 - Chlamydiosis component: every year.
 - Rhinotracheitis, calicivirosis and panleucopenia components: at intervals of up to three years.

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

No effect other than those already mentioned in section 3.6 "Adverse events" have been observed, except hyperthermia that may exceptionally last 5 days.

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

For administration only by a veterinarian.

3.12 Withdrawal periods

Not applicable.

4. IMMUNOLOGICAL INFORMATION

4.1 ATCvet code: QI06AJ03 (live feline rhinotracheitis virus + inactivated feline calicivirus antigen + live feline panleucopenia virus / parvovirus + live chlamydia).

Vaccine against feline viral rhinotracheitis, feline calicivirosis, chlamydiosis and feline panleucopenia. Stimulates active immunity against feline rhinotracheitis herpesvirus, feline calicivirus, *Chlamydophila felis* and feline panleucopenia virus.

The product was shown to reduce excretion of feline calicivirus at onset of immunity and for one year after vaccination.

5. PHARMACEUTICAL PARTICULARS

5.1 Major incompatibilities

Do not mix with Boehringer Ingelheim adjuvanted vaccine against rabies.

5.2 Shelf life

Shelf life of the veterinary medicinal product as packaged for sale: 18 months. Shelf life after reconstitution according to directions: use immediately.

5.3 Special precautions for storage

Store and transport refrigerated (2 °C - 8 °C). Protect from light. Do not freeze.

5.4 Nature and composition of immediate packaging

Type I glass bottle containing 1 dose of lyophilisate and type I glass bottle containing 1 ml or 0.5 ml of solvent, both closed with a butyl elastomer closure and sealed with an aluminium or plastic cap.

Plastic box containing 10 bottles of 1 dose of lyophilisate and 10 bottles of 1 ml of solvent. Plastic box containing 50 bottles of 1 dose of lyophilisate and 50 bottles of 1 ml of solvent. Plastic box containing 10 bottles of 1 dose of lyophilisate and 10 bottles of 0.5 ml of solvent. Plastic box containing 50 bottles of 1 dose of lyophilisate and 50 bottles of 0.5 ml of solvent.

Not all pack sizes may be marketed.

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

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

Boehringer Ingelheim Vetmedica GmbH

7. MARKETING AUTHORISATION NUMBER(S)

EU/2/04/050/001-004

8. DATE OF FIRST AUTHORISATION

Date of first authorisation: 23/02/2005

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

DD/MM/YYYY

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.

ANNEX II

OTHER CONDITIONS AND REQUIREMENTS OF THE MARKETING AUTHORISATION

None.

ANNEX III

LABELLING AND PACKAGE LEAFLET

A. LABELLING

PARTICULARS TO APPEAR ON THE OUTER PACKAGE

Plastic box of 10 bottles of lyophilisate and 10 bottles of solvent Plastic box of 50 bottles of lyophilisate and 50 bottles of solvent

1. NAME OF THE VETERINARY MEDICINAL PRODUCT

Purevax RCPCh lyophilisate and solvent for suspension for injection

2. STATEMENT OF ACTIVE SUBSTANCES

Per dose of 1 ml or 0.5 ml:	
FHV (F2 strain)	$\geq 10^{4.9} \text{ CCID}_{50}$
FCV (431 and G1 strains)	
Chlamydophila felis (905 strain)	$\geq 10^{3.0} \text{ EID}_{50}$
FPV (PLI IV)	$\geq 10^{3.5} \text{ CCID}_{50}.$

3. PACKAGE SIZE

Lyophilisate (10 x 1 dose) + solvent (10 x 1 ml) Lyophilisate (50 x 1 dose) + solvent (50 x 1 ml) Lyophilisate (10 x 1 dose) + solvent (10 x 0.5 ml) Lyophilisate (50 x 1 dose) + solvent (50 x 0.5 ml)

4. TARGET SPECIES

Cats

5. INDICATIONS

6. ROUTES OF ADMINISTRATION

Subcutaneous use.

7. WITHDRAWAL PERIODS

8. EXPIRY DATE

Exp. {dd/mm/yyyy} Once reconstituted use immediately.

9. SPECIAL STORAGE PRECAUTIONS

Store and transport refrigerated. Protect from light. Do not freeze.

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

Boehringer Ingelheim Vetmedica GmbH

14. MARKETING AUTHORISATION NUMBERS

EU/2/04/050/001 Lyophilisate (10 x 1 dose) + solvent (10 x 1 ml) EU/2/04/050/002 Lyophilisate (50 x 1 dose) + solvent (50 x 1 ml) EU/2/04/050/003 Lyophilisate (10 x 1 dose) + solvent (10 x 0.5 ml) EU/2/04/050/004 Lyophilisate (50 x 1 dose) + solvent (50 x 0.5 ml)

15. BATCH NUMBER

Lot {number}

MINIMUM PARTICULARS TO APPEAR ON SMALL IMMEDIATE PACKAGING UNITS

Lyophilisate bottle

1. NAME OF THE VETERINARY MEDICINAL PRODUCT

Purevax RCPCh



2. QUANTITATIVE PARTICULARS OF THE ACTIVE SUBSTANCES

1 dose

3. BATCH NUMBER

Lot

4. EXPIRY DATE

Exp. (dd/{mm/yyyy}

MINIMUM PARTICULARS TO APPEAR ON SMALL IMMEDIATE PACKAGING UNITS

Solvent bottle

1. NAME OF THE VETERINARY MEDICINAL PRODUCT

Purevax RCPCh solvent



2. QUANTITATIVE PARTICULARS OF THE ACTIVE SUBSTANCES

1 ml or 0.5 ml

3. BATCH NUMBER

Lot

4. EXPIRY DATE

Exp. {mm/yyyy}

B. PACKAGE LEAFLET

PACKAGE LEAFLET:

1. Name of the veterinary medicinal product

Purevax RCPCh lyophilisate and solvent for suspension for injection

2. Composition

Per dose of 1 ml or 0.5 ml:

Active Substances:

Attenuated feline rhinotracheitis herpesvirus (FHV F2 strain)	$\geq 10^{4.9} \text{ CCID}_{50}^{1}$
Inactivated feline calicivirus (FCV 431 and FCV G1 strains) antigens	\geq 2.0 ELISA U.
Attenuated Chlamydophila felis (905 strain)	$\geq 10^{3.0} \text{ EID}_{50}^2$
Attenuated feline panleucopenia virus (PLI IV)	$\geq 10^{3.5} \text{ CCID}_{50}^{1}$

¹ cell culture infective dose 50% ² egg infective dose 50%

Solvent:

Lyophilisate: homogeneous beige pellet. Solvent: clear colourless liquid.

3. Target species

Cats

4. Indications for use

Active immunisation of cats aged 8 weeks and older:

- against feline viral rhinotracheitis to reduce clinical signs,
- against calicivirus infection to reduce clinical signs,
- against *Chlamydophila felis* infection to reduce clinical signs,
- against feline panleucopenia to prevent mortality and clinical signs.

Onsets of immunity have been demonstrated 1 week after primary vaccination course for rhinotracheitis, calicivirus, *Chlamydophila felis* and panleucopenia components.

Duration of immunity:

- Rhinotracheitis, calicivirosis and panleucopenia components: 1 year after primary vaccination and 3 years after the last re-vaccination
- Chlamydophila felis component: 1 year after the last re-vaccination.

5. Contraindications

None.

6. Special warnings

Vaccinate healthy animals only.

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.

This vaccine should not be handled by persons who are immunodeficient or taking immunosuppressive medicinal products. If self-injection occurs, immediate medical advice should be sought and the doctor informed that self-injection with a living chlamydial vaccine has occurred.

Pregnancy and lactation:

Do not use during the whole pregnancy and lactation.

Interactions with other medicinal products and other forms of interaction:

Safety and efficacy data are available which demonstrate that this vaccine can be mixed and administered with Boehringer Ingelheim non-adjuvanted vaccine against feline leukaemia and/or administered the same day but not mixed with Boehringer Ingelheim adjuvanted vaccine against rabies.

Safety and efficacy data are available which demonstrate that this vaccine can be mixed and administered with Boehringer Ingelheim non-adjuvanted vaccine against rabies.

No information is available on the safety and efficacy of this vaccine when used with any other veterinary medicinal product except the products mentioned above. 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.

Overdose:

No effect other than those already mentioned in section on "Adverse events" have been observed after the administration of several doses, except hyperthermia that may exceptionally last 5 days.

<u>Special restrictions for use and special conditions for use:</u> For administration only by a veterinarian.

Major incompatibilities:

Do not mix with Boehringer Ingelheim adjuvanted vaccine against rabies.

7. Adverse events

Cats:

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

Transient apathy, anorexia, and hyperthermia¹ (observed during safety and field studies).

Injection site reactions (slight pain at palpation, itching or limited oedema)² (observed during safety and field studies)

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

Hypersensitivity reaction³ (observed in field studies)

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

Emesis⁴, transient hyperthermia and lethargy, sometimes associated with lameness⁵ (based on post-marketing experience)

¹ lasting usually for 1 or 2 days

² disappearing within 1 or 2 weeks at most

³ may require appropriate symptomatic treatment.

⁴ mostly within 24 to 48 hours

⁵ observed 1 to 3 weeks following booster vaccination in adult cats

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 local representative of the marketing authorisation holder using the contact details at the end of this leaflet, or via your national reporting system.

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

Subcutaneous route.

After reconstitution of the lyophilisate with 0.5 ml or 1 ml of the solvent (depending on the presentation chosen), inject one dose of vaccine according to the following vaccination scheme:

Primary vaccination course:

- first injection: from 8 weeks of age,
- second injection: 3 to 4 weeks later.

Where high levels of maternal antibodies against rhinotracheitis, calicivirosis, panleucopenia or *Chlamydophila* components are expected to be present (e.g. in kittens of 9 to 12 weeks of age born from queens which were vaccinated before pregnancy and/or with known or suspected previous exposure to the pathogen(s)), the primary vaccination course should be delayed until 12 weeks of age.

Revaccination:

- the first revaccination must be carried out for all components one year after the primary vaccination course,
- subsequent revaccinations must be carried out:
 - Chlamydiosis component: every year.
 - Rhinotracheitis, calicivirosis and panleucopenia components: at intervals of up to three years.

9. Advice on correct administration

Reconstitute gently the vaccine in order to obtain a uniform suspension with limited foam formation. Visual appearance after reconstitution: clear slightly yellow suspension.

10. Withdrawal periods

Not applicable.

11. Special storage precautions

Keep out of the sight and reach of children. Store and transport refrigerated (2 °C – 8 °C). Protect from light. Do not freeze. Do not use this veterinary medicinal product after the expiry date which is stated on the bottle after Exp. Shelf life after reconstitution according to directions: use immediately.

12. Special precautions for disposal

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.

13. Classification of veterinary medicinal products

Veterinary medicinal product subject to prescription.

14. Marketing authorisation numbers and pack sizes

EU/2/04/050/001-004

Plastic box containing: 10 x 1 dose of lyophilisate and 10 x 1 ml of solvent or 50 x 1 dose of lyophilisate and 50 x 1 ml of solvent or 10 x 1 dose of lyophilisate and 10 x 0.5 ml of solvent or 50 x 1 dose of lyophilisate and 50 x 0.5 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.

16. Contact details

Marketing authorisation holder: Boehringer Ingelheim Vetmedica GmbH 55216 Ingelheim/Rhein Germany

Manufacturer responsible for batch release: Boehringer Ingelheim Animal Health France SCS Laboratoire Porte des Alpes Rue de l'Aviation 69800 Saint Priest France

Local representatives and contact details to report suspected adverse reactions

België/Belgique/Belgien

Boehringer Ingelheim Animal Health Belgium SA Tél/Tel: + 32 2 773 34 56

Република България

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Deutschland Boehringer Ingelheim Vetmedica GmbH Tel: 0800 290 0 270

Eesti

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17. Other information

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Slovenská republika Boehringer Ingelheim RCV GmbH & Co KG, o.z. Tel: +421 2 5810 1211 Suomi/Finland Vetcare Oy Puh/Tel: + 358 201443360 Sverige Boehringer Ingelheim Animal Health Nordics A/S Tlf: +46 (0)40-23 34 00 United Kingdom (Northern Ireland) Boehringer Ingelheim Animal Health UK Limited

Tel: + 44 1344 746957

The product was shown to reduce excretion of feline calicivirus at onset of immunity and for one year after vaccination.