



dr. van haeringen laboratorium b.v.

a VHLGenetics company

## Inzendformulier Hond (Pagina 1 van 5)

### Instructies:

- Met één voorblad kunnen meerdere dieren ingezonden worden;
- Per ingezonden monster één diergegevensblad meesturen;
- Op het monster identificatienummer en naam van het dier vermelden;
- Factuur en uitslag worden altijd naar hetzelfde adres gestuurd;
- Via [www.vhlgenetics.com](http://www.vhlgenetics.com) dient u zelf te controleren of de gewenste analyse rasafhankelijk is;
- Onvolledige inzendingen worden niet in behandeling genomen.

### Adres voor factuur en uitslag

|                 |                       |         |  |
|-----------------|-----------------------|---------|--|
| Bedrijfsnaam    | Indien van toepassing |         |  |
| Contactpersoon* | Dhr./mevr.            |         |  |
| Adres*          |                       |         |  |
| Postcode*       |                       |         |  |
| Woonplaats*     |                       |         |  |
| E-mailadres*    |                       |         |  |
| Telefoon        |                       | Fax     |  |
| Klant nr.       |                       | BTW nr. |  |

\* Deze velden verplicht invullen

#### Administratiekosten

Bij inzendingen via dit inzendformulier wordt per monster €2,95 (excl. BTW) administratiekosten in rekening gebracht. Bij bestellingen via onze webshop betaalt u géén administratiekosten. Bezoek onze webshop op [www.vhlgenetics.com](http://www.vhlgenetics.com).

Aantal ingezonden monsters: \_\_\_\_\_ (verplicht)

Hierbij bevestig ik dat deze gegevens correct zijn ingevuld en dat ik door ondertekening van dit formulier de Algemene Voorwaarden VHL 2017 - waaronder een beperking van de aansprakelijkheid - accepteer.

Plaats \_\_\_\_\_ Datum \_\_\_\_\_ Naam \_\_\_\_\_ Handtekening \_\_\_\_\_



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| Diergegevens (Pagina 2 van 5) |   |   |       |
|-------------------------------|---|---|-------|
| Naam                          |   |   |       |
| Registratie nr.               |   |   |       |
| Chip/Tatoeage                 |   |   |       |
| Geboortedatum                 | dd-mm-jjjj                                  | Geslacht  | V / M |
| Ras                           |   |   |       |
| VHL-ID                        | Indien analyse op eerder ingezonden monster |   |       |
| Naam vader                    |   |   |       |
| Reg.nr. vader                 |   |   |       |
| Naam moeder                   |   |   |       |
| Reg.nr. moeder                |   |   |       |
| DNA Certificaat               | Ja / Nee                                    | Indien aangevraagd 'Verwantschap, Afstamming of Identiteit' |       |

## Gewenste analyse(s) hond

### Genoom Scan

- H692 CanineHD Whole-Genome Genotyping

### Test op aanvraag

- H340 Brachycefal syndroom  H417 Leukoencephalomyelopathie  
 H341 Cerebellaire Ataxie (SDCA2)  H376 Schedelvorm Diversiteit  
 H813 Glycogeenstapeling GSD Type IIIa (GSDIIIa)

### Verwantschap, Afstamming en Identiteit

- H202 Moederschapsverificatie Hond  H205 Profiel vastleggen Hond  
 H200 Ouderschapsverificatie Hond  H201 Vaderschapsverificatie Hond

### Combinatiepakketten

- H321 CombiBreed Australian Shepherd  H480 CombiBreed FCI Rasgroep 07  
 H319 CombiBreed Border Collie  H481 CombiBreed FCI Rasgroep 08  
 H320 CombiBreed Duitse Herder  H482 CombiBreed FCI Rasgroep 09  
 H474 CombiBreed FCI Rasgroep 01  H483 CombiBreed FCI Rasgroep 10  
 H475 CombiBreed FCI Rasgroep 02  H322 CombiBreed Golden Retriever  
 H476 CombiBreed FCI Rasgroep 03  H323 CombiBreed Labrador Retriever  
 H477 CombiBreed FCI Rasgroep 04  H398 Combinatiepakket Partnerlab  
 H478 CombiBreed FCI Rasgroep 05  H469 Combinatiepakket Vachtkleur  
 H479 CombiBreed FCI Rasgroep 06

### Erfelijke Ziekten

- H396 Adult Onset Neuropathy (AON)  H701 Koperstapeling



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## Diergegevens (Pagina 3 van 5)

| Naam            |  |
|-----------------|--|
| Registratie nr. |  |

|          |  |          |   |
|----------|--|----------|---|
| [ ] H625 | Aangeboren afwijking van verhoorning               | [ ] H724 | L2-HGA                                      |
| [ ] H626 | Aangeboren Myastheen 2 syndroom (CMS)              | [ ] H395 | 'Limb girdle' spierdystrofie (LGMD) 1       |
| [ ] H627 | Aberrant Autophagy (LSD)                           | [ ] H317 | Maculaire dystrofie van het hoornvlies      |
| [ ] H386 | Achromatopsia 2 (cone degeneration, hemeralopia)   | [ ] H746 | Maligne Hyperthermie                        |
| [ ] H387 | Achromatopsia 3 (cone degeneration, hemeralopia)   | [ ] H629 | MDR1 (partner lab)                          |
| [ ] H338 | Acute Respiratory Distress Syndroom (ARDS)         | [ ] H723 | MDR1 (patent eigenaar)                      |
| [ ] H351 | Amelogenesis Imperfecta                            | [ ] H427 | MTM   |
| [ ] H450 | Bleeding disorder due to P2RY12 defect             | [ ] H418 | Mucopolysaccharidose Type IIIa              |
| [ ] H487 | Brachyurie (Bobtail)                               | [ ] H748 | Mucopolysaccharidose Type VII               |
| [ ] H412 | C3 Deficientie                                     | [ ] H451 | Mucopolysaccharidose Type VII - 2           |
| [ ] H749 | Centronucleaire Myopatie (CNM, hiervoor HMLR)      | [ ] H306 | Multifocale Retinopatie 3 (cmr3) 1          |
| [ ] H413 | Cerebellaire Abiotrofie (NCCD)                     | [ ] H307 | Multifocale Retinopatie 3 (cmr3) 2          |
| [ ] H410 | Cerebellaire Ataxie                                | [ ] H493 | Musculaire hypertrofie (dubbele bespierung) |
| [ ] H331 | Cerebellaire Ataxie (SDCA1)                        | [ ] H359 | Muscular Dystrofie, Duchenne type (MDM)     |
| [ ] H804 | Cerebellaire Ataxie / NCL-A                        | [ ] H419 | Muscular Dystrofie, Duchenne type (MDM)     |
| [ ] H653 | Cerebellaire Ataxie 2                              | [ ] H424 | Musladin-Lueke Syndroom (MLS)               |
| [ ] H411 | Cerebellaire Ataxie, progressieve early-onset      | [ ] H391 | Myasthenia gravis-achtige ziekte            |
| [ ] H318 | Cerebellaire Corticale degeneratie                 | [ ] H368 | Myopathie                                   |
| [ ] H355 | Cerebellaire Hypoplasie                            | [ ] H738 | Myotonia Congenita                          |
| [ ] H356 | Chondrodysplasie                                   | [ ] H498 | Myotonia Congenita 2                        |
| [ ] H346 | Chondrodystrofie en ziekte van tussenwervelschijf  | [ ] H707 | Narcolepsie 1                               |
| [ ] H709 | CLAD, type I                                       | [ ] H697 | Narcolepsie 2                               |
| [ ] H484 | CLAD, Type III                                     | [ ] H698 | Narcolepsie 3                               |
| [ ] H871 | CMR1 (Canine Multifocal Retinopathy)               | [ ] H812 | Neonatale Encephalopatie                    |
| [ ] H730 | CMR2 (Canine Multifocal Retinopathy)               | [ ] H370 | Nephritis                                   |
| [ ] H705 | Collie Eye Anomaly (CEA_CH, partnerlab)            | [ ] H428 | Neuroaxonal Dystrofie                       |
| [ ] H710 | Collie Eye Anomaly (CEA_CH, patent eigen. Optigen) | [ ] H494 | Neuronal ceroid lipofuscinosis (NCL) 1      |
| [ ] H918 | Cone Degeneration                                  | [ ] H429 | Neuronal ceroid lipofuscinosis (NCL) 10     |
| [ ] H357 | Cone Rod Dystrofie 1 (crd1)                        | [ ] H499 | Neuronal ceroid lipofuscinosis (NCL) 2      |
| [ ] H358 | Cone Rod Dystrofie 2 (crd2)                        | [ ] H721 | Neuronal ceroid lipofuscinosis (NCL) 5      |
| [ ] H416 | Congenitaal Hypothyroidisme (CHG) 1                | [ ] H380 | Neuronal ceroid lipofuscinosis (NCL) 5 GR   |
| [ ] H485 | Congenitaal Hypothyroidisme (CHG) 2                | [ ] H330 | Neuronal ceroid lipofuscinosis (NCL) 6      |
| [ ] H488 | Congenitaal Hypothyroidisme (CHG) 3                | [ ] H652 | Neuronal ceroid lipofuscinosis (NCL) 8      |
| [ ] H425 | Congenitaal Myasthenic Syndroom                    | [ ] H337 | Neuronal ceroid lipofuscinosis NCL, 8       |
| [ ] H339 | Congenital Myasthenic Syndroom                     | [ ] H309 | Obesitas                                    |
| [ ] H312 | Craniomandibulaire Osteopathie                     | [ ] H393 | Oculocutaneous Albinisme                    |
| [ ] H856 | crd3   | [ ] H430 | Osteogenesis imperfecta                     |
| [ ] H766 | crd4-PRA (voorheen cord1-PRA)                      | [ ] H431 | Osteogenesis imperfecta 2                   |
| [ ] H728 | CSNB (Congenital Stationary Night Blindness)       | [ ] H381 | Osteogenesis imperfecta 3                   |
| [ ] H703 | Cystinuria I - A - 1                               | [ ] H305 | PAP-PRA1 1                                  |
| [ ] H644 | Cystinuria, type II - A                            | [ ] H301 | PAP-PRA1 2                                  |
| [ ] H643 | Cystinuria, type II - A - 1                        | [ ] H717 | PFK (Phosphofruktokinase Deficientie)       |



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## Diergegevens (Pagina 4 van 5)

| Naam            |  |
|-----------------|--|
| Registratie nr. |  |

|          |   |          |  |
|----------|---|----------|--|
| [ ] H645 | Cystinuria, type II - B                         | [ ] H509 | Polycysteuze Nieren (PKD1)                         |
| [ ] H389 | De ziekte van Alexander                         | [ ] H438 | Polycythemia                                       |
| [ ] H308 | Degeneratieve Myelopathie 2 (DM2 Berner Sennen) | [ ] H914 | Polyneuropathie 1                                  |
| [ ] H673 | Degeneratieve Myelopathie, DM (partner lab)     | [ ] H495 | Polyneuropathie 2                                  |
| [ ] H806 | Degeneratieve Myelopathie, DM (patenteigenaar)  | [ ] H379 | Polyneuropathy 3 (LPN1)                            |
| [ ] H489 | Dermatofibrose                                  | [ ] H342 | Polyneuropathie 4 (LPN2)                           |
| [ ] H434 | Dilated Cardiomyopathy (DCM)                    | [ ] H371 | PRA  |
| [ ] H739 | Dominant PRA                                    | [ ] H394 | PRA BBS4   |
| [ ] H913 | Dry Eye Curly Coat Syndroom                     | [ ] H372 | PRA crdPRA   |
| [ ] H385 | Ectodermal Dysplasia X-linked                   | [ ] H382 | PRA erd  |
| [ ] H497 | Epidermolysis bullosa, dystrophic (RDEB)        | [ ] H373 | PRA type 3   |
| [ ] H486 | Epilepsie, BFJ                                  | [ ] H390 | PRA2   |
| [ ] H674 | Episodic Falling Disease (partner lab)          | [ ] H704 | prcd PRA (partnerlab)                              |
| [ ] H883 | Episodic Falling Disease (patenteigenaar)       | [ ] H700 | prcd PRA (patent eigenaar Optigen)                 |
| [ ] H809 | Erfelijk Cataract (HC) - HSF4                   | [ ] H439 | Prekallikrein Deficientie                          |
| [ ] H699 | Erfelijk Cataract 2 (HC) - HSF4                 | [ ] H383 | Primair Glaucoom                                   |
| [ ] H672 | Exercise Induced Collapse, EIC (partner lab)    | [ ] H414 | Primaire Ciliary Dyskinesia                        |
| [ ] H805 | Exercise Induced Collapse, EIC (patenteigenaar) | [ ] H374 | Primaire Hyperoxalurie                             |
| [ ] H607 | Factor IX Deficiëntie                           | [ ] H849 | Primaire Lens Luxatie (PLL)                        |
| [ ] H435 | Factor VII deficiency                           | [ ] H740 | Pyruvaat Dehydrogenase Phosphatase 1 (PDP1)        |
| [ ] H324 | FBN2  | [ ] H741 | Pyruvaatkinase Deficientie (PKDef)                 |
| [ ] H676 | FN ,Familiaire Nephropatie (partner lab)        | [ ] H454 | Pyruvaatkinase Deficientie (PKDef) 2               |
| [ ] H729 | FN, Familiaire Nephropatie (patenteigenaar)     | [ ] H455 | Pyruvaatkinase Deficientie (PKDef) 3               |
| [ ] H375 | Fragiele Huid                                   | [ ] H768 | rcd1 PRA   |
| [ ] H736 | Fucosidosis                                     | [ ] H769 | rcd1a PRA  |
| [ ] H360 | Galblaas Mucocelie                              | [ ] H801 | rcd2 PRA   |
| [ ] H361 | Gangliosidosis, GM2, Ttype I (B variant)        | [ ] H770 | rcd3 PRA   |
| [ ] H490 | Gangliosidosis, GM2, type II                    | [ ] H511 | rcd4 PRA   |
| [ ] H496 | Glaucoma (POAG)                                 | [ ] H794 | Retinale Dysplasie Retinale Vouwing RD OSD 1       |
| [ ] H737 | Globoid Cell Leukodystrofie / Krabbes Disease   | [ ] H426 | Retinale Dysplasie Retinale Vouwing RD OSD 2       |
| [ ] H415 | Glycogeenstapeling GSD Type I                   | [ ] H423 | SCID   |
| [ ] H702 | GM1   | [ ] H456 | SCID 2   |
| [ ] H915 | gPRA  | [ ] H388 | Sensorische Neuropathie                            |
| [ ] H868 | GR_PRA1   | [ ] H336 | Sensorische neuropathie 3                          |
| [ ] H473 | GR_PRA2   | [ ] H510 | Skeletal Dysplasia 2 (SD2)                         |
| [ ] H752 | Gray Collie Syndroom (Cyclische Neutropenie)    | [ ] H747 | Spierdystrofie (GRMD)                              |
| [ ] H347 | GSDII (Ziekte van Pompe)                        | [ ] H377 | Spinaal dysrafisme                                 |
| [ ] H491 | Haemofilie A (Factor VIII) (partner lab)        | [ ] H303 | Spinocerebellaire Ataxie                           |
| [ ] H436 | Haemofilie A (Factor VIII) (patent eigenaar)    | [ ] H328 | Spinocerebellar ataxie (2)                         |
| [ ] H392 | Hemorragische diathese (Scott Syndroom)         | [ ] H327 | Tandheelkundige Hypomineralisatie (Raine Syndrome) |
| [ ] H919 | Heuplaxiteit 1                                  | [ ] H440 | Thrombasthenia                                     |
| [ ] H421 | Heuplaxiteit 2                                  | [ ] H447 | Thrombasthenia 2                                   |



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## Diergegevens (Pagina 5 van 5)

|                 |  |
|-----------------|--|
| Naam            |  |
| Registratie nr. |  |

|                               |  |                               |  |
|-------------------------------|--|-------------------------------|--|
| <input type="checkbox"/> H675 | HNPK (partner lab)                                 | <input type="checkbox"/> H441 | Thrombocytopaenia                        |
| <input type="checkbox"/> H443 | HNPK (patenteigenaar)                              | <input type="checkbox"/> H448 | Thrombopathia 2                          |
| <input type="checkbox"/> H492 | Hyperkeratose, palmoplantaire                      | <input type="checkbox"/> H449 | Thrombopathia 3                          |
| <input type="checkbox"/> H363 | Hyperkeratosis, epidermolytisch                    | <input type="checkbox"/> H442 | Thrombopatie                             |
| <input type="checkbox"/> H811 | Hyperuricemie (HUU)                                | <input type="checkbox"/> H787 | Trapped Neutrophil Syndrome (TNS)        |
| <input type="checkbox"/> H364 | Hypocatalasie                                      | <input type="checkbox"/> H432 | Tremor, X-linked                         |
| <input type="checkbox"/> H872 | Hypofysaire dwerggroei                             | <input type="checkbox"/> H771 | Type A PRA                               |
| <input type="checkbox"/> H365 | Hypomyelinisatie                                   | <input type="checkbox"/> H344 | Vergroeide lip/gehemelte en tenen (CLPS) |
| <input type="checkbox"/> H873 | Ichthyosis 2                                       | <input type="checkbox"/> H433 | Vitamin D-deficientie rickets, type II   |
| <input type="checkbox"/> H304 | Ichthyosis 3                                       | <input type="checkbox"/> H642 | Von Willebrand disease 3 - 2             |
| <input type="checkbox"/> H378 | Ichthyosis 4                                       | <input type="checkbox"/> H677 | Von-Willebrands Disease Type 1           |
| <input type="checkbox"/> H384 | Ichthyosis 5                                       | <input type="checkbox"/> H743 | Von-Willebrands Disease Type 2           |
| <input type="checkbox"/> H366 | IGS (Selectieve Cobalamine Malabsorptie) 1         | <input type="checkbox"/> H744 | Von-Willebrands Disease Type 3           |
| <input type="checkbox"/> H367 | IGS (Selectieve Cobalamine Malabsorptie) 2         | <input type="checkbox"/> H345 | Von-Willebrands Ziekte Type 2-2          |
| <input type="checkbox"/> H332 | IGS (Selectieve Cobalamine Malabsorptie) 3         | <input type="checkbox"/> H696 | Warburg Micro Syndroom 1 (WARBM1)        |
| <input type="checkbox"/> H335 | Juvenile Myoclonische Epilepsie                    | <input type="checkbox"/> H772 | X Gebonden PRA1 (XL PRA1)                |
| <input type="checkbox"/> H329 | Juvenile Laryngeal Paralysis Polyneuropathy (JLPP) | <input type="checkbox"/> H745 | X-SCID                                   |

### Overige erfelijke kenmerken

|                               |                                    |                               |  |
|-------------------------------|------------------------------------|-------------------------------|--|
| <input type="checkbox"/> H765 | Haarlengte                         | <input type="checkbox"/> H316 | Vachtkleur H-locus (Harlekijn)         |
| <input type="checkbox"/> H848 | Improper Coat/Furnishings          | <input type="checkbox"/> H819 | Vachtkleur K-locus                     |
| <input type="checkbox"/> H921 | Vacht - Gekrulde Vacht             | <input type="checkbox"/> H630 | Vachtkleur Merle (partner lab)         |
| <input type="checkbox"/> H820 | Vachtkleur A-locus                 | <input type="checkbox"/> H930 | Vachtkleur Merle (patenteigenaar)      |
| <input type="checkbox"/> H733 | Vachtkleur B-locus                 | <input type="checkbox"/> H354 | Vachtkleur Panda White Spotting        |
| <input type="checkbox"/> H847 | Vachtkleur D-locus Improved (MLPH) | <input type="checkbox"/> H326 | Vachtkleur Piebald (bont)              |
| <input type="checkbox"/> H734 | Vachtkleur E-locus                 | <input type="checkbox"/> H353 | Vachtkleur Saddle tan vs black-and-tan |
| <input type="checkbox"/> H818 | Vachtkleur Em-locus                |                               |  |