

REFERENCE NO.: 2020 - 35004**OWNER:**KAČA KACIAN
BUKEVJE 25C
HR-10411 ORLE
CROATIA**SAMPLE TAKEN BY:** VLATKA-ANTONIJA CSIK, DVM**SAMPLE INFORMATION****ANIMAL NAME/LABEL:** GALLIVANT LIVE TO RIDE**SPECIES:** DOG**BREED:** LABRADOR RETRIEVER**SEX:** MALE**MICROCHIP NO.:** 956000010780773**TATOO NO.:** NOT PROVIDED**PEDIGREE NO.:** SS14478308**SAMPLE TYPE:** BLOOD**SAMPLING DATE:** 01.09.2020

DNA PROFILE REPORT

MARKER	GENOTYPE	MARKER	GENOTYPE
Amelogenin:	Y / X	INRA21:	97 / 101
AHT121:	102 / 102	INU005:	124 / 126
AHT137:	149 / 153	INU030:	144 / 150
AHTh130:	125 / 129	INU055:	208 / 208
AHTh171:	223 / 223	REN105LO3:	235 / 235
AHTh260:	240 / 246	REN162C04:	202 / 202
AHTk211:	95 / 97	REN169D01:	210 / 214
AHTk253:	288 / 288	REN169O18:	164 / 168
CXX0279:	116 / 124	REN247M23:	268 / 268
FH2054:	152 / 152	REN54P11:	226 / 232
FH2848:	232 / 244	REN64E19:	145 / 153

The nomenclature is based on the standard of ISAG Comparison Test of 2015.

AUTHORIZED SIGNATURE:

MARIBOR, 22.09.2020

REFERENCE NO.: 2020 - 35004**OWNER:**KAČA KACIAN
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GALLIVANT LIVE TO RIDE

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GENETIC REPORT

SAMPLE: BLOOD**SAMPLE TAKEN BY:** VLATKA-ANTONIJA CSIK, DVM**REQUESTED TEST:** PROGRESSIVE RETINAL ATROPHY (PRA-PRCD)**RESULT:** CLEAR (WT/WT)**COMMENT :**

The test examines presence or absence of PRCD gene mutation (c.5G>A) described as the cause of one form of progressive retinal atrophy (PRA) in several dog breeds. PRA-PRCD is a late onset disease characterized by progressive degeneration of retinal cells. PRCD gene defect is inherited as an autosomal recessive trait.

Regarding to the presence of tested mutation animals are classified in three groups:

- Clear (wt/wt) - mutation is not present, normal genotype
- Carrier (mut/wt) - one of two alleles carries tested mutation, disease is not clinically manifested
- Affected (mut/mut) - both alleles carry tested mutation, disease is clinically manifested

For each group different breeding strategies should be followed. Breeding of affected and carrier animals should be avoided. If particularly valuable animal is classified as affected, it should be bred only with clear animal. In such case, all first generation siblings will be carriers. If a carrier is bred with clear animal, 50% of siblings are expected to be clear. In case two carriers are bred, 25% of siblings are expected to be clear and 50% are expected to be carriers. However, 25% of siblings are expected to be affected, therefore such breeding practice is discouraged.

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GENETIC REPORT

SAMPLE: BLOOD**SAMPLE TAKEN BY:** VLATKA-ANTONIJA CSIK, DVM**REQUESTED TEST:** EXCERSISE INDUCED COLLAPSE (EIC)**RESULT:** CLEAR (WT/WT)**COMMENT :**

The test examines presence or absence of DNM1 gene mutation (c.767G>T) described as the cause of exercise induced collapse (EIC) in several dog breeds. EIC is a syndrome characterized by collapse episodes following strenuous exercise. DNM1 gene defect is inherited as an autosomal recessive trait.

Regarding to the presence of tested mutation animals are classified in three groups:

- Clear (wt/wt) - mutation is not present, normal genotype
- Carrier (mut/wt) - one of two alleles carries tested mutation, disease is not clinically manifested
- Affected (mut/mut) - both alleles carry tested mutation, disease is clinically manifested

For each group different breeding strategies should be followed. Breeding of affected and carrier animals should be avoided. If particularly valuable animal is classified as affected, it should be bred only with clear animal. In such case, all first generation siblings will be carriers. If a carrier is bred with clear animal, 50% of siblings are expected to be clear. In case two carriers are bred, 25% of siblings are expected to be clear and 50% are expected to be carriers. However, 25% of siblings are expected to be affected, therefore such breeding practice is discouraged.

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GENETIC REPORT

SAMPLE: BLOOD**SAMPLE TAKEN BY:** VLATKA-ANTONIJA CSIK, DVM**REQUESTED TEST:** CENTRONUCLEAR MYOPATHY (CNM)**RESULT:** CLEAR (WT/WT)**COMMENT :**

The test examines presence or absence of PTPLA gene mutation (c.191_192ins236bp) described as the cause of centronuclear myopathy (CNM) in Labrador Retriever. The disease is characterized by hypotonia, generalized muscle weakness, abnormal postures, stiff hopping gait, exercise intolerance and increased collapse when exposed to cold. CNM is inherited as an autosomal recessive trait.

Regarding to the presence of tested mutation animals are classified in three groups:

- Clear (wt/wt) - mutation is not present, normal genotype
- Carrier (mut/wt) - one of two alleles carries tested mutation, disease is not clinically manifested
- Affected (mut/mut) - both alleles carry tested mutation, disease is clinically manifested

For each group different breeding strategies should be followed. Breeding of affected and carrier animals should be avoided. If particularly valuable animal is classified as affected, it should be bred only with clear animal. In such case, all first generation siblings will be carriers. If a carrier is bred with clear animal, 50% of siblings are expected to be clear. In case two carriers are bred, 25% of siblings are expected to be clear and 50% are expected to be carriers. However, 25% of siblings are expected to be affected, therefore such breeding practice is discouraged.

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GENETIC REPORT

SAMPLE: BLOOD**SAMPLE TAKEN BY:** VLATKA-ANTONIJA CSIK, DVM**REQUESTED TEST:** NARCOLEPSY - LABRADOR RETRIEVER**RESULT:** CLEAR (WT/WT)**COMMENT :**

The test examines presence or absence of HCRT2 gene mutation (c.1103+5G>A) described as the cause of narcolepsy in Labrador Retrievers. The disease is characterized by daytime sleepiness, cataplexy, and striking transitions from wakefulness into rapid eye movement sleep. Narcolepsy is inherited as an autosomal recessive trait.

Regarding to the presence of tested mutation animals are classified in three groups:

- Clear (wt/wt) - mutation is not present, normal genotype
- Carrier (mut/wt) - one of two alleles carries tested mutation, disease is not clinically manifested
- Affected (mut/mut) - both alleles carry tested mutation, disease is clinically manifested

For each group different breeding strategies should be followed. Breeding of affected and carrier animals should be avoided. If particularly valuable animal is classified as affected, it should be bred only with clear animal. In such case, all first generation siblings will be carriers. If a carrier is bred with clear animal, 50% of siblings are expected to be clear. In case two carriers are bred, 25% of siblings are expected to be clear and 50% are expected to be carriers. However, 25% of siblings are expected to be affected, therefore such breeding practice is discouraged.

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GENETIC REPORT

SAMPLE: BLOOD**SAMPLE TAKEN BY:** VLATKA-ANTONIJA CSIK, DVM**REQUESTED TEST:** E LOCUS**RESULT:** E/E (E/E)**COMMENT :**

Locus E is examined for MC1R gene mutation (c.914C>T) or e allele that enables expression of other coat colour loci and causes black coat colour to change to yellow-red coat colour which is inherited autosomal recessive.

The dog carries two copies of dominant E allele enabling expression of other loci that will determine coat colour. Yellow-red coat colour (breed specific: yellow, red, cream, apricot) encoded by e allele will not be expressed. The dog is homozygote for dominant E allele and will always transfer one copy to its offspring. All of the offspring will have normal expression of other coat colour loci.

For additional information we are available on our phone during working days between 9 a.m. and 3 p.m. or e-mail.

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GENETIC REPORT

SAMPLE: BLOOD**SAMPLE TAKEN BY:** VLATKA-ANTONIJA CSIK, DVM**REQUESTED TEST:** D LOCUS**RESULT:** D/D**COMMENT :**

Locus D is examined for MLPH gene mutation (c.22G>A) or d allele that causes coat colour dilution and is inherited autosomal recessive.

The dog has two copies of dominant D allele therefore the coat colour is undiluted. The dog is homozygous for D allele and will always transfer one copy of this allele to its offspring. Due to dominance of D allele the entire offspring will express normal undiluted coat colour.

For additional information we are available on our phone during working days between 9 a.m. and 3 p.m. or e-mail.

AUTHORIZED SIGNATURE:

MARIBOR, 22.09.2020

REFERENCE NO.: 2020 - 35004**OWNER:**KAČA KACIAN
BUKEVJE 25C
HR-10411 ORLE
CROATIA**NAME/LABEL:**

GALLIVANT LIVE TO RIDE

SPECIES: DOG**BREED:** LABRADOR RETRIEVER**SEX:** MALE**MICROCHIP NO.:** 956000010780773**TATOO NO.:** NOT PROVIDED**PEDIGREE NO.:** SS14478308

GENETIC REPORT

SAMPLE: BLOOD**SAMPLE TAKEN BY:** VLATKA-ANTONIJA CSIK, DVM**REQUESTED TEST:** COAT LENGTH (FGF5)**RESULT:** N/N**COMMENT :**

The test examines presence or absence of FGF5 gene mutation (c.284G>T), which was described as the cause for increased hair growth on whole body. All dogs homozygous for this mutation have long hair. Tested FGF5 gene mutation is inherited as an autosomal recessive trait.

Regarding to the presence of tested mutation animals are classified in three groups:

- N/N - mutation is not present, the dog has short hair
- N/FGF5- the dog has short hair, but carries one copy of the variant gene which may be transmitted to offspring
- FGF5/FGF5- the dog has long hair

For each group different breeding strategies should be followed. If an animal with two mutated alleles is bred with an animal without a mutation, all siblings are expected to have short hair and all of them will carry a mutation. If an animal with one affected allele is bred with an animal without a mutation, all siblings are expected to have short hair and 50% of them will carry a mutation. If two animals with one mutated allele are bred, 25% of siblings are expected to have short hair, 50% of siblings are expected to be mutation carriers with short hair and 25% of siblings are expected to have long hair. If an animal with both mutated alleles is bred with an animal with one mutated allele, 50% of siblings are expected to be carriers with short hair and 50% of siblings will have long hair.

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GALLIVANT LIVE TO RIDE

SPECIES: DOG**BREED:** LABRADOR RETRIEVER**SEX:** MALE**MICROCHIP NO.:** 956000010780773**TATOO NO.:** NOT PROVIDED**PEDIGREE NO.:** SS14478308

GENETIC REPORT

SAMPLE: BLOOD**SAMPLE TAKEN BY:** VLATKA-ANTONIJA CSIK, DVM**REQUESTED TEST:** SKELETAL DYSPLASIA 2 (SD2) - DWARFISM**RESULT:** CLEAR (WT/WT)**COMMENT :**

The test examines presence or absence of COL11A2 gene mutation (c.143G>C) described as the cause of skeletal dysplasia 2 (SD2) in Labrador Retriever. The disease is characterized by a very subtle phenotype where mild dwarfism with short-legged phenotype is observed. COL11A2 gene defect is inherited as an autosomal recessive trait.

Regarding to the presence of tested mutation animals are classified in three groups:

- Clear (wt/wt) - mutation is not present, normal genotype
- Carrier (mut/wt) - one of two alleles carries tested mutation, disease is not clinically manifested
- Affected (mut/mut) - both alleles carry tested mutation, disease is clinically manifested

For each group different breeding strategies should be followed. Breeding of affected and carrier animals should be avoided. If particularly valuable animal is classified as affected, it should be bred only with clear animal. In such case, all first generation siblings will be carriers. If a carrier is bred with clear animal, 50% of siblings are expected to be clear. In case two carriers are bred, 25% of siblings are expected to be clear and 50% are expected to be carriers. However, 25% of siblings are expected to be affected, therefore such breeding practice is discouraged.

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CROATIA

DATE:22.09.2020

TEST REPORT NO. 204283

TEST: HEREDITARY NASAL PARAKERATOSIS (HNPK)

MUTATION: c.972 T>G in SUV39H2 gene

RESULT: CLEAR (NORMAL/NORMAL)

ANIMAL NAME: GALLIVANT LIVE TO RIDE

SPECIES: DOG

BREED: LABRADOR RETRIEVER

MICROCHIP NO.: 956000010780773

PEDIGREE NO.: SS14478308

SAMPLE TYPE: BLOOD

SAMPLE TAKEN BY: VLATKA-ANTONIJA CSIK, DVM

RESULT COMMENT:

Clear (normal/normal): tested mutation is not present, normal genotype.

Carrier (normal/mutation): one allele carries tested mutation, disease is not clinically manifested.

Affected (mutation/mutation): both alleles carry tested mutation, disease is clinically manifested.

AUTHORIZED SIGNATURE:



Results are valid for laboratory analysed samples only.

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BUKEVJE 25C
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CROATIA

NAME/LABEL:

GALLIVANT LIVE TO RIDE

SPECIES: DOG

BREED: LABRADOR RETRIEVER

SEX: MALE

MICROCHIP NO.: 956000010780773

TATOO NO.: NOT PROVIDED

PEDIGREE NO.: SS14478308

GENETIC REPORT

SAMPLE: BLOOD

SAMPLE TAKEN BY: VLATKA-ANTONIJA CSIK, DVM

REQUESTED TEST: RETINAL DYSPLASIA/OCULOSKELETAL DYSPLASIA (RD/OSD)

RESULT: CLEAR (WT/WT)

COMMENT :

The test examines presence or absence of COL9A3 gene mutation (g.49,699,847insG) described as the cause of Retinal Dysplasia/Oculoskeletal Dysplasia (RD/OSD) in Labrador retriever. The disease is characterized by short-limbed dwarfism and ocular defects.

Regarding to the presence of tested mutation animals are classified in three groups:

- Clear (wt/wt) - mutation is not present, normal genotype
- Carrier (mut/wt) - one of two alleles carries tested mutations, partial expression of RD/OSD
- Affected (mut/mut) - both alleles carry tested mutations, full expression of RD/OSD

Carrier dogs typically show symptoms of Retinal Dysplasia - partial expression of RD/OSD. Affected dogs show symptoms of dwarfism and ocular defects - full expression of RD/OSD. Heterozygote animals pass the mutation to their siblings therefore mating of two carriers should be avoided, as 25% of puppies will be affected, 50% of puppies will be carriers and 25% of puppies will be clear.

AUTHORIZED SIGNATURE:

MARIBOR, 22.09.2020

AMERICAN KENNEL CLUB · FOUNDED 1884

Certified Pedigree

Upis u HR 20354 LP
Zagreb, 03.09.2020.
Vod.rod.knjige HR: 11511752



Sire
GALLIVANT ADAMANTLY BROWN
SS07908803 (11-19) EYE7 CHLT AKC DNA
#V906809

GALLIVANT LIVE TO RIDE

SS14478308
LABRADOR RETRIEVER MALE CHLT
Microchip: 956000010780773
Date Whelped: 08/30/2019
Breeder: THOMAS FLAHERTY/FABIAN
NEGRON/PATRICK COADY

Dam
GALLIVANT'S LIKE A PRAYER
SS03962306 (01-20) CHLT



AMERICAN
KENNEL CLUB®

Erica D. Lando
Executive Secretary

The Seal of The American Kennel Club affixed hereto certifies that this pedigree was compiled from official Stud Book records on December 23, 2019.

GCH CH SHEABOURNE'S GRIZZLY ADAMS
SR69734602 (08-13) OFA24G OFEL24
CHLT AKC DNA #V711752

GCHS CH PARADOCS OBSIDIAN
SR56731701 (04-11) OFA24G OFEL24
EYE111 BLK AKC DNA #V631419

CH SHEABOURNE'S PRECIOUS ANGEL
SR22661101 (02-09) OFA33G OFEL33 BLK

CH PARADOCS HUNTERLEIGH QUINN
SR33388702 (09-09) OFA29G BLK

GCH CH DRY CREEK PARADOCS ONYX
SR27932708 (12-08) OFA24G OFEL24 BLK

CH HIGHTIDE BONA VENTURE CAPIN JACK
SN80190104 (12-04) OFA24G OFEL24 BLK AKC
DNA #V305916

TABATHA'S ANGEL
SN80230903 (04-04) OFA31G OFEL31 BLK

GCHG CH EPOCH'S MOCCASIN JOE
SR68203801 (09-12) OFA24G OFEL24 CHLT
AKC DNA #V694690

GCHB CH QUAIL CHASE BROADWAY JOE
SN4751508 (12-09) OFA28F OFEL25 CHLT AKC
DNA #V672273
EPOCH'S TREASURE
SR52301903 (09-11) OFA24E OFEL24 CHLT

GOFETCH MOLASSES COADT
SR86751601 (01-19) CHLT (CAN) AKC
DNA #V866512

BEECHCROFT JERSEY STAR
SR63677704 OFA36G OFEL36 BLK

CH BEECHCROFT STUDY'S TOP SECRET
SR28057104 (06-07) OFA25G OFEL25 BLK AKC
DNA #V510530
BEECHCROFT'S SUMMER RAIN
SR35967808 (11-10) OFA24E OFEL24 BLK AKC
DNA #V605171

CH ASTER LIBERTI LABRO HAMBURG
SR92948401 (01-17) CHLT (RUS) AKC DNA
#V788349

NIGHT SINGLE GREAT GENUS WINNERS
RKF RKF3466526 BLK (RUS)
ASTER LIBERTI LABRO GOLLANDIYA
RKF RKF3063706 CHLT (RUS)

CH GALLIVANT BLACK RUSSIAN
SR95240602 (06-18) OFA27G OFEL27
EYE29 BLK AKC DNA #V847031

CH GALLIVANT BIGPINES KENYA
SR84819301 (01-17) EYE41 BLK

GCH CH PARADOCS BELLWEATHER HEATH
SR77968402 (10-14) OFA24G OFEL24 EYE72 BLK
AKC DNA #V745732
GCH CH BIG PINE FAITH IN AFRICA
SR64542901 (01-15) OFA35E OFEL35 BLK

HEYBERN'S RHUMBA JUMPS
SR51090803 (03-13) CHLT

CH WILCARE LEISURE SUIT LARRY JH
SR25847301 (01-07) OFA24G OFEL24 CHLT AKC
DNA #V436767
CH HEYBERN'S COTILLION RN
SN90662406 (03-06) OFA35E OFEL35 BLK AKC
DNA #V290033

HEYBERN'S WAIT TILL YOUR FATHER GETS HOME
SR75573501 (09-16) OFA38G OFEL38
CHLT

CH HEDGELAWN PEMBROKE DOC HIGGINS
SR12814306 (04-06) OFA24G OFEL24 BLK AKC
DNA #V400885

HEYBERN'S NEW BALANCE
SR35093202 (03-13) OFA42F BLK AKC DNA
#V679031

CH ERINILL'S HEYBERN DEBUTANTE CD RN JH
SN56930104 (07-02) OFA29G OFEL29 BLK AKC
DNA #V53028

Thank you for ordering a Certified Pedigree from the American Kennel Club®. A pedigree contains a great deal of information, and abbreviations are an unfortunate necessity. On this sheet you will find the meanings of or links to web pages for the abbreviations used for titles, countries (if a dog was from outside the US), colors and markings; OFA certifications, and genetic clearance information. Should you have further questions, please call the AKC® Customer Service office at 919-233-9767.

Abbreviations – Used for Titles

For a complete list of titles visit

www.akc.org/events/titles/. Some titles are appended to the beginning of a dog's name, and others are appended after the dog's name. Note that titles earned in a foreign country, or under non-AKC rules and regulations, are not shown on the pedigree.

Appearing Before a Dog's Name (Prefix)

AFC	Amateur Field Champion
CCH	Bench Show Champion
CGCH	Bench Show Grand Champion
CH	Champion
CNC	Nite Champion
CSG	Bench Show Supreme Champion
CT	Champion Tracker
FC	Field Champion
FCB	Field Champion - Brace
FCGD	Field Champion - Gundog
FCLP	Field Champion - Large Pack
GCH	Grand Champion
GCHB	Grand Champion Bronze
GCHG	Grand Champion Gold
GCHP	Grand Champion Platinum
GCHS	Grand Champion Silver
HC	Herdling Champion
MACH	Master Agility Champion
OTCH	Obedience Trial Champion
PACH	Preferred Agility Champion

Appearing After a Dog's Name (Suffix)

ACT	Agility Course Test 1/2	MH	Master Hunter	OJP	Open Agility Jumper Preferred
AJP	Excellent Agility Jumper Preferred	MB	Master Bronze Jumper	OM	Obedience Master
AX	Agility Excellent	MJC	Master Century Jumper	PAX	Preferred Agility Excellent
AXJ	Excellent Agility Jumper	MJG	Master Gold Jumper	PCD	Preferred Companion Dog
AXP	Agility Excellent Preferred	MJP	Master Excellent Jumper Preferred	PCDX	Preferred Companion Dog Excellent
BCAT	Beginner Coursing Ability Test	MJPB	Master Bronze Jumper Preferred	PT	Pre-Trial Tested
BN	Beginner Novice	MJPC	Master Century Jumper Preferred	RA	Rally Advanced
CA	Coursing Ability	MJPG	Master Gold Jumper Preferred	RAE	Rally Advanced Excellent
CAA	Coursing Ability Advanced	MJPS	Master Silver Jumper Preferred	RATM	Master Barn Hunt
CAX	Coursing Ability Excellent	MMS	Master Silver Jumper	RATN	Novice Barn Hunt
CD	Companion Dog	MX	Master Agility Excellent	RATO	Open Barn Hunt
CDX	Companion Dog Excellent	MXB	Master Bronze Agility	RATS	Senior Barn Hunt
CGC	Canine Good Citizen	MXC	Master Century Agility	RE	Rally Excellent
CGCA	AKC Community Canine	MXF	Agility Master FAST Excellent	RN	Rally Novice
CGCU	AKC Urban Canine Good Citizen	MXG	Master Gold Agility	SC	Senior Courser
DJ	Dock Junior	MXJ	Master Excellent Jumper	SH	Senior Hunter
DN	Dock Novice	MXP	Master Agility Excellent Preferred	T2B	Time 2 Beat
DS	Dock Senior	MXPB	Master Bronze Agility Preferred	T2BP	Time 2 Beat Preferred
GN	Graduate Novice	MXPG	Master Gold Agility Preferred	TD	Tracking Dog
GO	Graduate Open	MXPS	Master Silver Agility Preferred	TDX	Tracking Dog Excellent
HLAdsc*	Herdling Intermediate Course A	MXS	Master Silver Agility	THD	Therapy Dog
HSAdsc*	Herdling Started Course A	NA	Novice Agility	THDA	Therapy Dog Advanced
HT	Herdling Tested	NAJ	Novice Agility Jumper	THDD	Distinguished Therapy Dog
HXAdsc*	Herdling Excellent Course A	NAP	Novice Agility Preferred	THDN	Therapy Dog Novice
IT	Instinct Tested	NF	Agility FAST Novice	THDX	Therapy Dog Excellent
JC	Junior Coucher	NFP	Agility FAST Novice Preferred	TKN	Trick Dog Novice
JE	Junior Earthdog	NJP	Novice Agility Jumper Preferred	TOX	Triple Q Excellent
JH	Junior Hunter	OA	Open Agility	UD	Utility Dog
MFB	Master Bronze FAST	OAJ	Open Agility Jumper	UDX	Utility Dog Excellent
MFP	Agility Master FAST Excellent Preferred	OAP	Open Agility Preferred	VER	Versatility
		OF	Agility FAST Open	XE	Agility FAST Excellent
		OPF	Agility FAST Open Preferred	XFP	Agility FAST Excellent Preferred

*dsc = (ducks, sheep, cattle)

GCBS CH SAMPLE DOG JE CGC RN00000001 (04-02) OFA25E OFEL22 WLD BR (NZL) AKC DNA #V299999

Dog's Registered Name

Dog's Registration Number

OFA Certification
Hips Excellent, Elbow

Country
New Zealand

Prefix Titles
Grand Champion Silver
Champion

Suffix Titles
Junior Earthdog
Canine Good Citizen

Month and year the dog's first
breeding was recorded in the
AKC Stud Book Register

Color
Withoor

DNA Profile Number

Abbreviations – OFA Certifications

OFA (age in months when certified) (grade) = Hips Certification (Grade is either "E," "G," or "F" for "excellent," "good," or "fair" respectively.) Note that the grade may be absent if the dog passed but the grade is unavailable or unpublished.

OFEL (age in months when certified) = Elbow Certification

EYE (age in months when certified) = Eye Certification. Note that this certification is only good for one calendar year.

For more information contact: Orthopedic Foundation for Animals (OFA) – www.ofa.org.

NOTE: Certification(s) may not appear in the 3rd and 4th generations due to space limitations.

Abbreviations – Genetic Clearance Information

DNA ((disease)/(from of disease)) Example: DNA (PRARCD1) = absence of rod-cone dysplasia (RCD1), the form of Progressive Retinal Atrophy (PRA) affecting Irish Setters.

Abbreviations – Colors and Markings

For a complete list of colors and markings go to www.akc.org/colors.

Abbreviations – Foreign Countries

Listed in parentheses, this indicates a foreign dog's country of origin. Note that foreign or other non-AKC domestic registry numbers are shown for a dog only if those numbers are on file with the AKC. To see a complete list of abbreviations for countries go to www.akc.org/countries.

www.akc.org

CPXGEN (4/18)

ZAVOD ZA RENDGENOLOGIJU
VETERINARSKOG FAKULTETA ZAGREB
NALAZ KUKOVA

HD A B C D E

Datum: 30.8.2020. Potpis:



ZAVOD ZA RENDGENOLOGIJU
VETERINARSKOG FAKULTETA ZAGREB
NALAZ LAKTOVA

ED 0 1 2 3

Datum: 30.8.2020. Potpis:

Handwritten signature and date



Sveučilište u Zagrebu
Veterinarski Fakultet, Tel. 2390111
Heinzelova 55, Zagreb
Odjel klinika
Zavod za rendgenologiju, ultrazvuk i fizikalnu
terapiju
Laboratorij za RTG i UZV

Vrijeme prijema: 30.8.2020. 9:41:13
Vrijeme unosa nalaza: 30.8.2020. 9:42:42

Nalaz rendgenološke i/ili UZV pretrage

RTG-5540/2020

Broj uputnice: **RTG-2241/20**

Uputio/la: **Branimir Škrlin, dr. med. vet.**

Datum: 30.8.2020.

Vlasnik: Kaća Kacian, OIB: 48297944941, 10411 ORLE, Bukevje 25c, tel. 0914455900

Pacijent: pas labrador retriever Gallivant Live To Ride, datum rođenja: 30.8.2019, ID: 956000010780773

Dijagnoza

HD-A; ED-0 (free);

Voditelj laboratorija

Prof. dr. sc. Damir Stanin

Ispisao:

Branimir Škrlin, dr. med. vet.

Predstojnik:

VETERINARSKI FAKULTET U ZAGREBU
Zavod za rendgenologiju, ultrazvučnu
dijagnostiku i fizikalnu terapiju

Prof. dr. sc. Damir Stanin

Nalaz se odnosi na dostavljeni i pretraženi uzorak te se ne smije umnažati bez pismenog odobrenja laboratorija.

Vrijeme ispisa: 30.8.2020. 9:42:54



VETERINARSKI FAKULTET SVEUCILISTA U ZAGREBU
ZAVOD ZA RENDGENOLOGIJU, ULTRAZVUČNU DIJAGNOSTIKU I FIZIKALNU TERAPIJU
FACULTY OF VETERINARY MEDICINE UNIVERSITY OF ZAGREB
DEPARTMENT OF RADIOLOGY, ULTRASOUND DIAGNOSTICS AND PHYSICAL THERAPY
HEINZLOVA 55, ZAGREB

TEL: +385 (0)1 23 90 401



POTVRDA - CERTIFICATE

DISPLAZIJA KUKOVA - HIP DYSPLASIA	X
DISPLAZIJA LAKTOVA - ELBOW DYSPLASIA	X
DATUM SNIMANJA - X-RAYS MADE ON:	30.08.2020.

BROJ PROTOKOLA - PROTOCOL NO:
5540/20

PODACI O PSU - DOG DATA

PASMINA - BREED:	Labrador Retriever
IME PSA - DOG NAME:	Gallivant Live To Ride
SPOL - SEX:	M
DATUM OSTENJENJA - DATE OF BIRTH:	30.08.2019.
BROJ RODOVNICE - PEDIGREE NUMBER:	SS14478308
BROJ MIKROČIPA - MICROCHIP NUMBER:	956000010780773
VLASNIK - OWNER:	Kača Kacian
ADRESA VLASNIKA - OWNER ADDRESS:	10411 ORLE, Bukovje 25c

Ocjena - CLASSIFICATION

KUKOVI - HIP	A	B	C	D	E
LAKTOVI - ELBOW	0	1	2	3	

Ocjenu izdao - THE EVALUATION WAS MADE BY:

Branimir Škrin DVM

Postupak je izveden prema FCI pravilniku.

The procedure has been performed according to the rules of FCI.

POTPIS - SIGNATURE

VETERINARSKI FAKULTET U ZAGREBU
Zavod za rendgenologiju, ultrazvučnu
dijagnostiku i fizikalnu terapiju



VETERINARSKA AMBULANTA
LJUBIMAC

Matije Ivanića 19, 10000 Zagreb
tel. : 01/3750-339
www.ljubimac.net
ljubimac.ambulanta@gmail.com

OFTALMOLOŠKI PREGLED - OPHTHALMIC EXAMINATION

OPIS ŽIVOTINJE / ANIMAL DESCRIPTION

IME ŽIVOTINJE / NAME : GALLIVANT LIVE TO RIDE

DATUM ROĐENJA / DATE OF BIRTH : 30. 8. 2019.

VRSTA / SPECIES : PAS SPOL / SEX : M

PASMINA / BREED : LABRADOR RETRIEVER BOJA / COLOR : ČOKOLADNA

MIKROČIP / MICROCHIP N° : 956000010780773 BROJ RODOVNIKA / PEDIGREE N° : SS 14478308

VLASNIK / OWNER

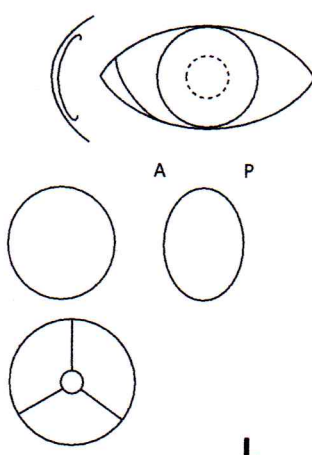
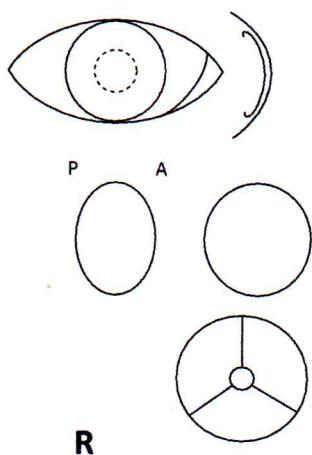
IME I PREZIME / FIRST NAME, SURNAME : KACA KACIAN

ADRESA / ADDRESS : BUZINSKI PRILAZ 17, 10010 ZAGREB, CROATIA

PREGLED OKA - EXAMINATION OF EYE

MYDRIATIC : ☒ OPHTHALMOSCOPY : DIRECT ☒ INDIRECT ☐ BIOMICROSCOPY ☒ OTHER

PARTS EXAMINED	ADNEXA	CORNEA	IRIS	LENS	VITREOUS	FUNDUS	CLINICALLY AFFECTED	CLINICALLY UNAFFECTED
CLINICALLY AFFECTED								
CLINICALLY UNAFFECTED								



COLLIE EYE ANOMALY	<input type="checkbox"/>	<input checked="" type="checkbox"/>
RD RETINAL DYSPLASIA	<input type="checkbox"/>	<input checked="" type="checkbox"/>
PPM PERSISTENT PUPILLARY MEMBRANE	<input type="checkbox"/>	<input checked="" type="checkbox"/>
PHPV PERSISTENT HYPERPLASTIC PRIMARY VITREUS	<input type="checkbox"/>	<input checked="" type="checkbox"/>
TUNICA VASCULOSA LENTIS	<input type="checkbox"/>	<input checked="" type="checkbox"/>
CATARACT	<input type="checkbox"/>	<input checked="" type="checkbox"/>
CORNEAL DYSTROPHY	<input type="checkbox"/>	<input checked="" type="checkbox"/>
PRIMARY LENS LUXATION	<input type="checkbox"/>	<input checked="" type="checkbox"/>
RETINAL DEGENERATION (PRA)	<input type="checkbox"/>	<input checked="" type="checkbox"/>
ENTROPION	<input type="checkbox"/>	<input checked="" type="checkbox"/>
ECTROPION	<input type="checkbox"/>	<input checked="" type="checkbox"/>
OPTIC NERVE HYPOPLASIA	<input type="checkbox"/>	<input checked="" type="checkbox"/>

KOMENTAR / COMMENTS :

DATUM / DATE OF EXAMINATION : 1. 9. 2020.

Vlatka-Antonija Csik
DMV

VLATKA-ANTONIJA CSIK
dr. med. vet.
OVLAŠTENI VETERINAR br. 204



HRVATSKI KINOLOŠKI SAVEZ CROATIAN KENNEL CLUB



Izložba: CACIB VARAŽDIN		
Pasma: LABRADOR RETRIVER / LABRADOR RETRIEVER	Kataloški broj: 656	
Ime psa: GALLIVANT LIVE TO RIDE	Spol: MUŽJAK / MALE	
Broj rodovnice: SS14478308	Oštenjen: 30.08.2019	Razred: MLADI / JUNIOR
Vlasnik: KACIAN KAČA	Datum izložbe: 13.09.2020	
Ime suca: AL DAGHISTANI VOJISLAV (RS)		

OPIS	OCJENA	
<i>12 m. star muškarac pravi zagriz, snažna grudi i konstitucija za cvrsto dobar lumbarski dio dobro usadeno uši dobro izbalansirano pravilne fronte dobri dlanovi</i> CROATIAN KENNEL CLUB INTERNATIONAL DOG SHOW VARAŽDIN - 13.09.2020.	PLASMAN (I - IV)	
	Odličan	X
	Vrlo dobar	
	Dobar	
	Dovoljan	
	Nedovoljan	
	Bez ocjene	
	Prvak razreda mladih	X
	CAC	
	R.CAC	
	CACIB	
	R.CACIB	
	Prvak razreda veterana	
	Najljepši mladi pas pasmine	X
	Najljepši pas suprotnog spola - BOS	
Prvak pasmine - BOB		
RAZRED ŠTENADI / NAJMLADIH		
PLASMAN (I - IV)		
Vrlo perspektivan		
Perspektivan		
Zadovoljava		
Ne zadovoljava		

Mladi klupski prvak	<input type="checkbox"/>	Klupski prvak	<input type="checkbox"/>	Klupski prvak veteran	<input type="checkbox"/>
---------------------	--------------------------	---------------	--------------------------	-----------------------	--------------------------

Potpis suca:

FCI JUDGE



Kataloški broj
Catalog Number

656

Datum
Date
13-09-2020

Sudac
Judge

Vojislav Al Daghlis
FCI JUDGE



Kataloški broj
Catalog Number

656

Datum
Date
13-09-2020

Sudac
Judge

Vojislav Al Daghlis
FCI JUDGE