PRECISE DIAGNOSTICS FOR IMPROVED CARE

Vcheck Product catalog_ver.10



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PRECISE DIAGNOSTICS FOR IMPROVED CARE

Vcheck is a multi-parametric fluorescent immunoassay analyzer providing rapid, accurate, and reliable results for quantitative, antibody titer, and infectious tests.





MULTIPLE TESTS ON A SINGLE ANALYZER

Point-Of-Care tests of various disease markers, viral antigens of infectious diseases, and antibody titer are possible with the Vcheck analyzers.



AUTO-CODING SYSTEM WITH 2D BARCODE TECHNOLOGY

All the test devices can be randomly accessible to the Vcheck analyzer without any pre-procedure. The analyzer recognizes each test device once inserted.



AUTOMATIC RECOGNITION OF HANDWRITING

A handwritten patient name or ID on the test device can be printed with the test result for user's convenience.



HIGH ACCURACY AND REPRODUCIBILITY

Strong correlation with the gold standard methods and reliability is one of the best strengths of Vcheck analyzers.

2 DIFFERENT MODELS TO MEET YOUR NEEDS



Choose the one that best suit your needs. V200 is a compact and convenient, all-in-one analyzer; V2400 has a high throughput and enables you to process large amounts of tests quickly.



RAPID, EASY TO USE AND COST EFFECTIVE

Save time, save money, and most importantly, save lives with Vcheck today.



The best way to reduce turnaround time and improve the service of your laboratory



Spe	cific	ation
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Model	: Vcheck V2400
Test capacity	: 24 tests at once / 70 tests per hour
Power	: AC/DC adaptor
Display	: 10" Color Touch Screen
Printer	: Built-in
Connectivity	: HL7 v2.6(PCD-01) / POCT1-A
Dimension	: 510 x 566 x 297 mm
Weight	: 20.0 kg

Product No.	Product Name	Storage temperature	Packing Unit
VC7403EA	V2400	15~30°C	1 EA

V200

Compact and convenient analyzer to expand your in-clinic testing



Specification	Model	: Vcheck V200
	Test capacity	: 1 test at a time
	Power	: AC/DC adaptor
	Display	: 7" Color Touch Screen
	Printer	: Built-in
	Connectivity	: HL7 v2.6(PCD-01) / POCT1-A
	Dimension	: 200 x 240 x 205 mm
	Weight	: 2.5 kg

Product No.	Product Name	Storage temperature	Packing Unit
VC7402EA	V200	15~30°C	1 EA

Feline Th Cardiac Troponin I

Quantitative marker of myocardial injury

Troponin consists of 3 subunits (troponin I, T, and C) which together function as the molecular switch of cardiomyocyte contraction. Among them, cardiac Troponin I (TnI) is a sensitive and specific circulating marker of cardiac injury for cats. Cardiac injury causes the release of TnI into the circulation, where its concentration is correlated to the severity of the damage.



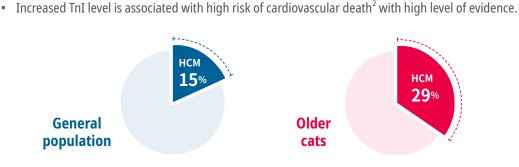
Clinical Application

Hypertrophic cardiomyopathy (HCM) is the most common heart disease and one of the 10 most common causes of death in cats. Measuring TnI concentrations can be useful in detecting subclinical HCM and predicting cardiac death in cats with HCM.

Detects HCM in apparently healthy cats

Predicts cardiac death in cats with HCM

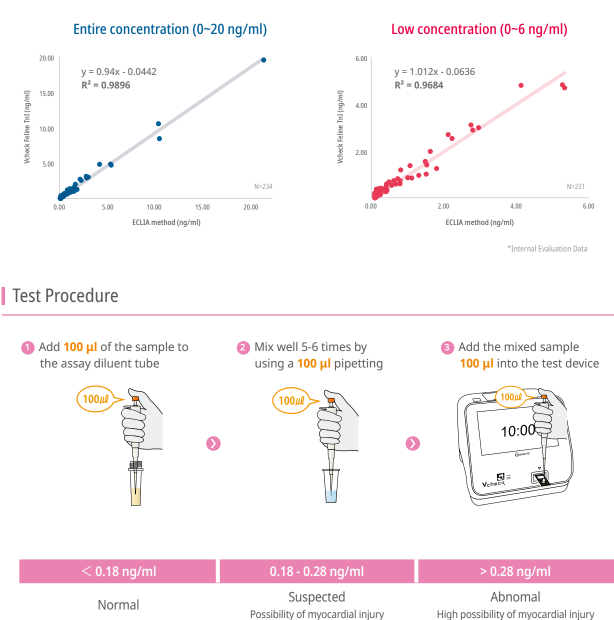
- Annual check-up, Prior to anesthesia, Cats suspected for heart diseases
- Differentiates between normal cats and cats with subclinical HCM¹



High prevalence of HCM even in apparently healthy cats³ Screen for the possibility of HCM with a cardiac biomarker, Troponin I

Reference : 1. J Vet Intern Med. 2019; May; 33(3): 1242-1250. 2. J Vet Intern Med. 2014; 28: 1731-1737. 3. J Vet Cardiol. 2015; Dec; 17 Suppl 1: S244-57.

Vcheck Feline TnI has a strong correlation (y=0.94x-0.0442, \mathbf{R}^2 =0.9896 in entire concentration; y=1.012x-0.0636, \mathbf{R}^2 =0.9684 in low concentration) with the ECLIA method from 'R' multinational healthcare company.



* TnI concentrations should not be used to either confirm or exclude primary cardiac disease without the simultaneous use of echocardiography.

Product No.	Product Name	Storage temperature	Packing Unit
VCF139DC	Vcheck Feline TnI	1~30°C	5 Tests/Kit

Canine Th Cardiac Troponin I

Quantitative marker of myocardial injury

Troponin consists of 3 subunits (troponin I, T, and C) which together function as a molecular switch of cardiomyocyte contraction. Among them, cardiac Troponin I (TnI) is a sensitive and specific circulating marker of cardiac injury for dogs. Cardiac injury causes the release of TnI into circulation, where its concentration is correlated to the severity of the damage.

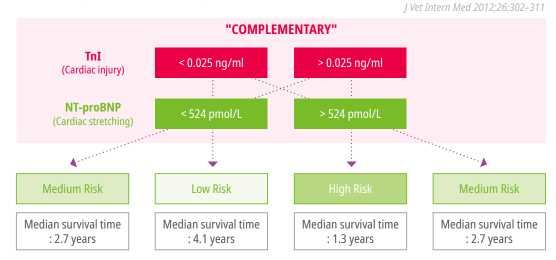


Clinical Application

Vcheck Canine TnI can provide important diagnostic and prognostic information in patients with cardiovascular or non-cardiac diseases as a cardiac injury marker of choice.

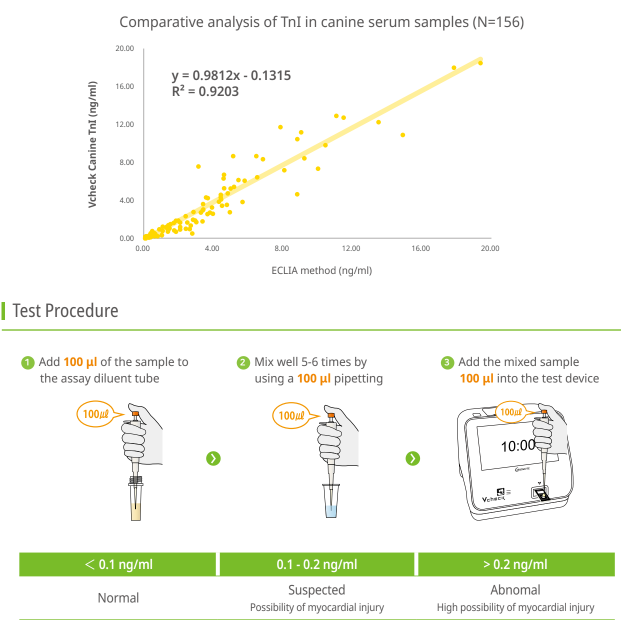
Cardiac Trauma - Detects or rules out significant blunt cardiac injury in frequent conditions

- **Primary Heart Disease** - Indicates ongoing myocyte damage in a chronic remodeling process
- Critically ill patients
 Provides prognostic information irrespective of underlying disease
- Combined measurement of TnI and NT-proBNP is prognostically superior to measuring each alone in dogs with MMVD.



Prognostic Algorithm

There is a high correlation (Y=0.9812X-0.1315, R^2 =0.92) with electrochemiluminescent immunoassay (ECLIA) from 'R' diagnostics.



* TnI concentrations should not be used to either confirm or exclude primary cardiac disease without the simultaneous use of echocardiography.

** When interpreting a slight increase of TnI in healthy dogs, biologic variability of TnI or old ages should be taken into account.

Product No.	Product Name	Storage temperature	Packing Unit
VCF137DC	Vcheck Canine TnI	1~30°C	5 Tests/Kit

Feline NT-proBNP

N-terminal pro-B type natriuretic peptide

Initial cardiac biomarker screening for heart disease in cats

NT-proBNP (N-terminal pro-B type natriuretic peptide) is cleaved from BNP which is produced by the muscle cells of the heart and increases with excessive stretching of the cells. NT-proBNP concentration reflects the degree of cardiac activation secondary to stimulus, such as stretching, allowing this marker to be used to assess the magnitude of cardiac muscle stretching.



Clinical
Application

To screen for occult heart disease

- Prior to anesthesia

- In apparently healthy cats with heart murmurs
- At risk breeds Maine Coon, Ragdoll, Birman, Persian

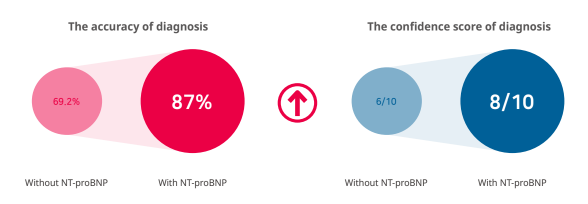
To determine Cardiac or Respiratory disease

- In cats with respiratory signs such as dyspnea, tachypnea, cough
- To differentiate cardiac and respiratory causes

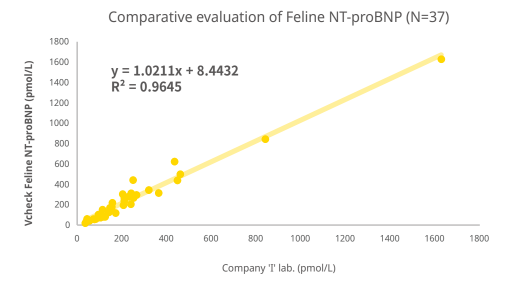
To determine the severity of heart disease

- For monitoring stabilization of CHF during hospitalization
- For predicting survival in cats with CHF

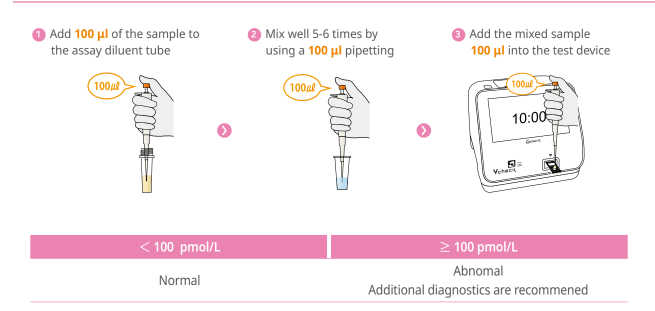




Strong correlation with company 'I' laboratories



Test Procedure



* A positive NT-proBNP test result should always be interpreted in combination and other diagnostic findings.

* In cats with respiratory signs, if the NT-proBNP is > 270 pmol/L, CHF is the most likely cause of the clinical signs.

Product No.	Product Name	Storage temperature	Packing Unit
VCF130DC	Vcheck Feline NT-proBNP	1~30°C	5 Tests/Kit

Canine NT-proBNP

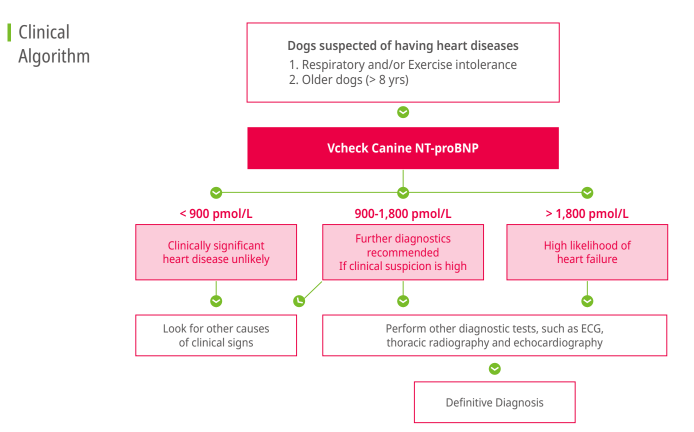
N-terminal pro-B type natriuretic peptide

New Cardiac Biomarker for dogs

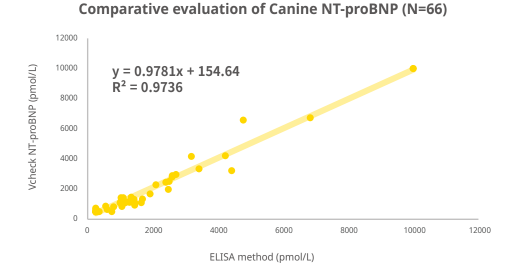
In dogs, NT-proBNP is correlated with heart size and systolic function, suggesting that the concentrations can be used to detect dogs with early disease.



- Clinical Application
- Distinguishes cardiac from respiratory disease
- Staging of Myxomatous Mitral Valve Degeneration (MMVD)
 - Detects Dilated Cardiomyopathy (DCM) in Large Breeds

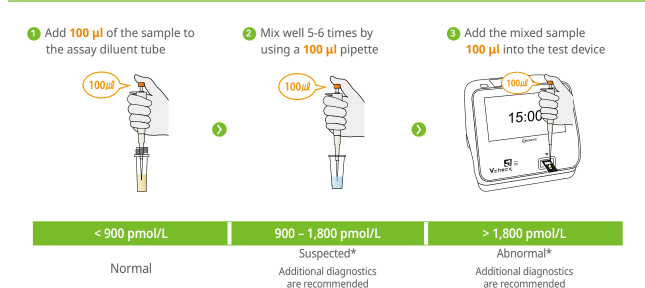


Strong correlation (R²=0.95) with an ELISA method (from company 'I' laboratories)



Test Procedure

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- * 'Abnormal' or 'Suspected' NT-proBNP test results should always be interpreted in combination and other diagnostic findings such as an echocardiogram.
- ** Concentration over 735 pmol/L in Doberman Pinschers indicates an increased risk for occult dilated cardiomyopathy.

Product No.	Product Name	Storage temperature	Packing Unit
VCF132DC	Vcheck Canine NT-proBNP	2~8°C	5 Tests/Kit

SDMA Symmetric Dimethylarginine

Biomarker for early detection of decreased renal function

SDMA is a methylated form of arginine and excreted almost exclusively by the kidneys. SDMA is a novel kidney biomarker that reflects glomerular filtration rate (GFR), increasing earlier than serum creatinine with acute kidney injury (AKI) and chronic kidney disease (CKD).

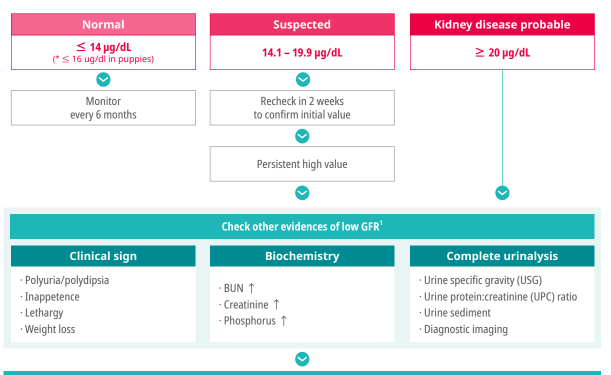
Species	Sample			0-
Dog, Cat	Serum/plasma (heparin) 100 μl		Veheck SDMA	2~8℃
Testing Time 11 min.	Measuring Range 10.0~100.0 µg/dL	• //	Sunnag	Gam

Clinical Application

- Early detection of kidney disease
- · Monitoring of patient with kidney disease
- SDMA is a novel biomarker for kidney function and more reliable than creatinine.
 But SDMA cannot replace creatinine and both are complementary to each other in diagnosing kidney dysfunction
- History, physical examination, CBC, chemistry profile including SDMA, creatinine, electrolytes, and urinalysis should be performed to evaluate kidney function

SDMA	Creatinine	Interpretation
Normal	Normal	 Normal renal function Early renal disease cannot be ruled out if SDMA and/or creatinine levels are at the upper end of the reference range.
Elevated	Normal	• Early renal disease probable
Normal	Elevated	 Not usual Possible if the lean body mass is high Further evaluation of renal function is recommended.
Elevated	Elevated	· Renal disease strongly suspected

Diagnostic Algorithm



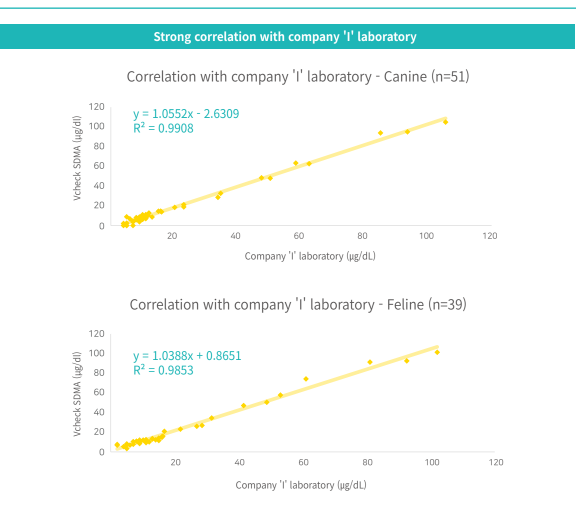
IRIS Staging of CKD (modified 2019)²

CKD Staging should be based on fasting creatinine or SDMA concentration or both measured (recommended) on at least 2 occasions in a hydrated and stable patient, preferably after 12h of fasting with free access to water.

IRES International Renal Interest Society CKD Staging	Stage 1 (No azotemia)	Stage 2 (Mild azotemia)	Stage 3 (Moderate azotemia)	Stage 4 (Severe azotemia)	
CANINE					
Creatinine mg/dL (µmol/L)	< 1.4 (< 125)	1.4 – 2.8 (125 - 250)	2.9 – 5.0 (251 - 440)	> 5.0 (> 440)	
SDMA µg/dL	< 18	18 - 35	36 - 54	> 54	
UPC ratio	< 0.2 (Non-proteinuric) 0.2–0.5 (Borderline) $>$ 0.5 (Proteinuric)				
Blood pressure	< 140 (Normotensive)	140-159 (Prehypertensive) 16	60-179 (Hypertensive) \geq 180 ((Severely hypertensive)	
FELINE					
Creatinine mg/dL (µmol/L)	< 1.6 (< 140)	1.6 – 2.8 (140 - 250)	2.9 – 5.0 (251 - 440)	> 5.0 (> 440)	
SDMA µg/dL	< 18	18 - 25	26 - 38	> 38	
UPC ratio	< 0.2 (Non-proteinuric) 0.2–0.4 (Borderline) $>$ 0.4 (Proteinuric)				
Blood pressure	$<$ 140 (Normotensive) 140-159 (Prehypertensive) 160-179 (Hypertensive) \geq 180 (Severely hypertensive)				

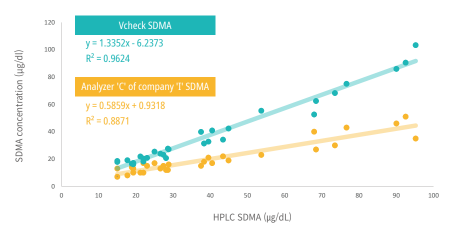
*In case of discrepancies in the interpretation of creatinine and SDMA, follow the result indicating a higher stage, and set the treatment methods accordingly.

Reference: 1. Sparkes, A. H., Caney, S., Chalhoub, S., et al. (2016) ISFM consensus guidelines on the diagnosis and management of feline chronic kidney disease. Journal of Feline Medicine and Surgery 18, 219-239 2. IRIS (International Renal Interest Society) Staging of CKD (Modified 2019).



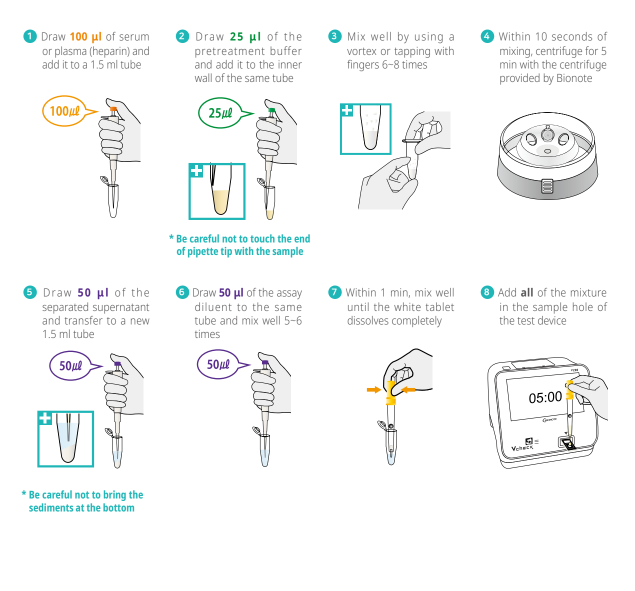
Higher correlation with the gold standard method (HPLC)

HPLC (High-Performance Liquid Chromatography): a Gold standard of SDMA



Correlation with HPLC SDMA (n=50)

Test Procedure



\leq 14 μ g/dL	14.1 – 19.9 μg/dL	≥ 20 μg/dL
Normal (≤ 16 μg/dL in puppies*)	Suspected (Check other evidence of kidney disease)	Kidney disease probable

* Mildly increased SDMA concentrations (14 – 16 μ g/dL) in puppies should be interpreted in light of the growth phase as well as other evidence of kidney disease.

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	Product No.	Product Name	Storage temperature	Packing Unit
	VCF125DD	Vcheck SDMA	2~8°C	10 Tests/Kit

D-dimer

Canine D-dimer

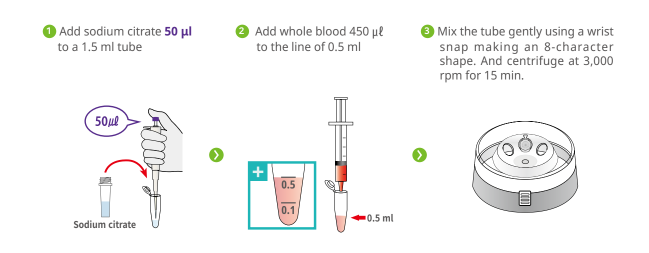
Highly sensitive marker for thromboembolism

D-dimer is a degradation fragment of cross-linked fibrin. This marker is specific for active coagulation and fibrinolysis, so increased D-dimer concentration indicates hypercoagulability. Measurement of plasma D-dimer concentration is useful for the diagnosis of systemic thrombosis, including pulmonary thromboembolism(PTE) and disseminated intravascular coagulation(DIC) in dogs.

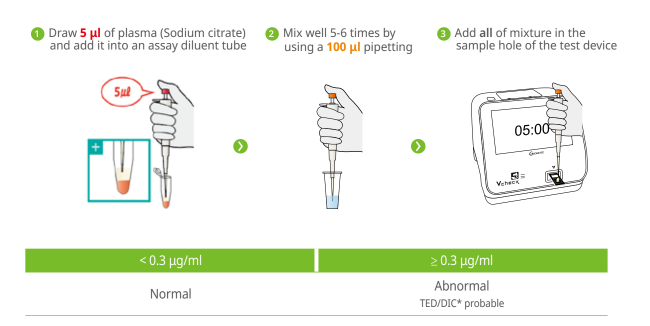
Species Dog	^{Sample} Plasma 5 μl (Sodium Citrate)	Veheck 2~8°C D-dimer
Testing Time 5 min.	Measuring Range 0.1~10 µg/ml	Contraction of the Contraction of Co

Clinical Application	 Early detection of hypercoagulability A good screening test for DIC (Disseminated intravascular Acute Thromboembolic Disease Assessment of pulmonary thromboo Monitoring of antithrombotic therap Prediction of survival prognosis after 	coagulation) embolism py
Risk Factors for Throm- boembolism	 Cancer Sepsis Pancreatitis Vascular diseases (i.e., heartworm) 	 Congestive heart failure Protein-losing disease Immune-mediated disease End/Exogenous Corticosteroids

Preparation of Sample



Test Procedure



* TED : Thromboembolic disease, DIC : Disseminated intravascular coagulation

 Product No.	Product Name	Storage temperature	Packing Unit
 VCF107DD	Vcheck D-dimer	2~8°C	10 Tests/Kit
 VCFI07DD	VCHECK D-UITIEI	2.00 C	TO TESIS/KIL

Canine CRP 2.0

C-Reactive Protein

Canine Real-Time Inflammation Marker

CRP exists at a very low concentration in healthy dogs. But it starts to increase 4 hours after inflammatory stimulation such as infection, trauma etc. If there is no further stimulation, the concentration returns to normal within a week. So CRP can be used as a real-time inflammatory marker.

Species Dog	^{Sample} Serum/Plasma (heparin) 5 μl	Veheck Canine CRP 2.0
Testing Time 5 min.	Measuring Range 10~200 mg/L	Description

Clinical	
Application	•

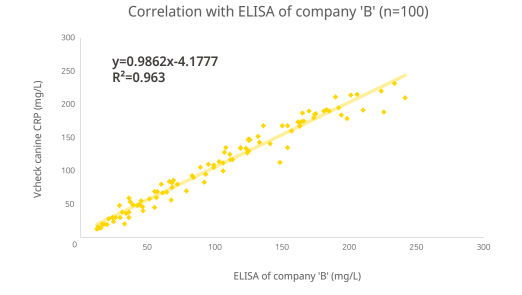
- · Earlier detection of acute inflammation : more sensitive than WBC
- Quantitative marker for inflammation : proportional to the severity of inflammation
- · Not affected by stress, steroids, NSAIDs or antibiotics unlike WBC count
- · Evaluation of treatment response, post-operative response and prognosis
- · Monitoring of recurrence of immune-mediated diseases

 CRP increases reported in dogs
 Infection / inflammation : pyometra, pneumonia, demodicosis, cystitis, periodontitis
 Tumors : hemangiosarcoma, lymphoma, nasal adenocarcinoma, cholangiocellular carcinoma

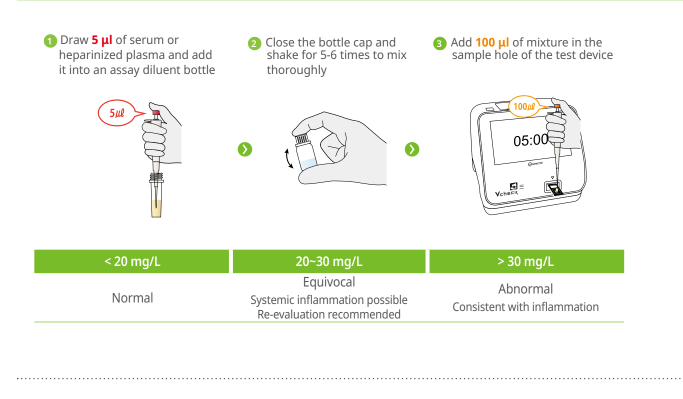
• Immune-mediated : idiopathic polyarthritis, IMHA, IMT

· Others

: acute pancreatitis, chronic hepatitis, cardiac tamponade, myelodysplastic syndrome



Test Procedure



Product No.	Product Name	Storage temperature	Packing Unit
VCF109DD	Vcheck Canine CRP 2.0	1~30°C	10 Tests/Kit

Feline SAA 3.0

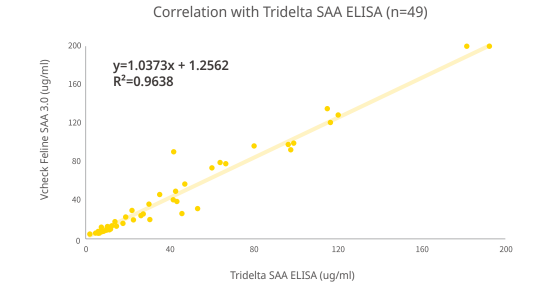
Serum Amyloid A

Feline real-time inflammation marker

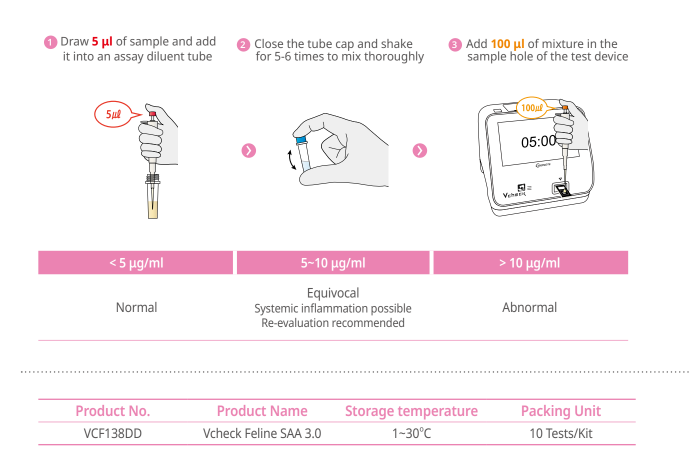
SAA exists at a very low concentration in healthy cats. But it starts to increase 4 hours after inflammatory stimulation such as infection, trauma etc. If there is no further stimulation, the concentration returns to normal within a week. So SAA can be used as a real-time inflammatory marker.

Species Cat	^{Sample} Serum/Plasma (heparin) 5 μl	Veteck Feline SAA 3.0
Testing Time 5 min.	Measuring Range 5~200 µg/ml	Comment

Clinical Application	 Differential diagnosis of diseases To evaluate severity of inflammation or infection - proportional to the severity of inflammation Differential diagnosis of FIP - SAA level highly increased compared to a feline enteric coronavirus infection Continual measurement to monitor disease progression and treatment response To evaluate recovery and complication after operations and estimate the time to hospital discharge Geriatric health checkup
SAA increases reported in cats	 Infection / inflammation acute pancreatitis, Feline Infectious Peritonitis, cholangitis, otitis media Tumors lymphoma, malignant mesothelioma Immune-mediated IMHA Others hyperthyroidism, Diabetes Mellitus, Chronic Kidney Disease



Test Procedure



CPL 2.0 Canine Pancreas-specific Lipase

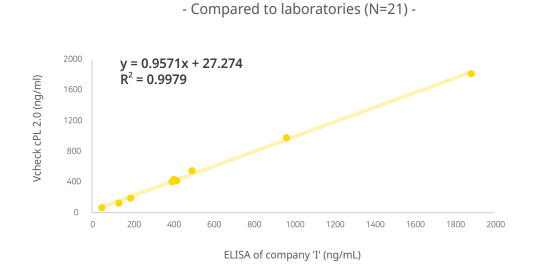
Canine pancreatitis diagnostic marker

Canine acute pancreatitis is often a life-threatening sudden and serious condition, but early diagnosis and treatment are not easy because the diagnosis is challenging and symptoms are not specific. cPL is considered to be the most specific enzyme that increases in dogs with pancreatitis and measurement of cPL is highly sensitive for a diagnosis of pancreatitis. Also cPL is little affected by other drugs or digestive disorders, thus it is useful for early diagnosis of pancreatitis. Continuous quantitative measurement also helps assess the treatment response of pancreatitis and secondary damage to pancreas caused by other digestive diseases.



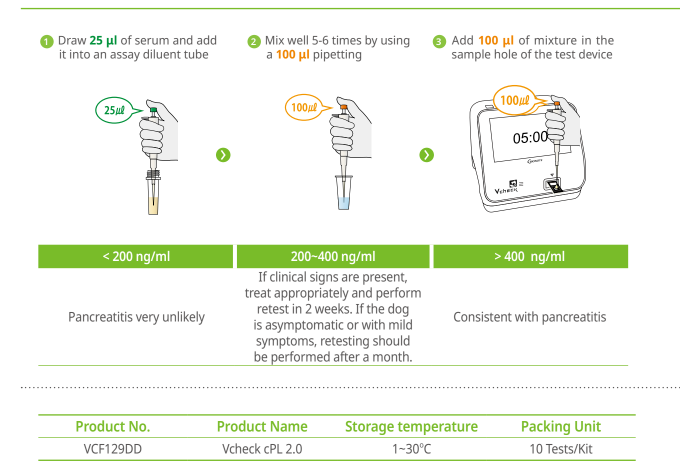
Clinical Application

- Clinical signs of acute pancreatitis: abdominal pain, anorexia, vomiting, dehydration, etc.
 - Treatment: when considering fluid therapy, analgesics, antiemetics, and antibiotics, etc.
- A specific enzyme released only from pancreas that enables early diagnosis of acute pancreatitis
- · To monitor the treatment response by continual testing
- To assess the secondary damage to pancreas in case of other digestive diseases such as cholecystitis or enteritis, etc.
- To evaluate the prognosis by measuring CRP simultaneously



Comparative Evaluation of Vcheck cPL 2.0

Test Procedure



fPL 2.0 Feline Pancreas-specific Lipase

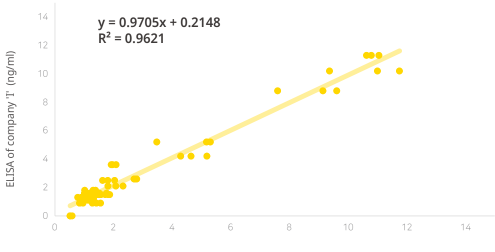
A diagnostic marker for feline pancreatitis

It is more difficult to diagnose feline pancreatitis with routine clinical chemistry tests or diagnostic imaging because the sensitivities and specificities of these diagnostic methods are low. fPL is a pancreas-specific lipase that increases in pancreatitis. Measurement of fPL has the highest sensitivity and likely the highest specificity and is the only reliable test for pancreatitis currently available in cats. Also, It helps to evaluate treatment response by continuous measurement.

Species	Sample			
Cat	Serum/Plasma (EDTA) 25 μl		Vetweck FPL 2.0	1~30°C
Testing Time	Measuring Range	- /	(111)	Comment
15 min.	1~50 ng/ml			

Clinical Application

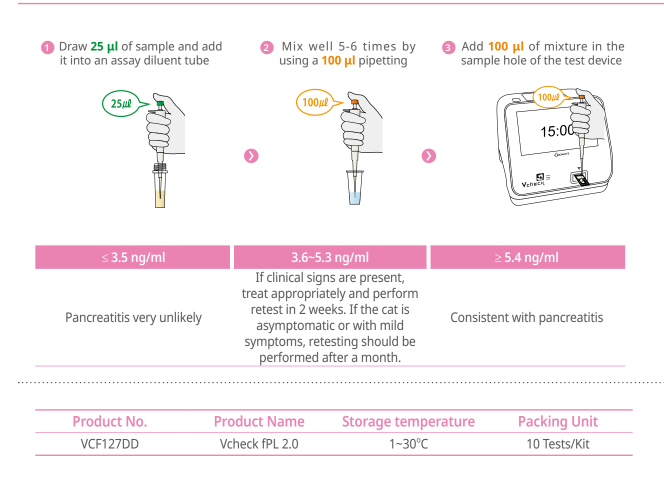
- Nonspecific clinical signs of pancreatitis: poor or absent appetite, lethargy, weight loss, dehydration, and diarrhea
 - Feline pancreas-specific lipase test correlates very well with pancreatic inflammation
 - The best overall sensitivity and specificity compared to other serum markers
 - To diagnose and rule out feline pancreatitis
 - · Time-course monitoring of pancreatitis in cats during recovery
 - To assess the secondary damage to pancreas in case of other digestive disease such as cholecystitis or enteritis, etc.



Correlation with ELISA of company 'I' (n=72)

Vcheck fPL 2.0 (ng/ml)

Test Procedure



cCortisol Canine Cortisol

Hormone Marker for hyperadrenocorticism /hypoadrenocorticism

Cortisol is secreted from the adrenal cortex and controls glucose and fat metabolism. In healthy dogs, cortisol concentration is within the normal ranges. But if there is a problem in related organs, the secretion can be excessive or insufficient. Hyperadrenocorticism (Cushing's disease) is one of the most common endocrinopathy in dogs. Measurement of cortisol level through ACTH stimulation test and LDDST, etc. can help to diagnose Cushing's disease.



Retest after 2-4

months if still

suspected

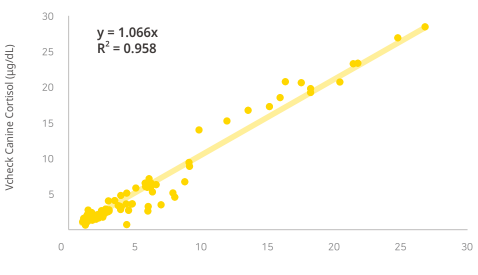
HDDST or abdominal

ultrasonography

AT

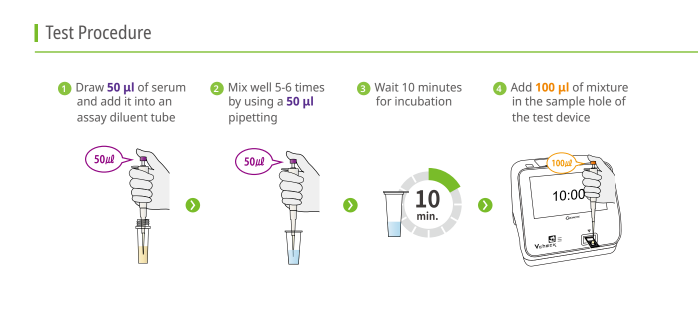
PDH

Goal of ACTH stimulation test in treatment is a post-ACTH cortisol concentration between 2-5 µg/dL (55.18 - 137.95 nmol/L).



Correlation with analyzer 'I' of company 'S' (n=50)

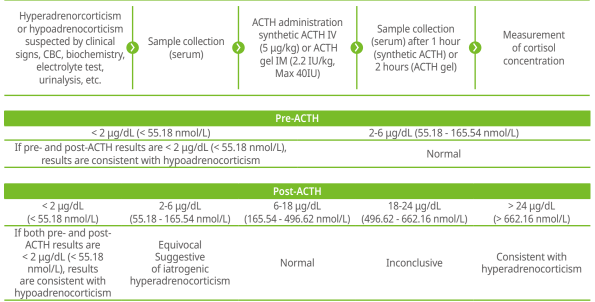
Analyzer 'I' of company 'S' (µg/dL)



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	Product No.	Product Name	Storage temperature	Packing Unit
	VCF105DD	Vcheck cCortisol	2~8°C	10 Tests/Kit

ACTH stimulation test

- ACTH stimulation test is the gold standard for diagnosis of hypoadrenocorticism, for identification of iatrogenic hyperadrenocorticism, for screening of hyperadrenocorticism and for monitoring of treatment of hyperadrenocorticism. ACTH stimulation test results do not distinguish between PDH and AT.
- Goal of ACTH stimulation test in treatment of Cushing's disease is a post-ACTH cortisol concentration between 2-5 μg/dL (55.18 137.95 nmol/L).



* 1 μg/dL is equal to 27.59 nmol/L.

Low-Dose Dexamethasone Suppression Test (LDDST)

 Results of LDDST can aid in diagnosing hyperadrenocorticism and discriminating PDH from AT in some cases

su clinic biochem	drenorcorticis spected by al signs, CBC, histry, electroly urinalysis, etc.	yte Sample c		xamethasone 01 mg/kg IV	(serui and dex	nple collection m) after 4 hours l 8 hours from kamethasone lministration	Measurement of cortisol concentration	
LDDST								
4-hour	-	1-1.4 μg/dL (27.59 - 38.63 nmol/L)	> 1.4 µg/dL (> 38.63 nmol/L) a > 50% of baselir	and (< 38.63	µg/dL nmol/L) or f baseline	> 1.4 μg/dL (> 38.63 nmol/L) or > 50% of baseline	<pre>< 1.4 µg/dL (< 38.63 nmol/L) or < 50% of baseline</pre>	
8-hour	< 1 μg/dL (< 27.59 nmol/L)	1-1.4 μg/dL (27.59 - 38.63 nmol/L)	> 1.4 µg/dL (> 38.63 nmol/L) a > 50% of baselir	and (> 38.63 r	µg/dL imol/L) and f baseline	> 1.4 µg/dL (> 38.63 nmol/L) and < 50% of baseline	> 1.4 µg/dL (> 38.63 nmol/L) and < 50% of baseline	
	Normal	Equivocal	Consistent with hyperadrenocortic	- р	DH	PDH	PDH	

* 1 μg/dL is equal to 27.59 nmol/L.

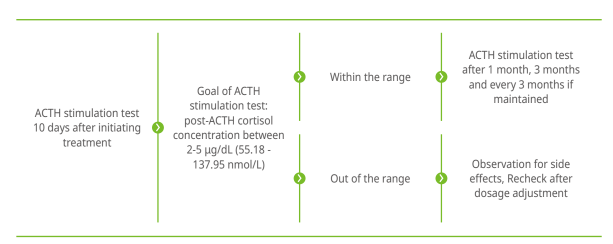
High-Dose Dexamethasone Suppression Test (HDDST)

• HDDST result can distinguish between PDH and AT in dogs with confirmed spontaneous hyperadrenocorticism. Abdominal ultrasonography can provide valuable information as well.

spont hyperadre by LDDS	th confirmed caneous enocorticism T or ACTH ation test	Sample co (serum) - E		Dexamethason 0.1 mg/kg i		Sample collection (serum) after 4 hou and 8 hours from dexamethasone administration	urs n 🜔	Measurement of cortisol concentration	
HDDST									
4-hour	<pre>< 1.4 µg/dL (< 38.63 nmol/L) ></pre>		1.5	> 1.4 µg/dL (> 38.63 nmol/L) and > 50% of baseline		< 1.4 µg/dL (< 38.63 nmol/L) or < 50% of baseline		> 1.4 µg/dL (> 38.63 nmol/L) and > 50% of baseline	
8-hour	> 1.4 μg/dL (> 38.63 nmol/L) and > 50% of baseline		< 1.4 µg/dL (< 38.63 nmol/ L) or < 50% of baseline		< 1.4 µg/dL (< 38.63 nmol/L) or < 50% of baseline		> 1.4 µg/dL (> 38.63 nmol/L) and > 50% of baseline		
	PDH		PDH		PDH		Additional testing required to differentiate PDH from ATH		

 \star 1 µg/dL is equal to 27.59 nmol/L.

Treatment Monitoring

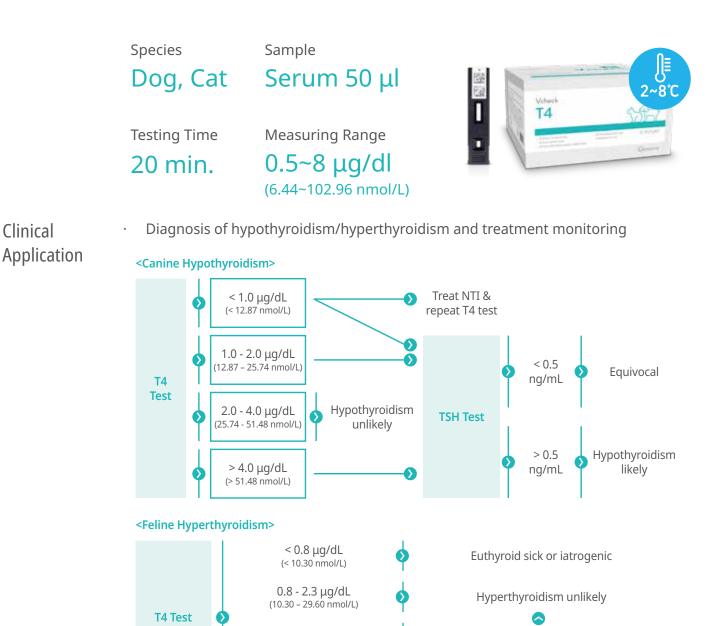


* 1 μ g/dL is equal to 27.59 nmol/L.



Hormone Marker for canine hypothyroidism and feline hyperthyroidism

T4 is a major thyroid hormone and important for normal regulation of metabolic rates and activity in various organs. Canine hypothyroidism is the common disease related to thyroid function in dogs and feline hyperthyroidism is the most common endocrine disease affecting old cats. T4 concentration level can be used to diagnose these diseases.



The prognosis of hyperthyroidism and hypothyroidism is excellent as long as they are diagnosed at early stage and the patients are treated and managed appropriately.

Additional tests including free T4 suggested

Hyperthyroidism likely

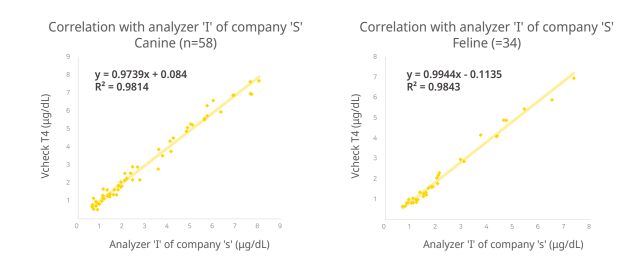
2.3 - 4.7 μg/dL

(29.60 - 60.49 nmol/L)

> 4.7 µg/dL

(> 60.49 nmol/L)

Clinical



Test Procedure

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< 1.0 µg/dL < 12.87 nmol/L)	1.0~2.0 μg/dL (12.87 – 25.74 nmol/L)	1.0~4.0 μg/dL (12.87 - 51.48 nmol/L)		> 4 μg/dL (> 51.48 nmol/L)	2.1~5.4 μg/dL (27.03 - 69.50 nmol/L)	
Low	Low normal	Norm	al	High	Therapeutic	
< 0.8 µg/dL (< 10.30 nmol/L)	0.8~4.7 μ (10.30 - 60.49		2.3~4.7 μg/dL (29.60 – 60.49 nmol/L)		> 4.7 μg/dL (> 60.49 nmol/L)	
Low	Norma	al	Gray zone		Consistent with hyperthyroidism	

Product No.	Product Name	Storage temperature	Packing Unit		
VCF106DD	Vcheck T4	2~8°C	10 Tests/Kit		

cTSH Thyroid-Stimulating Hormone

Hormone marker for canine hypothyroidism

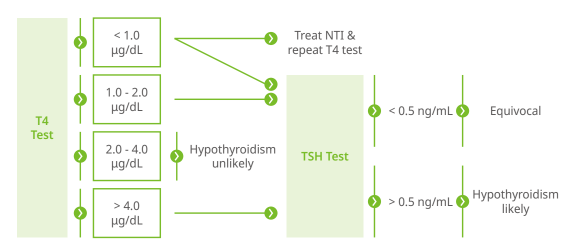
TSH is a glycoprotein produced by the anterior pituitary gland. Through its action on the thyroid gland, it plays a major role in maintaining normal circulating levels of the thyroid hormones, T4 and T3. Hypothyroidism is considered to be a common endocrine disorder in dogs, whereas hyperthyroidism in this species is rarely seen. Serum TSH is usually measured in dogs with nondiagnostic serum T4 test results, severe nonthyroidal illness, or both, and is a common component of canine thyroid panels.



Clinical Application

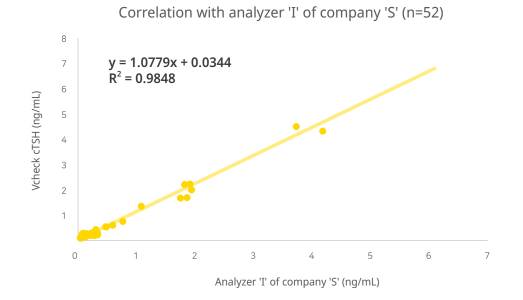
Diagnosis of canine hypothyroidism

- Most cases of canine hypothyroidism are primary in nature, involving impaired production of the thyroid hormones, T4 and T3. In this condition, elevated TSH levels are expected. Secondary or tertiary hypothyroidism, where thyroid hormone production is low as a consequence of hypothalamic or pituitary disease, is believed to account for less than 5% of canine hypothyroidism cases. And in these conditions, lowered levels of TSH would be expected.
- Serum TSH test results should always be interpreted in conjunction with results of serum T4, fT4, or both and should not be used alone in the diagnosis of hypothyroidism.

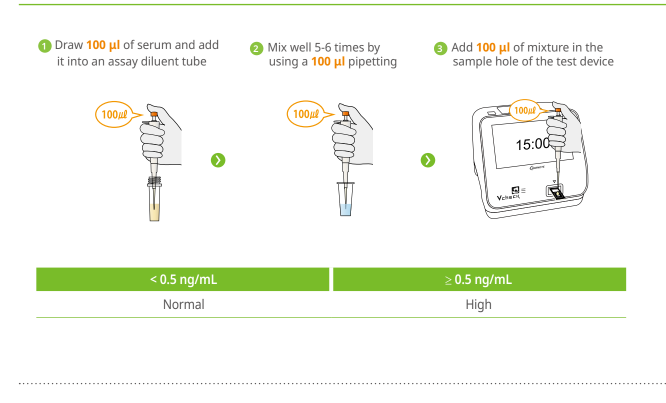


Therapeutic monitoring of canine hypothyroidism

Serum TSH concentrations are typically evaluated 4 to 6 hours after administration of levothyroxine in dogs. Ideally, the serum TSH concentration should be in the reference range.



Test Procedure



Product No.	Product Name	Storage temperature	Packing Unit
VCF118DC	Vcheck cTSH	2~8°C	5 Tests/Kit

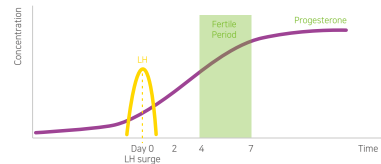
cProgesterone Canine Progesterone

Hormone

Progesterone is a steroid hormone produced primarily by the corpora luteum. Progesterone testing is used to determine when a bitch ovulates and thus when to breed. It also helps determine the timing of elective C-sections in pregnant dogs.

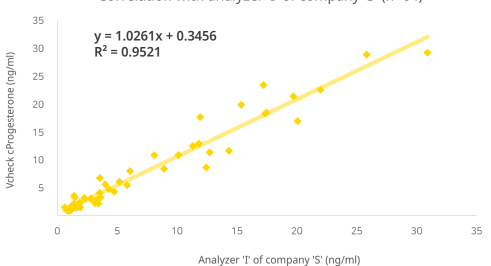


- Clinical Application
- To determine optimal breeding dates
- To predict parturition dates or time a Cesarean section
- To detect reproductive disorders such as split heats, delayed puberty, silent estrus or hypoluteidism



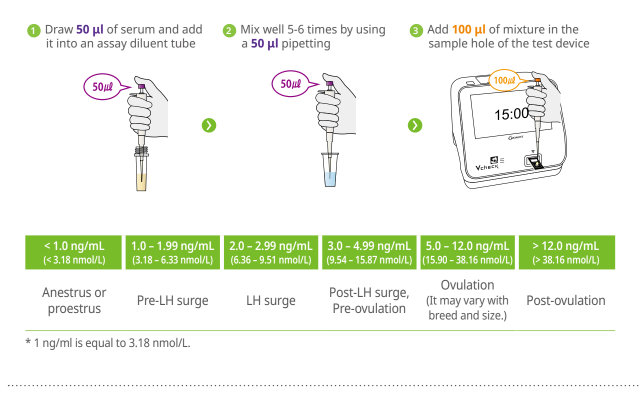
Peak fertility typically occurs 4-7 days after the LH surge (or 2-5 days after ovulation)

- Natural breeding : Ideally breed every other day while the female is showing signs of standing heat. If only 2 matings will be performed, attempt to breed 4 and 6 days after the progesterone predicted LH surge.
- Fresh or chilled semen : Ideally inseminate 3 and 5 days after the progesterone predicted LH surge.



Correlation with analyzer 'I' of company 'S' (n=64)

Test Procedure



Product No.	Product Name	Storage temperature	Packing Unit
VCF122DD	Vcheck cProgesterone	2~8°C	10 Tests/Kit



Equine real-time inflammation marker

SAA concentration increases in response to several clinical conditions in horses, and its measurement is useful for monitoring the response to treatment. Vcheck Equine SAA assay allows early detection of the presence of inflammation, monitors the post-operative effects and recovery, and serial monitoring of the response to treatment.

Species Horse	^{Sample} Serum/Plasma (Heparin) 5 μl	Veheck Equine SAA
Testing Time 5 min.	Measuring Range 10~1,000 mg/L	Comment

Early detection of the presence of inflammation

• SAA increases in the early stage of inflammation, enabling early detection of inflammation before clinical symptoms appear.

Monitoring the post-operative effects and recovery

SAA is useful for monitoring the occurrence of post-operative complications or relapse, and monitoring herd health.

Serial monitoring of the response to treatment

• SAA concentrations increase rapidly in response to inflammation and rapidly decline after the resolution of inflammation.

SAA increases reported in horses

Clinical

Application

Infection

- Bacterial: Sepsis, abscesses, strangles
- Viral: Equine herpesvirus-1 (EHV-1), Equine influenza virus (EIV)

Reproductive disease

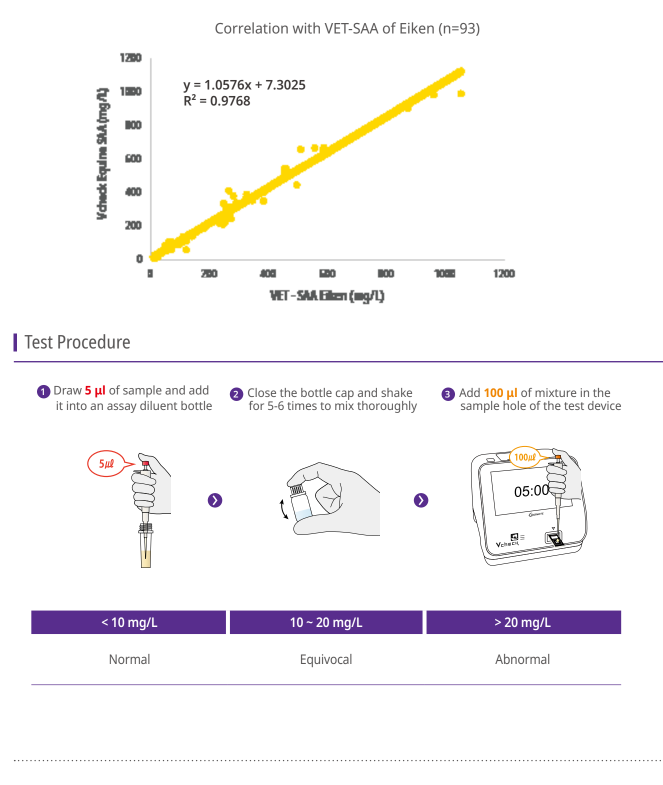
- · Septic abortion
- · Abortion of unknown aetiology

Gastrointestinal disease

- Diarrhoea and enteritis (foal)
- · Colic (adult horse)

Joint disease

- · Aseptic arthritis
- Infectious arthritis



Product No.	Product Name	Storage temperature	Packing Unit
VCF141DD	Vcheck Equine SAA	2~8°C	10 Tests/Kit



Hormone

Progesterone plays a crucial role in the maintenance of pregnancy until 120 days of gestation when the placenta becomes the main source. In addition, measuring progesterone helps find out mare's reproductive cycle and plan most effectively. Vcheck eProgesterone assay allows you to quickly analyze Equine progesterone in the field, evaluate corpus luteum in the early stages of pregnancy, and monitor progesterone during pregnancy.

Species

Horse

Sample

Serum/Plasma

(Heparin) 50 µl

Testing Time

15 min. 1~30 ng/ml

Measuring Range

(1 ng/ml is equal to 3.18 nmol/L)



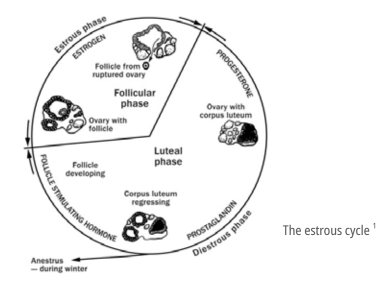
Clinical Application

In pregnant mares

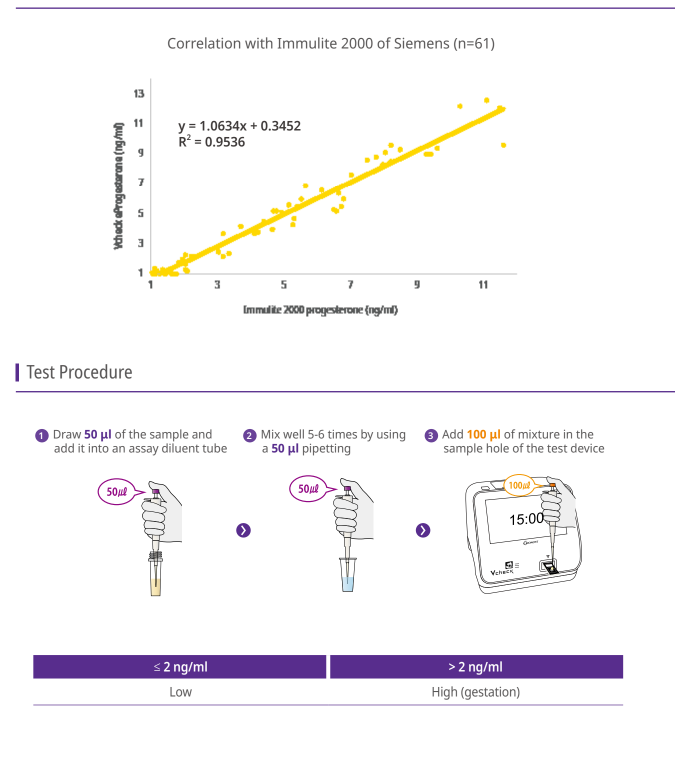
- To evaluate the maintenance of early pregnancy (Day 21~45)
- To monitor endogenous progesterone production in mares treated with supplemental hormones

In non-pregnant mares

To diagnose and treat the acyclic or irregularly cyclic mare (functional luteal tissue) (Day 21~)



Reference : 1. Anatomy, physiology and reproduction in the mare. 2010, https://www.ontario.ca/page/anatomy-physiologyand-reproduction-mare



Product No.	Product Name	Storage temperature	Packing Unit
VCF142DC	Vcheck eProgesterone	2~8°C	5 Tests/Kit

Vcheck Inf.

Infectious Test

Infectious disease test

Canine and feline infectious diseases can be diagnosed rapidly and precisely.

Specification

- Read the results within 10 minutes.
- Reading the RAPID test results visually can lead to ambiguous interpretation, especially for samples that have low levels of analyte. With Vcheck analyzer, a more precise and objective result is produced for better diagnosis.



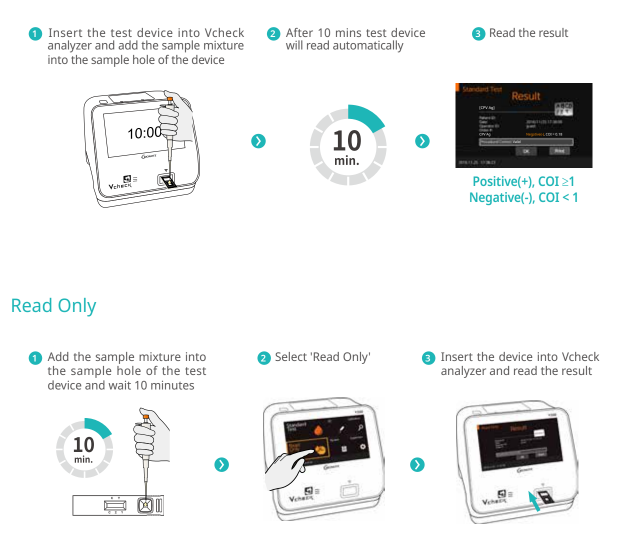
• Besides positive/negative result, COI value can help estimate the relative amount of antigen (The higher the COI value, the more antigen is present).

Products		Canine Corona Virus Antigen Vcheck CCV Ag	٥	Sample Feces
		Canine Distemper Virus Antigen Vcheck CDV Ag	٥	_{Sample} Conjunctival swab, Urine, Serum or Plasma
	Dog	Canine Parvo Virus Antigen Vcheck CPV Ag	٥	Sample Feces
		Canine Parvo/Corona Virus Antigen Vcheck CPV/CCV Ag (3 lines)	٥	Sample Feces
		Canine Heartworm Antigen Vcheck CHW Ag	٥	_{Sample} Whole blood, serum or plasma
	Cat	Feline Panleukopenia Virus Antigen Vcheck FPV Ag	•	Sample Feces

Evaluation		Vcheck CCV Ag	Vcheck CDV Ag	Vcheck CPV Ag	Vcheck FPV Ag
Data	Sensitivity	93.1 %	91.8 %	96.4 %	97 %
	Specificity	97.5 %	98.5 %	99.7 %	97.8 %

Test Procedure

Incubate and Read



Product No.	Product Name	Storage temperature	Packing Unit
VCF110DD	Vcheck CCV Ag	2~30°C	10 Tests/Kit
VCF111DD	Vcheck CDV Ag	2~30°C	10 Tests/Kit
VCF112DD	Vcheck CPV Ag	2~30°C	10 Tests/Kit
VCF114DD	Vcheck CPV/CCV Ag	2~30°C	10 Tests/Kit
VCF117DD	Vcheck CHW Ag	2~30°C	10 Tests/Kit
VCF113DD	Vcheck FPV Ag	2~30°C	10 Tests/Kit

Vcheck Ab Antibody Titer Test

Antibody Titer Test

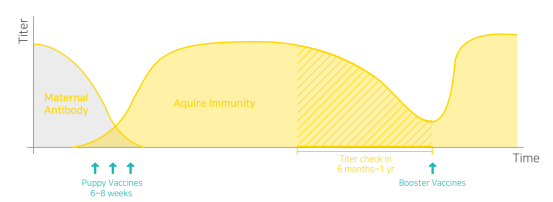
Immune status after core vaccination can be evaluated through the antibody titer test.



Clinical Application

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- To evaluate immune status after vaccination
- To optimize the primary vaccination protocol in consideration of maternallyderived antibody
- To schedule revaccination properly
- · To aid serological test and monitor treatment response



Evaluation	Vcheck CPV Ab	Compared with HI test (gold standard)	Sensitivity 100%	Specificity 85.7%
Data	Vcheck CDV Ab	Compared with VN test (gold standard)	Sensitivity 100%	Specificity 83.1%
	Vcheck CAV Ab	Compared with VN test (gold standard)	Sensitivity 87.8%	Specificity 98.2%
	Vcheck FHV Ab	Compared with VN test (gold standard)	Sensitivity 100%	Specificity 91.5%
	Vcheck FPV Ab	Compared with HI test (gold standard)	Sensitivity 100%	Specificity 95.2%
	Vcheck FCV Ab	Compared with VN test (gold standard)	Sensitivity 92.7%	Specificity 85.3%

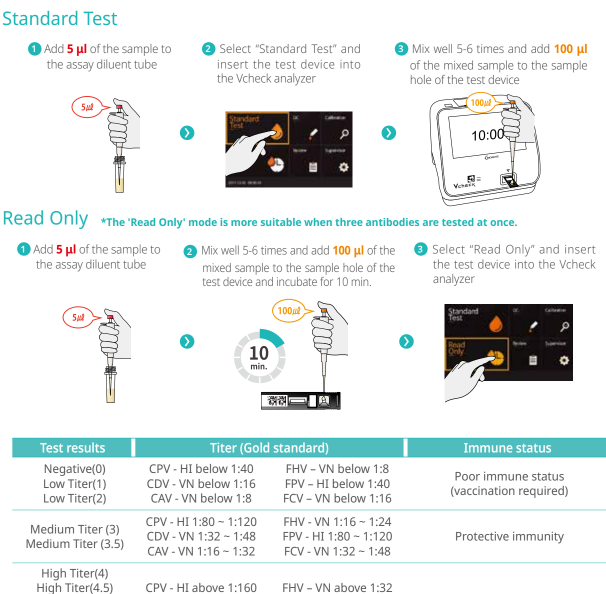
Test Procedure

High Titer(5)

High Titer(5.5) C/ High Titer(6)

CDV - VN above 1:64

CAV - VN above 1:64



FHV – VN above 1:32 FPV – HI above 1:160 FCV – VN above 1:64 Well with protective immunity

Product No.	Product Name	Storage temperature	Packing Unit
VCF115DD	Vcheck CDV Ab	2~30°C	10 Tests/Kit
VCF116DD	Vcheck CPV Ab	2~30°C	10 Tests/Kit
VCF126DD	Vcheck CAV Ab	2~30°C	10 Tests/Kit
VCF119DD	Vcheck FHV Ab	2~30°C	10 Tests/Kit
VCF120DD	Vcheck FPV Ab	2~30°C	10 Tests/Kit
VCF121DD	Vcheck FCV Ab	2~30°C	10 Tests/Kit





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