VETSCAN® HM5 Hospital Resource Guide









Welcome to the VETSCAN® HM5 Hospital Resource Guide

This guide is designed to give you everything you need to get the most out of the VETSCAN HM5 hematology system. Throughout the chapters listed, you will find links to supplemental resources to help address questions.

We hope you find this guide useful. And as always, contact Diagnostic Technical Support for further assistance at:



(888) 963-8471 (option 5)



dxsupport@zoetis.com

Need guidance on a treatment plan?

Confirm results and a path forward for complex cases with remote specialist consultations—at no additional charge for Zoetis Diagnostics customers.*



<u>ZoetisDx.com</u>

Contents

- How the VETSCAN HM5 Works
- Sample Handling
- Complete Hematology Picture
- Responsible Patient Trending
- > Histograms
- Interpretation Guide
- > Alternative Fluids
- Reference Intervals (Ranges)



How VETSCAN HM5 Works



How VETSCAN[®] HM5 Works

Sample Handling

- Patient preparation
- Sample collection
- Running a sample

Complete Hematology Picture

- Minimum database
- Automated CBC
- Blood smear
- PCV/TS

Responsible Patient Trending

Histograms

- Interpreting histograms
- Histogram examples

Interpretation Guide

- Macrothrombocytopenia
- Leukogram patterns
- Understanding anemia

Alternative Fluids

Reference Intervals (Ranges)

The VETSCAN HM5 is a fully automated, 5-part differential hematology analyzer displaying a comprehensive 22-parameter complete blood count (CBC) with cellular histograms on an easy-to-read touch screen. Its reliable performance, elegant design, ease of use and minimal maintenance with automated reminders make it the optimal veterinary hematology system.

Reliable performance for full CBC analysis



Automated CBC analysis on a wide range of species

- Flexibility to analyze different species
- Ideal for veterinary clinics, including mixed and large animal facilities
- Five-part differential for 6 species: alpaca, cat, cattle, dog, horse and llama
- Three-part differential for 9 species: ferret, goat, guinea pig, mouse, pig, primate,* rabbit, rat and sheep



Histograms to complement differentials

- Blood cell populations are graphically represented by cellular histograms
- Can easily verify differential cell counts, help identify uncommon disease processes or check sample integrity at a glance





How VETSCAN HM5 Works



How VETSCAN[®] HM5 Works

Sample Handling

- Patient preparation
- Sample collection
- Running a sample

Complete Hematology Picture

- Minimum database
- Automated CBC
- Blood smear
- PCV/TS

Responsible Patient Trending

Histograms

- Interpreting histograms
- Histogram examples

Interpretation Guide

- Macrothrombocytopenia
- Leukogram patterns
- Understanding anemia

Alternative Fluids

Reference Intervals (Ranges)

a combination of the two.

The VETSCAN HM5 uses impedance technology for the counting and categorization of cells.

Impedance technology

- In impedance counters, particles (cells) in a blood sample are suspended in an electrolyte solution and passed through an aperture that connects 2 chambers—1 containing a positive electrode and 1 containing a negative electrode
- Passage of a particle through the channel invokes a brief change in the electrical impedance between the electrodes that is proportional to the size of the particle
- Cells pass through the aperture to be classified by voltage emitted
- Measured values from impedance counters include RBC count, MCV, PLT count, WBC count and a 3- to 5-part WBC differential
- From the measured values, additional values—notably MCHC and HCT—are calculated



Most automated hematology analyzers used in veterinary medicine are either impedance counters or laser flow cytometers or contain

Hemoglobin (HGB) spectrophotometry measurement

• Spectrophotometry measures free HGB (from the lysed RBC and any amount that was already present in the plasma)



Used with permission from Alex Yartsev.





How VETSCAN HM5 Works



How VETSCAN[®] HM5 Works

Sample Handling

- Patient preparation
- Sample collection
- Running a sample

Complete Hematology Picture

- Minimum database
- Automated CBC
- Blood smear
- PCV/TS

Responsible Patient Trending

Histograms

- Interpreting histograms
- Histogram examples

Interpretation Guide

- Macrothrombocytopenia
- Leukogram patterns
- Understanding anemia

Alternative Fluids

Reference Intervals (Ranges)

VETSCAN HM5 supports 15 species

Species listed here have been validated on the VETSCAN HM5.

- Those not included are not currently supported by Zoetis Diagnostics
- The VETSCAN HM5 cannot be used for avian/reptilian species
- Primate reference intervals are available for research purposes only

The truth behind the 3- and 5-part differential

Automated CBC analyzers are generally classified as either 3-part or 5-part differential analyzers.







3-Part WBC Differential¹

Images obtained from VETSCAN IMAGYST™.

is a significant enhancement.





Sample Handling



How VETSCAN® HM5 Works

Sample Handling

Patient preparation

Sample collection

Running a sample

Complete Hematology Picture

Minimum database

Automated CBC

Blood smear

PCV/TS

Responsible Patient Trending

Histograms

Interpreting histograms

Histogram examples

Interpretation Guide

Macrothrombocytopenia

Leukogram patterns

Understanding anemia

Alternative Fluids

Reference Intervals (Ranges)

Patient preparation

These recommendations apply to all laboratories and instruments—whether point-of-care or send-out.

Before the appointment Rationale Image: Second Secon			
• Avoid feeding patients for 10 to 12 hours prior to appointment unless contraind cated • A postprandial sample may cause liperior • In horses and ruminants, fasting prior to hematology analysis is not required • A postprandial sample may cause liperior • Or sider timing of patient appointment relative to when hematologic testing will be completed • Age-related changes can lead to artifacts in sample "or hematology as maly site is not required" • Understand that certain medications may impact test results • Age-related dhanges can lead to artifacts in sample" or hematologic adverse drug events* • Avoid exercise and minimize excitement/fear prior to the appointment • Avoid exercise and minimize excitement/fear prior • Avoid exercise and minimize excitement/fear prior to the appointment • Consider the use of sodation and antianxiety medications in a set is "Physiological leukocytes s" • Minimize excitement/fear during the appointment • Or sider the use of sodation and antianxiety medications in the analyse set. • With a sick patient, anticipate that analyte results may be impacted • With a sick patient, anticipate that analyte results may single quality in may indicate the present • With a sick patient, anticipate that analyte results may be impacted • Visually inspect for dots that can falsely im courts and harminates (deal medicate the present) • With a sick patient, anticipate that analyte results may • Visually inspect for dots that can falsely im courts and harminates (deal medicate the present) •	Before the appointment		Rationale
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At the clinicRationaleWinimize excitement/fear during the appointment • Consider the use of sedation and antianxiety medications to help decrease stress for anxious animals and enable safer and gentler restraint, when appropriateCan cause: • Physiological leukocytosis³ • Transient hyperglycemia in cats°Image: Can cause: • Physiological leukocytosis 		 Avoid exercise and minimize excitement/fear prior to the appointment 	Can cause: • Physiological leukocytosis ³ • Transient hyperglycemia in cats ⁶
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		 Minimize excitement/fear during the appointment Consider the use of sedation and antianxiety medications to help decrease stress for anxious animals and enable safer and gentler restraint, when appropriate With a sick patient, anticipate that analyte results may be impacted 	 Can cause: Physiological leukocytosis³ Transient hyperglycemia in cats⁶ Visually inspect for clots that can falsely imcounts and harm the analyzer Visual assessment of the sample preanalyst highlight abnormalities (eg, hemolysis can sample quality) or may indicate the present

MCH=mean corpuscular hemoglobin; NSAID=nonsteroidal anti-inflammatory drug.



<u>Get more sample handling best practices</u> *>*



interference in hematology

n the blood BC crenation,

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npact cell

sis can indicate poor nce of disease

sed risk



Sample Handling



How VETSCAN[®] HM5 Works

Sample Handling

- Patient preparation
- Sample collection
- Running a sample

Complete Hematology Picture

- Minimum database
- Automated CBC
- Blood smear
- PCV/TS

Responsible Patient Trending

Histograms

- Interpreting histograms
- Histogram examples

Interpretation Guide

- Macrothrombocytopenia
- Leukogram patterns
- Understanding anemia

Alternative Fluids

Reference Intervals (Ranges)

Keys to successful sample collection

The quality of the sample analyzed is directly related to the quality of the result.

Avoid vein collapse when drawing samples	• Minin
Prevent hemolysis	 Use t Avoid Use n Remobility
Ensure the correct ratio of anticoagulant to blood	• Fill El • Imme (more
	 Select Ensure Alway ED Iov If it Blood fill orde
Prevent unwanted blood clotting	• Do no • For fe of a s
Do not allow samples to degrade	• Run s

Ca=calcium; EDTA=ethylenediaminetetraacetic acid; K⁺=potassium.



nize suction on the syringe, and do not draw back too quickly

- the largest vein and needle appropriate for blood collection
- d use of any needle smaller than a 23 gauge (though certain exotic species may require a smaller needle) minimal alcohol on fur/skin
- ove the needle from the syringe before dispensing into the blood tube unless using a closed vacuum collection system

DTA tube to manufacturer's sample fill line

- ediately after filling tube, cap the tube and invert 10-15 times to sufficiently mix with anticoagulant e inversions would be needed in case of 0.25 mL, 0.5 mL or 1.3 mL tubes)
- t tubes based on the testing requirements and size of patient (Microtainer[®], 1.3 mL, 3 mL, 5 mL) re tubes have not expired
- ys fill blood tubes in the correct order to avoid contamination
- TA contamination of chemistry samples may affect electrolyte results and cause a falsely v Ca and falsely high K⁺
- mproper tube-filling order occurs, the sample should be redrawn



- ot hold off the vein for more than a few seconds before venipuncture
- eline samples collected from the medial saphenous vein, a vacuum blood collection system instead syringe is recommended

samples as soon as possible after drawing







Sample Handling



How VETSCAN[®] HM5 Works

Sample Handling

- Patient preparation
- Sample collection
- Running a sample

Complete Hematology Picture

- Minimum database
- Automated CBC
- Blood smear
- PCV/TS

Responsible Patient Trending

Histograms

- Interpreting histograms
- Histogram examples

Interpretation Guide

- Macrothrombocytopenia
- Leukogram patterns
- Understanding anemia

Alternative Fluids

Reference Intervals (Ranges)

Running a sample on the VETSCAN HM5⁷

DO's

- Always use an EDTA collection tube—no other tubes are validated
- Always fill the collection tube to the manufacturer's sample fill line
- Test as soon as possible after sample collection
- Refrigerate if it will be >20 minutes until sample can be run to preserve cell morphology - Allow sample to warm to room temperature prior to running on the VETSCAN HM5
- Reinvert the tube 10 to 15 times after the sample is collected AND again just prior to running on the VETSCAN HM5
- <u>Create a blood smear</u> shortly after collection—no need to stain or evaluate slide right away • Check for blood clots before running the sample
- Select the correct species to be analyzed, as the algorithms and cell sizes differ by species - If the original species selected was incorrect (eg, if a canine sample was run as a feline), rerun the sample under the correct species

Spurious errors^{8,9}

Keep in mind that anything that spuriously affects RBC or MCV will in turn spuriously affect MCHC and HCT. Anything that spuriously affects HGB will also affect MCHC. Inaccurate results are possible if the analyzer is out of calibration.

Spurious results/errors	Cause	Prevention: Always include a blood smear to help with interpretation
HGB	Hemolysis, lipemia, icterus, Heinz bodies, severe leukocytosis	Prevent hemolysis and lipemia
MCV	Prolonged storage at room temperature, RBC swelling, autoagglutination, some regenerative anemia cases	Run sample immediately or store in refrigerator
MCHC	Hemolysis, lipemia, many Heinz bodies, storage at room temperature or false increase in HGB, marked spherocytosis	Most of these causes can be prevented with proper sampl handling techniques
PLT	PLT clumping, blood clots, macroplatelets	Prevent through proper sample handling, needle size and selection. Fill tube to manufacturer's sample fill line and in immediately after collection and just prior to running on a
RBC	Dehydration, macroplatelets	Consider patient health status and breed variations
	Blood clots, agglutination	Use proper sample handling to avoid blood clots. Never run a clotted sample

DON'Ts

- Freeze sample
- Use if tube not filled to manufacturer's sample fill line—an incorrect ratio of EDTA to blood can affect results
- Run a sample with a visible blood clot
- Run a sample straight from the refrigerator without warming to room temperature

vein nvert tube analyzer





Define your laboratory testing minimum database¹⁰

How VETSCAN[®] HM5 Works

Sample Handling

Patient preparation

Sample collection

Running a sample

Complete Hematology Picture

Minimum database

Automated CBC

Blood smear

PCV/TS

Responsible Patient Trending

Histograms

Interpreting histograms

Histogram examples

Interpretation Guide

Macrothrombocytopenia

Leukogram patterns

Understanding anemia

Alternative Fluids

Reference Intervals (Ranges)

Fecal testing is part of a minimum database when medical history and physical examination are indicative of gastrointestinal disease and as part of a preventive healthcare screening depending on patient age and lifestyle.

> Point-of-care infectious disease testing can complete a minimum database. Testing should be determined based on regional disease prevalence and patient lifestyle.

Chemistry tests assess the function and condition of various body systems. Interpretation of chemistry test results should always be associated with the analysis of other sources of data, such as: patient's signalment, history, clinical signs and results of other diagnostic testing, especially hematology and urinalysis.







How VETSCAN® HM5 Works

Sample Handling

Patient preparation

Sample collection

Running a sample

Complete Hematology Picture

Minimum database

Automated CBC

Blood smear

PCV/TS

Responsible Patient Trending

Histograms

Interpreting histograms

Histogram examples

Interpretation Guide

Macrothrombocytopenia

Leukogram patterns

Understanding anemia

Alternative Fluids

Reference Intervals (Ranges)

A complete hematologic picture includes the following CBC components:





Quantitative evaluation: automated CBC

The automated CBC is a diagnostic tool that classifies, enumerates and differentiates the different types of cells present in the peripheral blood. This **quantitative** evaluation of the blood provides different cell population counts and their associated indexes as well as graphic representations when performed on an automated analyzer.

An automated CBC also includes a differential WBC count, which is a breakdown of the amount of each subpopulation of the WBC present in the total WBC population.

Since each WBC has a very specific function, the differential count may be used to identify abnormal levels of specific WBC subpopulations and may offer diagnostic information about specific underlying health conditions.

Learn more about histograms 🗡

Blood smear (qualitative) Manual or using artificial intelligence (AI)



ZoetisDx

Clinic Name : Zoetis Demo Hospital Requesting Doctor : Dr. Smith

Owner Name : Jenna Jones Patient Name : Luna Patient ID : 12345

Hematology						
Test	Ref Range	Units	Graph	08/27/21 19:17	10/04/22	
WBC	6 - 17	10^9/I		13.79	13.93	
LYM	1 - 4.8	10^9/I		2.76	2.48	
MON	0.2 - 1.5	10^9/I		0.73	1.13	
NEU	3 - 12	10^9/I		10.03	10.14	
EOS	0 - 0.8	10^9/I		0.19	0.14	
BAS	0 - 0.4	10^9/I		0.07	0.04	
LYM%		%		20.0	17.8	
MON%		%		5.3	8.1	
NEU%		%		72.8	72.8	
EOS%		%		1.4	1.0	
BAS%		%		0.5	0.3	
RBC	5.5 - 8.5	10^12/I		8.11	7.48	
HGB	12 - 18	g/dl		16.8	15.6	
НСТ	37 - 55	%		52.56	53.71	
MCV	60 - 77	fl		65	72	
MCH	19.5 - 24.5	pg		20.7	20.9	
MCHC	31 - 39	g/dl		32.0	29.1	
RDWc	14 - 20	%		17.5	16.4	
RDWs		fl		40.6	43.0	
PLT	165 - 500	10^9/I		297	248	
MPV	3.9 - 11.1	fl		11.0	10.3	
PCT		%		0.33	0.25	
PDWc		%		40.0	40.0	
PDWs		fl		17.9	17.9	
	~ 28		12		9	









How VETSCAN[®] HM5 Works

Sample Handling

Patient preparation

Sample collection

Running a sample

Complete Hematology Picture

Minimum database

Automated CBC

Blood smear

PCV/TS

Responsible Patient Trending

Histograms

Interpreting histograms

Histogram examples

Interpretation Guide

Macrothrombocytopenia

Leukogram patterns

Understanding anemia

Alternative Fluids

Reference Intervals (Ranges)

Qualitative evaluation: blood smear

A blood smear is a **qualitative** part of the comprehensive CBC that is used to confirm results, assure quality, and may provide additional insights to guide diagnosis and treatment.¹¹⁻¹⁴ Microscopic examination of a blood smear is an essential part of the hematologic picture, as it can provide vital diagnostic information that is not identified on the automated CBC.

clinical instances⁸:

- Anemia (low RBC count)
- Thrombocytopenia (low PLT count)
- Neutrophilia or neutropenia (verify count and examine cells)
- Lymphocytosis
- Severe illness (eg, sepsis)
- Suspicion of parasites
- When certain warning flags are present on the automated CBC report

A blood smear evaluation should not be utilized as a replacement for an automated CBC, as automated analyzers count thousands of cells for more precise and accurate data than manual cell counting. Machines must be properly maintained for consistent precision and accuracy.¹⁵





Ideally, a blood smear evaluation should always be performed with every CBC, but it is vital that one be performed in the following



A **blood smear** enables the veterinarian to confirm results and assure quality and may provide additional insights to guide diagnosis and treatment.¹¹⁻¹⁴



Figure: VETSCAN IMAGYST[™] AI Blood Smear sample report









How VETSCAN[®] HM5 Works

Sample Handling

Patient preparation

Sample collection

Running a sample

Complete Hematology Picture

Minimum database

Automated CBC

Blood smear

PCV/TS

Responsible Patient Trending

Histograms

Interpreting histograms

Histogram examples

Interpretation Guide

Macrothrombocytopenia

Leukogram patterns

Understanding anemia

Alternative Fluids

Reference Intervals (Ranges)

Further quantitative evaluation: PCV/TS

PCV is the direct centrifugal measurement of the percentage of the blood that consists of RBC. It is an accurate measurement with minimal inherent error (+/-1%).¹⁶

HCT is a calculated value of the percentage of the blood that consists of RBC: HCT (%)=(RBC/ μ L) x MCV (fL/10).¹⁷ Unlike PCV, the potential for error is greater with the HCT method because it is subject to MCV variation that can occur with certain hematologic conditions (eg, agglutination) and/or improper sample handling (eg, excess EDTA, hemolysis or inadequate mixing).¹⁸

To verify that HCT was not affected by artifactual errors, perform a PCV measurement¹⁹

HCT (%) can also be estimated with this calculation, using the HGB result from the automated CBC report:

HGB x 3 (SI units: HGB x 0.3)

Zoetis Diagnostics delivers a complete hematology solution A streamlined workflow makes it easy to get to the harder diagnoses





Start with VETSCAN HM5 for quantitative CBC results

Al=artificial intelligence; HCT=hematocrit; MCV=mean cell volume; PCV=packed cell volume; TS=total solids. *If abnormalities are observed, expert review via digital image transfer is available. This can be done within the VETSCAN IMAGYST system. Additional costs may apply. ⁺Available in select markets.

Al Blood Smear





ZoetisDx⁺



Responsible Patient Trending



How VETSCAN[®] HM5 Works

Sample Handling

- Patient preparation
- Sample collection
- Running a sample

Complete Hematology Picture

- Minimum database
- Automated CBC
- Blood smear
- PCV/TS

Responsible Patient Trending

Histograms

- Interpreting histograms
- Histogram examples

Interpretation Guide

- Macrothrombocytopenia
- Leukogram patterns
- Understanding anemia

Alternative Fluids

Reference Intervals (Ranges)

Responsible Patient Trending—**why perform**?

Due to biological variations, the best reference values are a pet's own diagnostic values over time, encompassing breed, age, sex and individual variation

- on published reference values for chemistry and hematology²⁰

Senior patients

- The common occurrence of physical exam and laboratory abnormalities in apparently healthy senior dogs and cats emphasizes the need for regular health screening, including regular laboratory testing^{12,13}
- Visit/exam frequency and testing recommendations should be based on patient's age, breed and lifestyle
- Senior and geriatric dogs and cats should be examined at least semiannually to allow for earlier intervention of chronic disease
- Regular testing at geriatric equine annual examinations assesses overall health and may display any early signs of potentially serious disease, such as liver and kidney dysfunction or onset of metabolic disease²¹

• Most reference intervals represent results expected for 95% (19 of 20) of a healthy population, and therefore 1 of 20 healthy animals is expected to have a measured value outside of the reference interval¹¹

• For these reasons, individual patient trending is more sensitive and better at detecting pathologic changes than reliance





Responsible Patient Trending



How VETSCAN[®] HM5 Works

Sample Handling

- Patient preparation
- Sample collection
- Running a sample

Complete Hematology Picture

- Minimum database
- Automated CBC
- Blood smear
- PCV/TS

Responsible Patient Trending

Histograms

- Interpreting histograms
- Histogram examples

Interpretation Guide

- Macrothrombocytopenia
- Leukogram patterns
- Understanding anemia

Alternative Fluids

Reference Intervals (Ranges)

Keys to patient trending success

The best practice is to monitor a patient on the same analyzer using the same analytical methods

Whenever comparing or trending analyte results, it is important to trend using best practices and responsible trending to have a consistent comparison. This practice includes:

- Using the same analyzer every time, when possible
- Performing the test in the same way (sample type, number of hours pre- or posttreatment, fed or fasted state, etc)
- Keeping in mind that different assays and instruments have reference intervals that may differ among analyzers and/or laboratories
- Performing a quality check or verifying with a different test, methodology or laboratory if a value does not match the clinical picture

What is responsible trending?

Responsible Trending^M, available only on the ZoetisDx online platform, shows chronological test analyte results as a sequence of graphs. This visual format provides a clear story of each patient's trends in test results over time—with results from different analyzers displayed together but always relative to each analyte's reference interval on its respective analyzer.



Note: It is imperative when comparing results between different analyzers or labs to interpret the raw value with respect to the reference interval provided and not the raw number due to inherent methodology differences.





Histograms



How VETSCAN[®] HM5 Works

Sample Handling

Patient preparation

Sample collection

Running a sample

Complete Hematology Picture

Minimum database

Automated CBC

Blood smear

PCV/TS

Responsible Patient Trending

Histograms

Interpreting histograms

Histogram examples

Interpretation Guide

Macrothrombocytopenia

Leukogram patterns

Understanding anemia

Alternative Fluids

Reference Intervals (Ranges)

Introduction to histograms

Histograms are graphic representations of cell distribution and cell counts²²

Histograms are the graphs that display on your results screen or appear on the printed hematology report, depending on the practice management software used. They are easily understood and can help to:

Verify differential cell counts



The graphical display of data provides information not available in the automated CBC and allows verification of the CBC's numerical results

Histogram interpretation requires a thorough review of PLT, RBC and WBC curves for abnormalities

- Perform a blood smear for all patients who have any cell outside of the reference interval (range)
- Remember that a single hemogram is a snapshot in time and changes can occur rapidly
- response and response to treatment
- Always interpret results in conjunction with the clinical evaluation of the patient

Perform quality control check



Show sample integrity and potential abnormalities, so users can quickly decide whether results are meaningful or whether they need to draw a new sample

Identify uncommon disease processes



Identify uncommon disease processes that can affect cell morphology and distribution. These need to be confirmed visually with a blood smear

• Monitoring hemogram results frequently (every 12 to 24 hours) can be helpful in determining the course of an inflammatory







How VETSCAN[®] HM5 Works

Sample Handling

- Patient preparation
- Sample collection
- Running a sample

Complete Hematology Picture

- Minimum database
- Automated CBC
- Blood smear
- PCV/TS

Responsible Patient Trending

Histograms

Interpreting histograms

Histogram examples

Interpretation Guide

- Macrothrombocytopenia
- Leukogram patterns
- Understanding anemia

Alternative Fluids

Reference Intervals (Ranges)

Interpreting histograms

Histogram interpretation requires a thorough review of PLT, RBC and WBC curves for abnormalities.

Elements of a histogram²³













How VETSCAN® HM5 Works

Sample Handling

- Patient preparation
- Sample collection
- Running a sample

Complete Hematology Picture

- Minimum database
- Automated CBC
- Blood smear
- PCV/TS

Responsible Patient Trending

Histograms

Interpreting histograms

Histogram examples

Interpretation Guide

- Macrothrombocytopenia
- Leukogram patterns
- Understanding anemia

Alternative Fluids

Reference Intervals (Ranges)

Normal histograms

WBC Histogram





The EOS histogram varies and may be symmetrical or asymmetrical, jagged or smooth. Debris (lysed cells) from the EOS counting step may be seen to the left of the discriminator. It is not uncommon for the EOS discriminator to appear on the edge of the EOS peak.

- The eosinophils are counted separately from the other WBC types and are thus shown in a separate histogram
- Interpreting EOS by focusing on the numerical VETSCAN® HM5 data is recommended

RDWc=red cell distribution width (percent).



The peaks in the **WBC histogram**, separated by discriminators, correspond with LYM, MON and GRA. Debris (lysed RBC) from the WBC counting step may be seen to the left of the first discriminator.

LYM peak (left) is seen to the right of the first discriminator. In canines, the LYM peak starts on the low-to-mid portion along the y-axis, as shown, indicating lower populations of this cell type. In normal felines, peak starts on the mid portion of the y-axis due to higher relative number of LYM in cats

MON peak (center) is seen to the right of the second discriminator and is typically shorter due to a smaller population relative to other WBC

GRA peak (right) is seen to the right of the third discriminator and is predominantly composed of neutrophils. It is the tallest and widest peak, indicating it is the most numerous WBC population



The **RBC histogram** in normal dogs and cats should present as an almost symmetrical, bell-shaped curve. The PLT peak can be seen to the left of the discriminator.

- The width of the curve relates to the RDWc
- An increased RDWc would show on the RBC histogram as a wider peak and mean that some of the RBC are either larger and/or smaller than normal
- RDWc measures anisocytosis, or RBC size variation

PLT Histogram



The **PLT histogram** begins with a sharp increase to a peak and tapers downward as cell size increases. This indicates that most PLT are small, with fewer large PLT. The RBC peak can be seen starting to the right of the discriminator.

- In felines and equines, the histogram tapers downward more slowly, indicating more large cells, commonly associated with mild PLT clumping
- The reported PLT number is the minimum number of free PLT counted by the analyzer
- The PLT histogram should be evaluated whenever the PLT count is low and/or lymphocytes are elevated. A blood smear to confirm a low PLT count is also recommended





How VETSCAN[®] HM5 Works

Sample Handling

Patient preparation

Sample collection

Running a sample

Complete Hematology Picture

Minimum database

Automated CBC

Blood smear

PCV/TS

Responsible Patient Trending

Histograms

Interpreting histograms

Histogram examples

Interpretation Guide

Macrothrombocytopenia

Leukogram patterns

Understanding anemia

Alternative Fluids

Reference Intervals (Ranges)

Histogram examples

PLT clumping and anemia²⁶ Feline

High WBC	
could be	WBC
nucleated cells	LYM
or clumped PLT	MON
Elevated NEU	
	BAS
	LYM%
	MON%
	NEU%
	EOS%
	BAS%
	RBC
	HGB 🚺
	НСТ 🚺
	MCV
	MCH
	MCHC
In felines.	RDWc
RDWc >24%	RDWs
indicates	
increased	
variation of	
RBC size	FCI
(anisocytosis)	PDWC
	PDWs



Always confirm these results with a blood smear



Flattened PLT curve with an upward trend and low PLT count

Tail on RBC curve indicates higher amount of larger, immature RBC

Long tail on WBC curve







How VETSCAN[®] HM5 Works

Sample Handling

- Patient preparation
- Sample collection
- Running a sample

Complete Hematology Picture

- Minimum database
- Automated CBC
- Blood smear
- PCV/TS

Responsible Patient Trending

Histograms

- Interpreting histograms
- Histogram examples

Interpretation Guide

- Macrothrombocytopenia
- Leukogram patterns
- Understanding anemia

Alternative Fluids

Reference Intervals (Ranges)

Histogram examples



Classic leukemia²⁶ Canine

Very high total WBC

WBC			
LYM			
MON			
NEU			
EOS		\mathbf{i}	
BAS		\diamond	
LYM%		\bigcirc	
MON%		\diamond	
NEU%		\bigcirc	
EOS%		$\mathbf{\Diamond}$	
BAS%		\bigcirc	
RBC	\bigcirc		
HGB		\bigcirc	
НСТ		\diamond	
MCV		\bigcirc	
МСН			0
МСНС			\Diamond
RDWc		\mathbf{b}	
RDWs			
PIT			
PCI			
PDWc			
PDWs			

M warning flag may appear (refer to manual)



Always confirm these results with a blood smear and potential pathology review







Only one WBC peak

Clumping vs truly low PLT (ambiguous PLT curve)





How VETSCAN[®] HM5 Works

Sample Handling

- Patient preparation
- Sample collection
- Running a sample

Complete Hematology Picture

- Minimum database
- Automated CBC
- Blood smear
- PCV/TS

Responsible Patient Trending

Histograms

- Interpreting histograms
- Histogram examples

Interpretation Guide

- Macrothrombocytopenia
- Leukogram patterns
- Understanding anemia

Alternative Fluids

Reference Intervals (Ranges)

Histogram examples



Microcytic anemia²⁵ Canine





		WBC		\mathbf{O}
normall	ΥM	LYM	\diamond	
		MON		\bigcirc
Hiab -		NEU		\diamond
NFU +/-	MON	EOS	\diamond	
		BAS	\Diamond	
Normal F	ormal EAC	LYM%		
INUITIAI L				







Flat LYM peak

RDWc on the upper limit

narrowed RBC histogram







How VETSCAN[®] HM5 Works

Sample Handling

- Patient preparation
- Sample collection
- Running a sample

Complete Hematology Picture

- Minimum database
- Automated CBC
- Blood smear
- PCV/TS

Responsible Patient Trending

Histograms

- Interpreting histograms
- Histogram examples

Interpretation Guide

- Macrothrombocytopenia
- Leukogram patterns
- Understanding anemia

Alternative Fluids

Reference Intervals (Ranges)

Inherited macrothrombocytopenia^{14,27}

Inherited macrothrombocytopenia is due to a mutation in the gene encoding betal-tubulin. This mutation has been identified in approximately 90% of Cavalier King Charles spaniels (CKCSs), as well as other breeds, including bichons frises, boxers, cairn terriers, Chihuahuas, cocker spaniels, English toy spaniels, Havanese, Jack Russell terriers, labradoodles, Labrador retrievers, Maltese, mixed breeds, Norfolk terriers, poodles and shih tzu.

Inherited macrothrombocytopenia on the VETSCAN® HM5^{28,29}

Macrothrombocytes will not break into fragments when exposed to lyse solution and may cause inaccurate counts and/or instrument errors, particularly in analyzers that use impedance technology as the primary measurement method. When such samples are analyzed on the VETSCAN HM5, an "L" flag will often be displayed. While an "L" flag could suggest a potential lyse-resistant RBC anomaly or a problem with lyse delivery, in this case, the analyzer is reporting macroplatelets as nonlysed lymphocytes.



From chapter 7, Evaluation of hemostasis: coagulation and platelet disorders. Figure 7-26. In: Harvey JW, ed. Veterinary Hematology: A Diagnostic Guide and Color Atlas. 1st ed. Elsevier Inc.; 2012:211.

MPV=mean platelet volume.



	PID SID Needle	2 00057 -2mm	03/24/2016 06: Type Warning
	Necule	211111	warning
A CRC from the	WBC	16.17 10 ⁹ /	+ 1 - • :
A CDC HOIT THE	MON	0.08 10 ⁹ /	
VEISCAN HMISION d	NEU	7.83 10 ⁹ /	
patient with innerited	FOS	0.38 10 ⁹ /	
macrothrombocytopenia	BAS	0.00 10 ⁹ /	57 89
will typically show an	LYM%	48.6 %	
elevated LYM and a	MON%	0.5 %	
warning "L" flag	NEU%	48.5 %	
	EOS%	2.4 %	
	BAS%	0.0 %	
	RBC	5.96 10 ¹² /	79
	HGB	14.2 g/dl	
There will not be a	НСТ	37.73 %	
significant PLT neak	MCV	63 fl	
and the DLT count	МСН	23.8 pg	
will be reported	МСНС	37.6 g/dl	
	RDWc	15.4 %	14
as very low	RDWs	39.8 fl	
	PLT	9 10 ⁹ /	
The MPV may also be	MPV	6.9 fl	-
reported as normal	PCT	0.01 %	
	PDWc	34.1 %	
	PDWs	9.5 fl	14

Note: Not every CKCS has macrothrombocytopenia. Those individuals with normal-size PLT will not show the features mentioned above.









How VETSCAN[®] HM5 Works

Sample Handling

- Patient preparation
- Sample collection
- Running a sample

Complete Hematology Picture

Minimum database

Automated CBC

Blood smear

PCV/TS

Responsible Patient Trending

Histograms

Interpreting histograms

Histogram examples

Interpretation Guide

Macrothrombocytopenia

Leukogram patterns

Understanding anemia

Alternative Fluids

Reference Intervals (Ranges)

Understanding classic leukogram patterns

Changes in total and differential leukocyte counts are usually grouped into patterns that facilitate interpretation. These patterns are:

		NEU	Left Shift	Toxic Δ^*	LYM	MON
Stress leukogram	 A result of cortisol released by the adrenal gland Occurs due to a wide range of processes Systemic illness; metabolic disturbance; pain Mimicked by corticosteroid therapy 	ſ	No	No	¥	canine > feline
Physiological leukocytosis	 A result of epinephrine or norepinephrine release Also called a flight-or-fight response Most often seen in cats (of any age) and possibly in the young of other species Usually transient and generally resolves about 30 minutes after the patient relaxes 	t	No	No	f (mostly feline)	Normal
	 Represents the balance between tissue demand and bone marrow supply May vary depending on source and severity of inflammation and timing of sample collection NEU numbers may vary from severely depressed to markedly increased A left shift indicates the presence of immature NEU 			Mild/Chronic	: Inflammat	tion
		Ť	+/-	No	Normal or ↓	(chronic)
		Acute Inflammation				
Inflammatory leukogram		Ť	Ť	Frequent	↓	Normal or
	— Usually, but not always, indicates an inflammatory leukogram	Overwhelming Inflammation				
	 Inflammation is possible in patients without an inflammatory leukogram 		↑ to ↑↑	Present	↓ ↓	No
Leukemoid reaction	 Characterized by a marked neutrophilic leukocytosis (>50,000 cells/µL) with a concurrent, orderly left shift Toxic changes may or may not be present Resembles granulocytic leukemia but is caused by another process Also referred to as extreme neutrophilic (granulocytic) leukocytosis 		+/-	Occasional	Normal or ↓	Normal or

 $^{*}\Delta$ =change.







Inflammation



+/-

Unusual

Hopefully





?





How VETSCAN® HM5 Works

Sample Handling

- Patient preparation
- Sample collection
- Running a sample

Complete Hematology Picture

- Minimum database
- Automated CBC
- Blood smear
- PCV/TS

Responsible Patient Trending

Histograms

- Interpreting histograms
- Histogram examples

Interpretation Guide

- Macrothrombocytopenia
- Leukogram patterns
- Understanding anemia

Alternative Fluids

Reference Intervals (Ranges)

Understanding anemia

Anemia is one of the most common hematologic abnormalities encountered in veterinary clinical practice. It is the manifestation of an underlying disorder, like a fever, and not a diagnosis. It can be a primary sign of disease (eg, hemorrhage or immune-mediated hemolytic anemia) or a marker of underlying disease (eg, cancer or chronic kidney disease). Therefore, even mild, asymptomatic anemia should be investigated thoroughly to diagnose and treat the primary problem.

Anemia is defined by a PCV, HCT, HGB or RBC count below the reference intervals for that species. Anemia can be mild, moderate or severe and could be caused by an acute disease process or have been ongoing for a long time due to a chronic condition.

When evaluating an anemic patient, hematology testing MUST include BOTH quantitative automated cell count + gualitative blood smear evaluation

Once we receive abnormal RBC, HCT, HGB or PCV results, how do we proceed?

- as the minimum database and other diagnostic tests
- Consider the potential for laboratory or sampling error
- If an automated count is performed and anemia detected:
- Confirm with a PCV, since this is the direct measurement of the proportion of blood comprised of RBC
- Perform a blood smear to examine the RBC morphology and confirm automated cell counts to aid in determining a diagnosis and prognosis
- Remember, a comprehensive CBC includes an automated cell count along with a blood smear evaluation

Note: Anemia can be masked by concomitant dehydration. Decreased measured erythrocyte parameters may also be observed when the total-body erythrocyte mass is normal but there is an expansion of the vascular space faster than the expansion of the total-body erythrocyte mass (relative anemia).³⁰⁻³²



• Evaluate the hematology results in the context of the entire patient, including the patient's signalment and clinical status as well







How VETSCAN[®] HM5 Works

Sample Handling

- Patient preparation
- Sample collection
- Running a sample

Complete Hematology Picture

- Minimum database
- Automated CBC
- Blood smear
- PCV/TS

Responsible Patient Trending

Histograms

- Interpreting histograms
- Histogram examples

Interpretation Guide

- Macrothrombocytopenia
- Leukogram patterns
- Understanding anemia

Alternative Fluids

Reference Intervals (Ranges)

Further diagnostic testing to determine the underlying cause of anemia³³

Blood smear examination provides information about blood cell pathology and the potential for blood parasites not available with automated analyzers. In addition, evaluation of RBC morphology can help pinpoint a diagnosis, determine the recommended treatment and monitor the response to treatment for anemia.

Common RBC morphologies

IMHA	Regenerative anemia	RBC damage due to microangiopathy*	Oxidative damage	Iron deficien
Spherocytes	Anisocytosis	Schistocytes	Eccentrocytes	Schistocytes
Agglutination	Howell-Jolly bodies	Acanthocytes	Heinz bodies	Microcytes
Ghost cells	Polychromasia	Keratocytes	Spherocytes	Leptocytes

*Associated with neoplasia, disseminated intravascular coagulation, glomerulonephritis or vasculitis.

Consider the whole patient

Because anemia is a manifestation of an underlying disorder and not a diagnosis, further diagnostic testing is usually necessary to determine the underlying cause. Additional diagnostic tests listed below should be utilized based on the differential diagnosis suggested by the classification of anemia (see **Anemia algorithm** on page 26).

Additional diagnostic tests

- Clinical chemistry profile/urinalysis +/- endocrine testing
- Virology, serology if infection is likely (eg, fever, lymphadenopathy, etc)
- dyserythropoiesis, leukemia, metastatic neoplasia, myelodysplastic syndromes, etc)

IMHA=immune mediated hemolytic anemia.



• Bone marrow examination may reveal many diagnoses (eg, myelofibrosis, aplastic anemia, bone marrow necrosis/inflammation,







How VETSCAN[®] HM5 Works

Sample Handling

- Patient preparation
- Sample collection
- Running a sample

Complete Hematology Picture

- Minimum database
- Automated CBC
- Blood smear
- PCV/TS

Responsible Patient Trending

Histograms

- Interpreting histograms
- Histogram examples

Interpretation Guide

- Macrothrombocytopenia
- Leukogram patterns
- Understanding anemia

Alternative Fluids

Reference Intervals (Ranges)

Two ways to classify anemia: bone marrow responsiveness

- Evaluation must be interpreted relative to the duration and severity of the anemia³²
- the erythroid response
- the automated RETIC count

RETIC counts can be interpreted by either absolute or corrected counts to determine if regeneration exists



VETSCAN IMAGYST[™] can identify polychromatophils





• Classification of anemia in accordance with bone marrow responsiveness is based on the presence or absence of an increased number of immature erythrocytes in circulation (known as reticulocytosis, polychromasia) or erythroid hyperplasia in the bone marrow.

• In most species, a RETIC count is considered the easiest, most reliable measure of marrow responsiveness. A notable exception is the horse, which releases few to no immature RBC into circulation; therefore, a bone marrow sample must be used to determine

- Note: an automated RETIC count should always be verified with a blood smear to examine RBC morphology and to confirm

• Interpretation must be made relative to the duration and severity of the anemia. Simply relying on a reference interval may lead to misinterpretation of the erythroid response. See **Anemia algorithm** on page 26 for examples

> VETSCAN IMAGYST supports using a PCM count to classify the bone marrow response to an anemia. RETIC and PCM are both immature red blood cells and have the same function but are identified using different staining methodologies.

> Image obtained from VETSCAN IMAGYST at 400x magnification, showing Wright's-stained blood smear of RBC. Immature RBC are shown in a purple-blue color.







How VETSCAN[®] HM5 Works

Sample Handling

- Patient preparation
- Sample collection
- Running a sample

Complete Hematology Picture

- Minimum database
- Automated CBC
- Blood smear
- PCV/TS

Responsible Patient Trending

Histograms

- Interpreting histograms
- Histogram examples

Interpretation Guide

- Macrothrombocytopenia
- Leukogram patterns
- Understanding anemia

Alternative Fluids

Reference Intervals (Ranges)

Two ways to classify anemia: bone marrow responsiveness



*Evaluation of the adequacy of the bone marrow regenerative response in the individual patient should also include consideration of severity and chronicity of the anemia, suspected cause of the anemia and potential for multiple causes contributing to the patient's anemia. Trending the anemia and RETIC/PCM count through sequential CBC may be helpful.

See the full anemia algorithm 🖊

*RETIC/PCM counts supporting regeneration can be seen in nonanemic patients. This may reflect a normal physiological response or a response to an increased need. Serial evaluations of the CBC should be done to rule out an emerging anemia in these patients. RETIC or elevated PCM counts in the absence of anemia (RAA) may indicate recovery from anemia or may be associated with nonanemic chronic hypoxia (eg, cardiovascular disease, pulmonary disease). RAA has also been observed in patients with gastrointestinal, inflammatory and neoplastic disorders and in dogs with osteoarthritis or receiving osteoarthritis treatments (eg, anti-inflammatory drugs, nutraceuticals).











How VETSCAN[®] HM5 Works

Sample Handling

- Patient preparation
- Sample collection
- Running a sample

Complete Hematology Picture

- Minimum database
- Automated CBC
- Blood smear
- PCV/TS

Responsible Patient Trending

Histograms

- Interpreting histograms
- Histogram examples

Interpretation Guide

- Macrothrombocytopenia
- Leukogram patterns
- Understanding anemia

Alternative Fluids

Reference Intervals (Ranges)

Two ways to classify anemia: red blood cell indexes

- See Anemia algorithm on page 26 for additional information

Table: MCV classifications

MCV	Description
Decreased	Microcytic
Normal	Normocytic
Increased	Macrocytic

The 3 most important and relevant anemia diagnostic patterns using RBC indexes are³⁴:

Microcytic Hypochromic



Usually due to iron deficiency anemias

Images obtained from VETSCAN IMAGYST™.



• In addition to RETIC or PCM counts, it is important to review the pertinent RBC parameters found on the automated CBC report: MCV and MCHC to describe trends in RBC size and HGB concentration to aid in classification of the anemia

Common pathology

- Iron deficiency
- Hepatic portocaval vascular shunts
- Normal breed variation (eg, Shiba Inu, Akita)
- Usually nonregenerative, poorly or early regenerative
- "Early regenerative" refers to blood loss or blood destruction anemia in which evidence of regeneration is not yet apparent because the bone marrow has not had time to respond to acute loss
- Regeneration: bone marrow is responding and is releasing PCM/RETIC that are larger than normal
- Congenital poodle macrocytosis
- Hereditary stomatocytosis
- Myelodysplasia
- FeLV

Normocytic Normochromic



Nonregenerative anemias with residual normal erythrocytes

Macrocytic Hypochromic



Regenerative anemias with large, young erythrocytes that are not fully hemoglobinized







Anemia classification by red blood cell indexes³⁵

How VETSCAN[®] HM5 Works

Sample Handling

- Patient preparation
- Sample collection
- Running a sample

Complete Hematology Picture

- Minimum database
- Automated CBC
- Blood smear
- PCV/TS

Responsible Patient Trending

Histograms

- Interpreting histograms
- Histogram examples

Interpretation Guide

- Macrothrombocytopenia
- Leukogram patterns
- Understanding anemia

Alternative Fluids

Reference Intervals (Ranges)



Microcytic

- Chronic iron deficiency
- Portosystemic shunts (often not anemic)
- Anemia of inflammatory disease (usually normocytic)
- Hepatic lipidosis in cats (usually normocytic)
- Normal Akita and Shiba dogs (not anemic)
- Familial dyserythropoiesis of English springer spaniels (rare)
- Hereditary elliptocytosis in dogs (rare)
- Spurious when PLT are included in erythrocyte histograms
- Spurious in dogs with persistent hyponatremia (not typically anemic)









Alternative Fluids



How VETSCAN[®] HM5 Works

Sample Handling

- Patient preparation
- Sample collection
- Running a sample

Complete Hematology Picture

- Minimum database
- Automated CBC
- Blood smear
- PCV/TS

Responsible Patient Trending

Histograms

- Interpreting histograms
- Histogram examples

Interpretation Guide

- Macrothrombocytopenia
- Leukogram patterns
- Understanding anemia

Alternative Fluids

Reference Intervals (Ranges)

Running alternative fluids on the VETSCAN HM5

Identify in-clinic nucleated cell counts* in alternative fluids (including pleural, peritoneal and synovial fluids) with the VETSCAN HM5. Characterize exudate samples to diagnose and identify potential inflammation, infections and neoplasia.



Synovial fluid procedure

- 1. Add a flake of hyaluronidase[‡] (enzyme concentrate) to sample in a serum top tube, mix and store at room temperature for 10 minutes
- 2. Run the undiluted sample under the appropriate species setting on the VETSCAN HM5
- **3.** Observe the total WBC number, which is useful in the determination of whether the WBC count is normal or abnormal
- 4. Microscopically evaluate a stained smear for a leukocyte differential and identification of nonleukocyte cells and/or infectious organisms
- 5. Perform a Soak Cleaning cycle (blue HM5) using HemaClean after every synovial fluid sample run

Contact Diagnostic Technical Support 24/7 for assistance with interpreting VETSCAN HM5 troubleshooting reports



(888) 963-8471 (option 5) \bigcirc

- by Zoetis Diagnostics.
- [‡]Hyaluronidase from Sigma Chem Co, Part No. H3506, 100 mg bottle.



Pleural and peritoneal fluid procedure

- 1. Run undiluted sample in a serum tube under the appropriate species setting on the VETSCAN HM5
- 2. Observe the total WBC number, which is useful for classification of fluids and provides the total nucleated cell count
- 3. Perform a total protein concentration on a refractometer
- 4. Microscopically evaluate a stained smear for a leukocyte differential and identification of nonleukocyte cells and/or infectious organisms

dxsupport@zoetis.com

*VETSCAN HM5 analyzers are not intended to provide accurate differential cell counts for alternative fluid analysis samples.

⁺For results for species and/or fluids not validated, fluid samples may be run. However, precision and accuracy are not available, and results will not be supported





VETSCAN HM5 reference intervals, SI units⁷

How VETSCAN®	HM5
Works	

Sample Handling

- Patient preparation
- Sample collection
- Running a sample

Complete Hematology Picture

- Minimum database
- Automated CBC
- Blood smear
- PCV/TS

Responsible Patient Trending

Histograms

- Interpreting histograms
- Histogram examples

Interpretation Guide

- Macrothrombocytopenia
- Leukogram patterns
- Understanding anemia

Alternative Fluids

Reference Intervals (Ranges)

Parameter	Units	DOG	کرکر CAT ³⁷	HORSE	CATTLE	SHEEP*
WBC	10 ⁹ cells/l	6.0 - 17.0	3.5 - 20.7	5.4 - 14.3	4.0 - 12.0	4.0 - 12.0
LYM	10 ⁹ cells/l	1.00 - 4.80	0.83 - 9.10	1.50 - 7.70	2.50 - 7.50	2.00 - 9.00
MON	10 ⁹ cells/l	0.20 - 1.50	0.09 - 1.21	0 - 1.50	0 - 0.84	0 - 0.75
NEU (GRA)	10 ⁹ cells/l	3.00 - 12.00	1.63 - 13.37	2.30 - 9.50	0.60 - 6.70	(0.70 - 7.30)
EOS	10 ⁹ cells/l	0 - 0.80	0.02 - 0.49	0 - 1.00	0.10 - 1.00	
RBC	10 ¹² cells/l	5.5 - 8.5	7.7 - 12.8	6.8 - 12.9	5.0 - 10.0	9.0 - 15.8
НСТ	%	37.0 - 55.0	33.7 - 55.4	32.0 - 53.0	24.0 - 46.0	27.0 - 45.0
HGB	g/l	120 - 180	100 - 170	110 - 190	80 - 150	90 - 150
MCV	fL	60 - 77	35 - 52	37 - 59	40 - 60	28 - 40
МСН	pg	19.5 - 24.5	10.0 - 16.9	12.3 - 19.7	11.0 - 17.0	8.0 - 12.0
MCHC	g/l	310 - 390	270 - 350	310 - 390	300 - 360	310 - 340
RDWc	%	14.0 - 20.0	18.3 - 24.1	N/A	N/A	N/A
PLT	10º cells/l	165 - 500	125 - 618	100 - 400	100 - 800	100 - 800
MPV	fL	3.9 - 11.1	8.6 - 14.9	N/A	N/A	N/A

Reference intervals are determined from a population of healthy adult animals of a given species for a given test.

---: EOS not included in 3-part differential. N/A: Reference interval data not available for these parameters *3-part differential with GRA.

VETSCAN HM5 reference intervals continue on the next page.

GOAT*	
4.0 - 13.0	
2.00 - 9.00	
0 - 0.50	
(1.20 - 8.00)	
5.5 - 8.5	
37.0 - 55.0	
120 - 180	
60 - 77	
19.5 - 24.5	
310 - 340	
N/A	
200 - 500	
N/A	



How VETSCAN[®] HM5 Works

Sample Handling

- Patient preparation
- Sample collection
- Running a sample

Complete Hematology Picture

- Minimum database
- Automated CBC
- Blood smear
- PCV/TS

Responsible Patient Trending

Histograms

- Interpreting histograms
- Histogram examples

Interpretation Guide

- Macrothrombocytopenia
- Leukogram patterns
- Understanding anemia

Alternative Fluids

Reference Intervals (Ranges)

VETSCAN HM5 reference intervals, SI units⁷ (cont'd)

Parameter	Units	RABBIT*	FERRET*	GUINEA PIG*	RAT*	MOUSE*†	PIG*	ALPACA	LLAMA
WBC	10º cells/l	3.0 - 11.5	2.0 - 10.0	5.0 - 17.0	2.1 - 19.5	6.0 - 15.0	11.0 - 22.0	6.0 - 30.0	8.0 - 23
LYM	10 ⁹ cells/l	2.00 - 9.10	0.40 - 6.50	2.00 - 15.00	2.00 - 14.10	3.40 - 7.44	5.50 - 11.10	1.00 - 20.00	1.00 - 6.00
MON	10º cells/l	0 - 0.50	0.10 - 0.70	N/A	0 - 0.98	0 - 0.60	0.66 - 1.32	N/A	N/A
NEU (GRA)	10 ⁹ cells/l	(0 - 2.80)	(0.80 - 4.50)	(1.00 - 11.00)	(0.10 - 5.40)	(0.50 - 3.80)	(5.00 - 10.00)	3.00 - 20.00	5.00 - 24.00
EOS	10 ⁹ cells/l							N/A	N/A
RBC	10 ¹² cells/l	5.0 - 9.0	7.8 - 13.0	4.8 - 6.3	5.3 - 10.0	7.0 - 12.0	5.0 - 8.0	8.0 - 20.0	10.0 - 17.0
НСТ	%	36.0 - 50.0	36.0 - 56.0	30.0 - 44.0	35.0 - 52.0	35.0 - 45.0	32.0 - 50.0	25.0 - 45.0	25.0 - 50.0
HGB	g/l	130 - 160	120 - 180	80 - 150	140 - 180	120 - 160	100 - 160	90 - 210	110 - 180
MCV	fL	57 - 70	40 - 48	50 - 90	50 - 62	45 - 55	50 - 68	15 - 35	20 - 35
МСН	pg	17.5 - 23.5	13.5 - 16.5	12.0 - 13.0	16.0 - 23.0	11.1 - 12.7	17.0 - 21.0	7.5 - 13.5	10.0 - 14.0
MCHC	g/l	300 - 380	320 - 350	300 - 360	310 - 400	220 - 320	300 - 340	300 - 450	300 - 450
RDWc	%	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A
PLT	10 ⁹ cells/l	218 - 641	96 - 776	200 - 600	500 - 1000	200 - 450	325 - 715	N/A	N/A
MPV	fL	N/A	N/A	N/A	N/A	N/A	N/A	N/A	N/A

---: EOS not included in 3-part differential.

N/A: Reference interval data not available for these parameters *3-part differential with GRA.

⁺Different mouse models may have varied intervals, and these reference intervals should be used only as a guideline.

VETSCAN HM5 reference intervals continue on the next page.

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VETSCAN HM5 reference intervals, common units⁷

How VETSCAN[®] HM5 Works

Sample Handling

- Patient preparation
- Sample collection
- Running a sample

Complete Hematology Picture

- Minimum database
- Automated CBC
- Blood smear
- PCV/TS

Responsible Patient Trending

Histograms

- Interpreting histograms
- Histogram examples

Interpretation Guide

- Macrothrombocytopenia
- Leukogram patterns
- Understanding anemia

Alternative Fluids

Reference Intervals (Ranges)

Parameter	Units	DOG	رکر CAT ³⁷	HORSE	CATTLE	SHEEP*
WBC	10º cells/l	6.0 - 17.0	3.5 - 20.7	5.4 - 14.3	4.0 - 12.0	4.0 - 12.0
LYM	10 ⁹ cells/l	1.00 - 4.80	0.83 - 9.10	1.50 - 7.70	2.50 - 7.50	2.00 - 9.00
MON	10 ⁹ cells/l	0.20 - 1.50	0.09 - 1.21	0 - 1.50	0 - 0.84	0 - 0.75
NEU (GRA)	10 ⁹ cells/l	3.00 - 12.00	1.63 - 13.37	2.30 - 9.50	0.60 - 6.70	(0.70 - 7.30)
EOS	10 ⁹ cells/l	0 - 0.80	0.02 - 0.49	0 - 1.00	0.10 - 1.00	
RBC	10 ¹² cells/l	5.5 - 8.5	7.7 - 12.8	6.8 - 12.9	5.0 - 10.0	9.0 - 15.8
НСТ	%	37.0 - 55.0	33.7 - 55.4	32.0 - 53.0	24.0 - 46.0	27.0 - 45.0
HGB	g/dl	12 - 18	10 - 17	11 - 19	8 - 15	9 - 15
MCV	fL	60 - 77	35 - 52	37 - 59	40 - 60	28 - 40
MCH	pg	19.5 - 24.5	10.0 - 16.9	12.3 - 19.7	11.0 - 17.0	8.0 - 12.0
MCHC	g/dl	31 - 39	27 - 35	31 - 39	30 - 36	31 - 34
RDWc	%	14.0 - 20.0	18.3 - 24.1	N/A	N/A	N/A
PLT	10º cells/l	165 - 500	125 - 618	100 - 400	100 - 800	100 - 800
MPV	fL	3.9 - 11.1	8.6 - 14.9	N/A	N/A	N/A

Reference intervals are determined from a population of healthy adult animals of a given species for a given test.

---: EOS not included in 3-part differential. N/A: Reference interval data not available for these parameters *3-part differential with GRA.

VETSCAN HM5 reference intervals continue on the next page.

GOAT*
4.0 - 13.0
2.00 - 9.00
0 - 0.50
(1.20 - 8.00)
5.5 - 8.5
37.0 - 55.0
12 - 18
60 - 77
19.5 - 24.5
31 - 34
N/A
200 - 500
N/A



How VETSCAN® HM5 Works

Sample Handling

- Patient preparation
- Sample collection
- Running a sample

Complete Hematology Picture

- Minimum database
- Automated CBC
- Blood smear
- PCV/TS

Responsible Patient Trending

Histograms

- Interpreting histograms
- Histogram examples

Interpretation Guide

- Macrothrombocytopenia
- Leukogram patterns
- Understanding anemia

Alternative Fluids

Reference Intervals (Ranges)

VETSCAN HM5 reference intervals, common units⁷ (cont'd)

Parameter	Units	RABBIT*	FERRET*	GUINEA PIG*	RAT*	MOUSE*†	PIG*	ALPACA
WBC	10 ⁹ cells/l	3.0 - 11.5	2.0 - 10.0	5.0 - 17.0	2.1 - 19.5	6.0 - 15.0	11.0 - 22.0	6.0 - 30.0
LYM	10 ⁹ cells/l	2.00 - 9.10	0.40 - 6.50	2.00 - 15.00	2.00 - 14.10	3.40 - 7.44	5.50 - 11.10	1.00 - 20.00
MON	10 ⁹ cells/l	0 - 0.50	0.10 - 0.70	N/A	0 - 0.98	0 - 0.60	0.66 - 1.32	N/A
NEU (GRA)	10 ⁹ cells/l	(0 - 2.80)	(0.80 - 4.50)	(1.00 - 11.00)	(0.10 - 5.40)	(0.50 - 3.80)	(5.00 - 10.00)	3.00 - 20.00
EOS	10 ⁹ cells/l							N/A
RBC	10 ¹² cells/l	5.0 - 9.0	7.8 - 13.0	4.8 - 6.3	5.3 - 10.0	7.0 - 12.0	5.0 - 8.0	8.0 - 20.0
НСТ	%	36.0 - 50.0	36.0 - 56.0	30.0 - 44.0	35.0 - 52.0	35.0 - 45.0	32.0 - 50.0	25.0 - 45.0
HGB	g/dl	13 - 16	12 - 18	8 - 15	14 - 18	12 - 16	10 - 16	9 - 21
MCV	fL	57 - 70	40 - 48	50 - 90	50 - 62	45 - 55	50 - 68	15 - 35
MCH	pg	17.5 - 23.5	13.5 - 16.5	12.0 - 13.0	16.0 - 23.0	11.1 - 12.7	17.0 - 21.0	7.5 - 13.5
MCHC	g/dl	30 - 38	32 - 35	30 - 36	31 - 40	22 - 32	30 - 34	30 - 45
RDWc	%	N/A	N/A	N/A	N/A	N/A	N/A	N/A
PLT	10 ⁹ cells/l	218 - 641	96 - 776	200 - 600	500 - 1000	200 - 450	325 - 715	N/A
MPV	fL	N/A	N/A	N/A	N/A	N/A	N/A	N/A

Need help? Contact Diagnostic Technical Support 24/7

(888) 963-8471 (option 5)

@ <u>dxsupport@zoetis.com</u>

---: EOS not included in 3-part differential.

N/A: Reference interval data not available for these parameters *3-part differential with GRA.

⁺Different mouse models may have varied intervals, and these reference intervals should be used only as a guideline.

LLAMA 8.0 - 23 1.00 - 6.00 N/A 5.00 - 24.00 N/A 10.0 - 17.0 25.0 - 50.0 11 - 18 20 - 35 10.0 - 14.0 30 - 45 N/A N/A N/A

References



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PCV/TS

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Interpreting histograms

Histogram examples

Interpretation Guide

Macrothrombocytopenia

Leukogram patterns

Understanding anemia

Alternative Fluids

Reference Intervals (Ranges)

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