

1 **Characteristics of a novel NMR-based metabolomics platform for dogs**

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21 **Abstract**

22 Metabolomics has proven itself an invaluable research tool, providing comprehensive insight to systemic metabolism.
23 However, the lack of scalable and quantitative methods with known reference intervals and documented reproducibility
24 has prevented the use of metabolomics in the clinical setting. This study describes the development and validation of a
25 quantitative nuclear magnetic resonance (NMR) -based metabolomics platform for canine serum and plasma samples.
26 Altogether 8247 canine samples were analyzed using a Bruker's 500 MHz NMR spectrometer. Using statistical
27 approaches derived from international guidelines, we defined reference intervals for 123 biomarkers, studied method
28 precision, analyte storage stability, the effect of prolonged contact to red blood cells, differences of blood collection

29 tubes, interference of lipemia, hemolysis and bilirubinemia, method comparison, and demonstrated the method's
30 practical relevance in a hyperglycemic cohort. Owing to the advantages of quantitative results, high reproducibility, and
31 scalability, this canine metabolomics platform holds great potential for numerous clinical and research applications to
32 improve canine health and well-being.

33

34 **Keywords: canine, metabolomics, NMR, reference intervals, precision, stability**

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36 Metabolomics is an omics-based approach that generates comprehensive information on metabolism, enabling an
37 extensive view on the current state of systemic metabolism. Metabolomics has become increasingly popular in canine
38 studies. It is especially suitable in characterizing metabolic effects of multiple environmental and inter- and intra-
39 individual¹ factors including feeding²⁻⁶, aging⁷, inter-breed differences^{8,9}, drug action^{10,11}, behavior^{12,13}, exercise¹⁴,
40 genetic factors¹⁵ and pathological processes¹⁶⁻²⁶.

41 Mass spectrometry (MS) and proton nuclear magnetic resonance (NMR) spectroscopy are the two main technologies for
42 metabolic profiling, both in human and animal studies²⁷. Proton NMR spectroscopy offers quantitative data with high
43 reproducibility, high throughput and excellent scalability in a non-destructive and cost-effective manner²⁷. NMR
44 spectroscopy is well suited for the scientific use of large cohorts and biobanks^{28,29} and is highly suitable for studies
45 combining different omics technologies²⁹⁻³¹. Due to the quantitative and highly reproducible nature of NMR
46 spectroscopy, it can be easily utilized as a diagnostic tool and research findings can be easily applied to clinical use.

47 Metabolomics holds great potential in clinical diagnostics. It is a promising tool for highlighting the metabolic
48 alterations associated with the emergence and progression of diseases, early disease detection and individualization of
49 treatment³²⁻³⁴. Clinical diagnostics and decision-making have formerly relied on the use of single clinical chemistry
50 biomarkers. The routinely used approach is largely unusable for risk prediction, since changes are typically only seen
51 when the diseased state has already been reached^{33,35}. Thus, preventive and supportive care measures cannot be
52 undertaken before organ failure is already present. Interpretation of the results by the traditional approach might also be
53 challenging, since multiple simultaneous metabolic phenomena can affect the interpretation of these results and multiple
54 diseases can be associated with similar laboratory results^{22,35-37}. Moreover, evaluations of disease severity lack accuracy
55 when only one affected metabolic pathway is taken into account^{34,37}. Furthermore, traditional diagnostic approaches do
56 not take the metabolic state of the patient into account, although it might have an impact on the best treatment of the
57 patient^{34,38}.

58 The prerequisites for the utility of a particular method include documentation of the method's reproducibility,
59 understanding of confounding factors that might affect the interpretation of the results, and the formation of reference
60 intervals³⁹. All of these factors have previously been largely unpublished in the canine metabolomics field. In this study,
61 we describe the development and validation of a novel NMR-based metabolomics platform for dogs, demonstrate its
62 performance characteristics, publish the determined dog-specific reference intervals for its biomarkers and demonstrate
63 its utility in a practical situation.

64

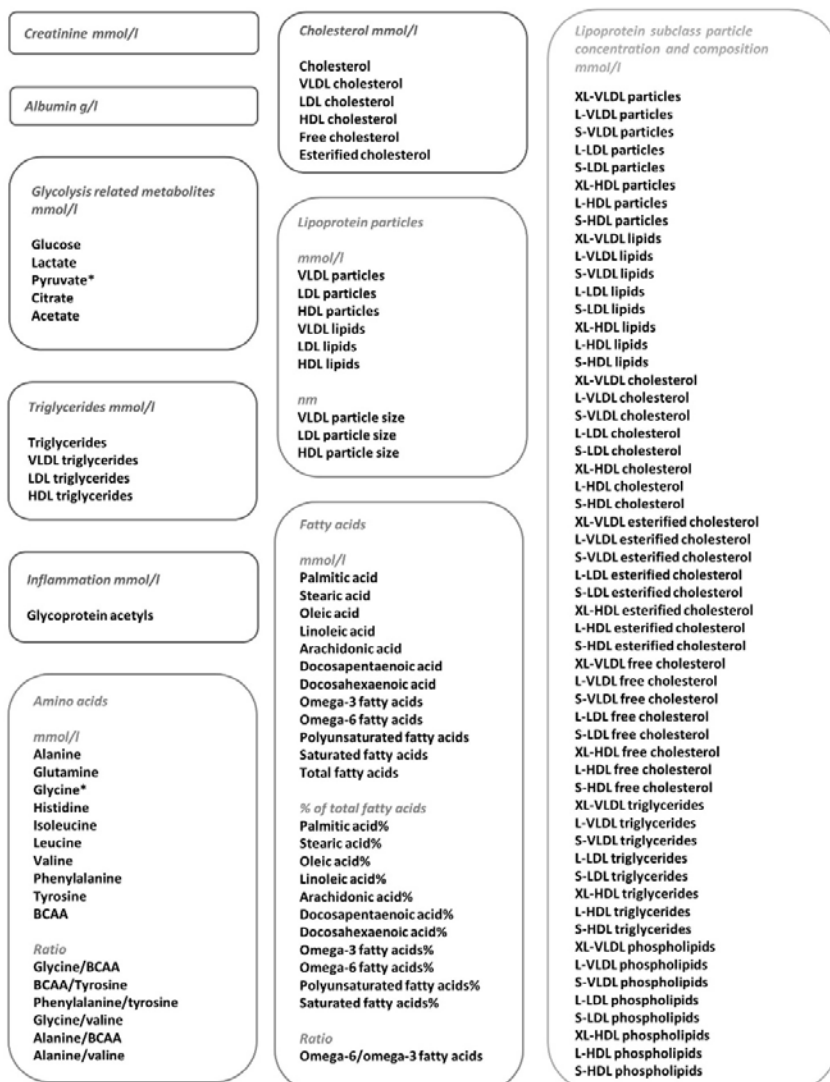
65 **Results**

66 To develop and validate a quantitative NMR-based approach for canine metabolomics, a total of 8247 serum or plasma
67 samples were collected and analyzed. Thirty of these were pooled samples, 912 were replicate samples of client-owned
68 dogs, 999 leftover clinical laboratory samples, and 6306 were single samples of client-owned dogs (Supplementary
69 Figure 1). The number of individual client-owned dogs with samples collected during this project was 6164
70 (Supplementary Table 1). Eighteen percent of the dogs were under one year of age, 61% aged 1-7, and 21% over 7
71 years of age. Males consisted 40% of the population and samples were taken from 256 breeds.

72 **Proton NMR spectroscopy quantitates 123 biomarkers**

73 All samples were analyzed with Bruker's 500 MHz NMR spectrometer with spectral data interpreted by in-house
74 scripts following principles demonstrated for human samples⁴⁰. The NMR method was able to quantify 123 biomarkers,
75 including extensive lipoprotein profiling, fatty acids, amino acids, albumin, creatinine and glycolysis related
76 metabolites, in canine serum/plasma (Fig 1).

BIOMARKERS



77

78 **Fig 1.** The biomarkers quantified by the NMR metabolomics testing platform.

79 *: not available from EDTA-plasma samples.

80 BCAA: Branched-chain amino acids

81 XL-VLDL: Chylomicrons and very large VLDL-particles

82 L-VLDL: large VLDL particles

83 S-VLDL: small VLDL-particles

84 L-LDL: large LDL particles

85 S-LDL: small LDL particles

86 XL-HDL: very large HDL particles

87 L-HDL: large HDL particles

88 S-HDL: small HDL particles

89

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91 **Reference intervals for puppies, adult and senior dogs**

92 We determined reference intervals (RI) (Table 1, Supplementary Table 2) with 90% confidence intervals (CI)

93 (Supplementary Table 2) for 123 metabolites in serum and heparin plasma samples.

94 In certain analytes, such as XL-VLDL variables, the concentration in healthy animals was very low, causing highly

95 skewed distributions. Also, automatic rejection of extremely low values by the platform's quality control, caused

96 inability to calculate CIs with the used non-parametric method. For these analytes, the lower reference limit was

97 rounded to 0. For a multitude of analytes, the 90% CI width was higher than 20% of the RI width, due to skewed or

98 heavy tailed distributions. This was especially observed in RI for puppies and senior dogs, where the n count was lower

99 ($n < 170$) than in adult dogs. However, for some analytes, it was also observed in RIs with very high n counts ($n > 800$).

100

101 RIs of puppies differed from the RIs of adult dogs in many analytes, such as glucose, lactate, creatinine, albumin,

102 glycine, glutamine, leucine, valine, branched chain amino acids (BCAA), VLDL particle size, and HDL and LDL

103 triglycerides. The higher reference limit (RL) of senior dogs exceeded the higher RL of adult dogs for alanine, tyrosine,

104 alanine/BCAA, glycoprotein acetyls (GlycA), oleic acid, docosahexaenoic acid%, HDL triglycerides, S-VLDL lipids

105 and L-HDL triglycerides.

106 Table 1. Established reference intervals for the validated biomarkers.

Analyte	HP all dogs	S all dogs	S puppy	S adult	S senior
Glucose mmol/l	4.3-6.8	4.4-6.8	4.7-7.3 ^a	4.4-6.6	4.0-6.1
Lactate mmol/l	0.7-3.0	1.1-3.6	1.3-2.9 ^b	1.1-3.6	1.1-4.2
Creatinine mmol/l	40-99	32-103	21-96 ^b	40-108	37-104
Albumin mg/dl	26-32	25-32	22-31 ^b	26-32	26-32
Cholesterol mmol/l	3.8-10.4	3.6-10.3	3.8-10.1	3.6-10.3	3.5-10.6
Triglycerides mmol/l	0.22-0.97	0.19-1.00	0.18-0.76	0.19-0.97	0.19-1.13
Pyruvate μ mol/l	29-155	11-107	9-96	11-111	13-109
Citrate μ mol/l	63-122	61-123	57-115	61-124	62-128
Acetate μ mol/l	19-36	21-37	22-40	21-37	20-37
Alanine μ mol/l	214-584	216-597	205-504	205-583	244-650 ^a
Glycine μ mol/l	147-466	130-454	145-594 ^{a,b}	127-381	128-403
Glutamine μ mol/l	570-919	640-1,015	642-997 ^b	659-1,028	616-1,014
Histidine μ mol/l	50-91	53-98	46-93	55-99	55-97

Isoleucine $\mu\text{mol/l}$	33-80	37-89	30-80 ^a	38-92	38-87
Leucine $\mu\text{mol/l}$	74-168	83-185	67-159 ^b	89-186	94-186
Valine $\mu\text{mol/l}$	107-245	113-251	80-236 ^b	119-260	124-253
Phenylalanine $\mu\text{mol/l}$	32-64	30-65	28-66	29-64	34-66
Tyrosine $\mu\text{mol/l}$	39-85	41-89	39-83	41-90	46-93 ^a
BCAA $\mu\text{mol/l}$	222-482	242-515	178-462 ^b	261-524	251-521
Glycine/BCAA	0.3-1.4	0.3-1.5	0.4-3.0 ^a	0.3-1.2	0.3-1.2
BCAA/Tyrosine	3.7-9.5	3.8-9.2	3.8-8.3 ^a	3.9-9.5	3.5-8.6
Phenylalanine/tyrosine	0.6-1.2	0.5-1.0	0.5-1.1	0.5-1.0	0.5-1.0
Glycine/valine	0.6-2.6	0.7-3.0	0.9-6.8 ^a	0.7-2.5	0.7-2.6
Alanine/BCAA	0.7-2.0	0.6-1.6	0.6-1.4	0.5-1.6	0.6-1.8 ^a
Alanine/valine	1.3-3.7	1.2-3.5	1.3-3.0	1.1-3.5	1.3-3.8
VLDL lipids mmol/l	0.1-1.1	0.1-1.2	0.1-0.8	0.1-1.1	0.1-1.3
VLDL particles nmol/l	15-54	12-54	14-45	12-51	12-67
LDL lipids mmol/l	0.5-3.7	0.7-3.7	0.7-3.7	0.7-3.6	0.7-4.5
LDL particles nmol/l	200-1,400	240-1,300	270-1,300	240-1,300	240-1,600
HDL lipids mmol/l	7.7-14.6	6.9-15.1	7.2-15.0	6.9-15.1	6.6-15.3
HDL particles nmol/l	32,000-57,000	30,000-58,000	33,000-58,000	30,000-59,000	28,000-58,000
VLDL particle size nm	35.8-43.6	35.2-43.8	35.1-41.7 ^a	35.3-43.8	35.4-44.4
LDL particle size nm	22.1-23.9	22.2-23.5	22.3-23.5	22.2-23.4	22.2-23.5
HDL particle size nm	10.1-10.7	10.1-10.7	10.1-10.6	10.0-10.6	10.1-10.7
VLDL cholesterol mmol/l	0.03-0.31	0.03-0.31	0.03-0.23	0.03-0.29	0.03-0.36
LDL cholesterol mmol/l	0.16-2.35	0.28-2.27	0.30-2.24	0.28-2.21	0.28-2.85
HDL cholesterol mmol/l	3.6-8.0	3.2-7.9	3.6-7.9	3.2-7.9	3.0-8.0
Esterified cholesterol mmol/l	3.1-8.2	2.9-8.1	3.2-8.0	2.9-8.1	2.9-8.4
Free cholesterol mmol/l	0.7-2.2	0.6-2.2	0.7-2.0	0.6-2.2	0.6-2.4
VLDL triglycerides mmol/l	0.02-0.70	0.00-0.70	0.00-0.44	0.00-0.69	0.00-0.86
LDL triglycerides mmol/l	0.13-0.30	0.13-0.31	0.15-0.33 ^a	0.13-0.29	0.14-0.27
HDL triglycerides mmol/l	0.01-0.08	0.00-0.08	0.00-0.04 ^b	0.00-0.07	0.01-0.09 ^a
Glycoprotein acetyls $\mu\text{mol/l}$	532-964	597-1,028	611-996	596-1,005	605-1,129 ^a
Palmitic acid mmol/l	1.9-3.5	1.8-3.6	1.8-3.4	1.8-3.6	1.8-3.9
Stearic acid mmol/l	1.9-3.8	1.7-3.8	1.8-3.7	1.7-3.9	1.7-4.0

Oleic acid mmol/l	1.1-2.5	1.3-2.8	1.3-2.5	1.3-2.8	1.3-3.2 ^a
Linoleic acid mmol/l	2.7-5.7	2.5-5.9	2.5-5.1	2.5-5.9	2.5-6.2
Arachidonic acid mmol/l	1.6-3.7	1.4-3.6	1.5-3.7	1.4-3.6	1.3-3.7
Docosapentaenoic acid mmol/l	0.1-0.3	0.1-0.4	0.1-0.3	0.1-0.4	0.1-0.4
Docosahexaenoic acid mmol/l	0.1-0.9	0.1-0.7	0.1-0.8	0.1-0.7	0.1-0.8
Omega-3 fatty acids mmol/l	0.4-1.8	0.4-1.6	0.4-1.6	0.4-1.6	0.5-1.8
Omega-6 fatty acids mmol/l	4.5-9.6	4.1-9.8	4.2-9.3	4.1-9.6	3.9-10.6
Polyunsaturated fatty acids mmol/l	5.2-10.8	4.7-11.1	4.8-10.6	4.7-11.0	4.6-12.0
Saturated fatty acids mmol/l	3.9-7.2	3.6-7.4	3.6-6.8	3.6-7.5	3.5-7.7
Total fatty acids mmol/l	10.5-20.4	9.7-21.0	9.8-20.0	9.7-21.2	9.3-22.5
Palmitic acid%	15.9-19.8	16-20	16-19	16-19	16-20
Stearic acid%	17.5-19.5	17.3-19.4	17.5-19.3	17.4-19.5	17.1-19.4
Oleic acid%	9.2-14.2	10.7-15.1	10.6-14.2 ^a	10.9-15.1	10.9-15.7
Linoleic acid%	23.9-29.4	24.1-28.6	23.3-27.5 ^{a,b}	24.5-28.6	24.7-28.7
Arachidonic acid%	13.6-20.7	13.2-20.1	15.0-20.7 ^{a,b}	13.1-19.9	12.7-19.2
Docosapentaenoic acid%	0.9-1.9	1.1-2.0	1.1-1.9	1.1-1.9	1.1-2.0
Docosahexaenoic acid%	0.9-6.2	0.7-4.8 ^b	1.0-5.2 ^b	0.7-4.7	0.6-5.0 ^a
Omega-3 fatty acids%	3-12	3-10	3-11	3-11	4-11
Omega-6 fatty acids%	43-49	42-48	42-48	42-48	42-47
Polyunsaturated fatty acids%	48-57	48-55	48-55 ^b	47-55	47-55
Saturated fatty acids%	34-39	34-38	34-38	34-38	33-38
Omega-6/omega-3 fatty acids	3.5-16.3	4.2-13.4	4.1-13.5	4.2-14.2	4.0-13.1
XL-VLDL particles nmol/l	0-2	0-1	0-1	0-1	0-2
L-VLDL particles nmol/l	1-15	0-16	0-11 ^a	0-15	0-19
S-VLDL particles nmol/l	11-38	10-39	13-34	10-37	10-47
L-LDL particles nmol/l	79-380	90-370	110-350	91-380	83-400
S-LDL particles nmol/l	89-1000	140-960	130-940	130-920	140-1,100
XL-HDL particles nmol/l	1-6,200	0-5,800	25-5,500	0-5,800	1-6,000
L-HDL particles nmol/l	21,000-34,000	20,000-34,000	20,000-34,000	19,000-35,000	18,000-34,000
S-HDL particles nmol/l	11,000-19,000	11,000-20,000	11,000-19,000	11,000-20,000	10,000-20,000

XL-VLDL lipids mmol/l	0-0.3	0-0.3	0-0.2	0-0.2	0-0.4
L-VLDL lipids mmol/l	0-0.6	0-0.6	0-0.4 ^a	0-0.6	0-0.7
S-VLDL lipids mmol/l	0.1-0.4	0.1-0.4	0.1-0.3	0.1-0.3	0.1-0.4 ^a
L-LDL lipids mmol/l	0.2-1.2	0.3-1.2	0.3-1.1	0.3-1.2	0.3-1.3
S-LDL lipids mmol/l	0.2-2.6	0.3-2.4	0.3-2.4	0.3-2.3	0.3-2.9
XL-HDL lipids mmol/l	0.6-5.2	0.6-5.0	0.6-4.7	0.6-5.0	0.7-5.2
L-HDL lipids mmol/l	5.4-8.1	5.0-8.3	5.0-8.1	4.9-8.3	4.8-8.3
S-HDL lipids mmol/l	1.3-2.1	1.3-2.2	1.3-2.1	1.3-2.2	1.1-2.2
XL-VLDL cholesterol mmol/l	0-0.06	0-0.06	0-0.04	0-0.06	0-0.08
L-VLDL cholesterol mmol/l	0-0.13	0-0.13	0-0.08	0-0.12	0-0.15
S-VLDL cholesterol mmol/l	0.01-0.14	0.02-0.15	0.02-0.11	0.02-0.14	0.02-0.18
L-LDL cholesterol mmol/l	0.03-0.70	0.07-0.68	0.11-0.61	0.07-0.68	0.07-0.76
S-LDL cholesterol mmol/l	0.09-1.64	0.16-1.53	0.16-1.50	0.16-1.48	0.18-1.88
XL-HDL cholesterol mmol/l	0.3-3.0	0.2-2.8	0.3-2.9	0.2-2.7	0.2-2.9
L-HDL cholesterol mmol/l	2.6-4.3	2.3-4.4	2.5-4.3	2.3-4.5	2.2-4.4
S-HDL cholesterol mmol/l	0.5-1.0	0.5-1.0	0.6-1.0	0.5-1.0	0.5-1.0
XL-VLDL esterified cholesterol mmol/l	0-0.03	0-0.03	0-0.02	0-0.03	0-0.03
L-VLDL esterified cholesterol mmol/l	0-0.05	0-0.06	0-0.04	0-0.05	0-0.07
S-VLDL esterified cholesterol mmol/l	0-0.09	0.01-0.09	0.01-0.08	0.01-0.09	0.01-0.11
L-LDL esterified cholesterol mmol/l	0.01-0.51	0.03-0.49	0.05-0.43	0.02-0.48	0.02-0.55
S-LDL esterified cholesterol mmol/l	0.05-1.14	0.10-1.09	0.10-1.06	0.10-1.06	0.12-1.35
XL-HDL esterified cholesterol mmol/l	0.2-2.3	0.2-2.1	0.2-2.1	0.2-2.1	0.2-2.3
L-HDL esterified cholesterol mmol/l	2.3-3.7	2.0-3.8	2.2-3.7	2.0-3.8	1.9-3.8
S-HDL esterified cholesterol mmol/l	0.4-0.8	0.4-0.8	0.5-0.8	0.4-0.8	0.4-0.8
XL-VLDL free cholesterol mmol/l	0-0.05	0-0.05	0-0.03	0-0.04	0-0.05
L-VLDL free cholesterol mmol/l	0-0.08	0-0.08	0-0.05 ^a	0-0.07	0-0.10
S-VLDL free cholesterol mmol/l	0.01-0.05	0.01-0.06	0.01-0.05	0.01-0.05	0.01-0.07
L-LDL free cholesterol mmol/l	0.03-0.20	0.04-0.20	0.05-0.18	0.04-0.20	0.04-0.21
S-LDL free cholesterol mmol/l	0.04-0.48	0.05-0.45	0.06-0.45	0.05-0.43	0.05-0.53
XL-HDL free cholesterol mmol/l	0.1-0.7	0.1-0.6	0.1-0.6	0.1-0.6	0.1-0.6
L-HDL free cholesterol mmol/l	0.3-0.6	0.3-0.6	0.3-0.6	0.3-0.6	0.3-0.6
S-HDL free cholesterol mmol/l	0.1-0.2	0.1-0.2	0.1-0.2	0.1-0.2	0.1-0.2

XL-VLDL triglycerides mmol/l	0-0.22	0-0.17	0-0.13	0-0.17	0-0.24
L-VLDL triglycerides mmol/l	0.01-0.41	0.01-0.42	0.00-0.27 ^b	0.01-0.39	0.01-0.49
S-VLDL triglycerides mmol/l	0.03-0.16	0.02-0.16	0.02-0.13	0.01-0.15	0.02-0.19
L-LDL triglycerides mmol/l	0.11-0.24	0.10-0.24	0.11-0.26 ^{a,b}	0.09-0.23	0.10-0.22
S-LDL triglycerides mmol/l	0.02-0.07	0.03-0.07	0.03-0.08	0.03-0.07	0.03-0.07
XL-HDL triglycerides mmol/l	0-0.02	0-0.02	0-0.02	0-0.02	0-0.03
L-HDL triglycerides mmol/l	0.01-0.03	0-0.03	0-0.02	0-0.03	0-0.04 ^a
S-HDL triglycerides mmol/l	0-0.04	0-0.03	0-0.02	0-0.03	0-0.04
XL-VLDL phospholipids mmol/l	0-0.05	0-0.05	0-0.04	0-0.05	0-0.09
L-VLDL phospholipids mmol/l	0-0.09	0-0.11	0-0.06	0-0.10	0-0.15
S-VLDL phospholipids mmol/l	0.01-0.07	0.01-0.08	0.01-0.06	0-0.08	0.01-0.10
L-LDL phospholipids mmol/l	0.08-0.33	0.08-0.32	0.09-0.33	0.07-0.30	0.08-0.35
S-LDL phospholipids mmol/l	0.08-0.87	0.12-0.84	0.12-0.80	0.12-0.80	0.12-1.01
XL-HDL phospholipids mmol/l	0.35-2.27	0.33-2.22	0.32-1.97	0.30-2.21	0.34-2.37
L-HDL phospholipids mmol/l	2.65-3.92	2.48-3.94	2.61-3.79	2.44-3.95	2.56-4.01
S-HDL phospholipids mmol/l	0.74-1.18	0.72-1.19	0.76-1.14 ^a	0.72-1.19	0.67-1.21

107 HP all dogs n = 269, S all dogs n =865, S puppy n =152, S adult n =545, S senior n =168.

108 HP: heparinized plasma

109 S: serum

110 BCAA: Branched-chain amino acids

111 ^a: no overlap between the 90% CI of the upper reference limit of this age group compared to adult dogs

112 ^b: no overlap between the 90% CI of the lower reference limit of this age group compared to adult dogs

113

114 **Precision of the platform**

115 We studied analysis precision using three biologically different dogs, with duplicate aliquots of each dog's sample

116 analyzed once a day during a twenty-day period (Supplementary Table 3). The aliquots from a senior dog suffering

117 from hyperadrenocorticism showed marked chylomicronemia, and was excluded from method precision estimations.

118 Outlier removal slightly affected result interpretation, thus results were evaluated with outliers removed. In the

119 primarily evaluated (non-chylomicronemic) samples, 110/123 biomarkers reached all of their precision goals. Albumin

120 did not reach its primary precision goal of ASVCP CV_{min} based on biological variation, but reached all its other

121 precision goals; CV% goal based on ASVCP Clinical Decision Limits and the S_T goal of under 1/8 reference width

122 (S_{Tmax}). Histidine, acetate, phenylalanine/tyrosine and BCAA/tyrosine met their primary precision goal of CV% under
123 20% (CV_{maxBG}), but did not meet their secondary precision goal of S_{Tmax} . 12/123 biomarkers did not reach their primary
124 precision goal of CV_{maxBG} , but reached the S_{Tmax} goal. All of these analytes were lipid analytes with high inter-
125 individual biological variation and large reference interval width, and were in low concentrations in the tested samples.
126 The CV_{maxBG} goal was thus regarded inappropriate for these analytes, since the variation in these analytes'
127 concentrations was not considered to affect clinical decision-making.

128 Chylomicronemia caused imprecision in multiple lipid measurements as well as in creatinine and the amino acids
129 leucine and phenylalanine. Precision of glutamine was non-evaluable, since the olefin oligomer gel in the sample tubes
130 inhibited correct glutamine estimation from the NMR signal. The precision of XL-VLDL phospholipids was non-
131 evaluable due to a large amount of missing observations caused by physiologically very low analyte concentrations in
132 healthy dogs.

133 **The effect of sample storage on biomarker stability**

134 All evaluated analytes remained stable in refrigerated temperature for 7 days (Table 2, Supplementary Table 4). Most of
135 the analytes remained also stable at room temperature for 7 days. Unstable metabolites in room temperature were the
136 amino acids histidine, isoleucine, glutamine, tyrosine and phenylalanine, acetate, triglycerides and certain lipoprotein
137 particle constituent concentrations as well as the fatty acid docosapentaenoic acid. The stability of all other amino acids
138 than glutamine was better in EDTA-plasma than in serum. Glutamine concentration changed significantly at 4 days of
139 storage at room temperature in EDTA-plasma samples and at 7 days of storage in serum samples. The acceptable
140 change limit (ACL) of glutamine and XL-VLDL phospholipids could not be calculated using the analysis mean
141 coefficient of variation (CVa), since these were not available from the precision study. The changes observed in
142 triglyceride concentrations at room temperature originate from the computational quantitation based on lipoprotein
143 particle distribution and the lipid quantity in the lipoprotein particle core and their outer surface. These are slightly
144 affected by sample storage at room temperature, thus affecting triglyceride quantitation.

145 All of the evaluated analytes were stable at -80°C for 12 months. Storage at -20°C affected certain lipoprotein particles
146 and their composition, certain fatty acids, citrate, acetate and GlycA. Citrate levels were significantly ($p < 0.05$)
147 changed already at one week of storage at -20°C.

148 Heparin plasma storage samples rarely made under the p-value of 0.05, even when the mean percentage deviation
149 (MPD) exceeded the ACL and statistically significant ($p < 0.05$) changes were noted in other sample types.

150 Outlier removal slightly affected result interpretation in serum and EDTA plasma, thus their results were evaluated after
 151 outlier removal. In heparin plasma, outliers could not be reliably identified due to the small sample size (n = 4).
 152 Table 2. Critical storage times for analytes sensitive to certain storage conditions in at least one sample type.

Analyte	Temperature	Storage time _{Serum} (p)	Storage time _{EDTA} (p)	Storage time _{HP} (p)
Glutamine	RT	7 days^a (0.00[*])	4 days^a (0.04[*])	4 days^a (0.11)
Histidine	RT	3 days (0.00[*])	NO	3 days (0.03[*])
Isoleucine	RT	2 days (0.04[*])	NO	4 days (0.20)
Phenylalanine	RT	3 days (0.00[*])	NO	7 days (0.06)
Tyrosine	RT	3 days (0.01[*])	NO	NO
HDL triglycerides	RT	2 days (0.01[*])	4 days (0.07)	4 days (0.20)
LDL triglycerides	RT	2 days (0.01[*])	4 days (0.02[*])	2 days (0.20)
VLDL triglycerides	RT	2 days (0.00[*])	2 days (0.04[*])	4 days (0.11)
Triglycerides	RT	24 hours (0.04[*])	2 days (0.02[*])	2 days (0.20)
Acetate	RT	4 days (0.02[*])	7 days (0.00[*])	4 days (0.03[*])
Docosapentaenoic acid	RT	7 days (0.03[*])	NE	7 days (0.20)
Citrate	-20°C	1 week (0.00[*])	NO	-
Acetate	-20°C	3 months (0.00[*])^b	6 months (0.00[*])	-
Glycoprotein acetyls	-20°C	1 month (0.00[*])	1 month (0.01[*])	-
Omega-3 fatty acids	-20°C	6 months (0.02[*])	NE	-
HDL particle size	-20°C	3 months (0.04[*])	3 months (0.03[*])	-
HDL particles	-20°C	6 months (0.00[*])	6 months (0.00[*])	-
LDL lipids	-20°C	NE	6 months (0.03[*])	-
LDL particles	-20°C	NE	6 months (0.03[*])	-
LDL cholesterol	-20°C	NE	6 months (0.03[*])	-
VLDL particles	-20°C	6 months (0.04[*])	6 months (0.01[*])	-

153 The storage time presented in the table represents the time point, where statistically significant changes (two-sided
 154 Wilcoxon exact test $p < 0.05$), with a mean percentage deviation (MPD) exceeding the acceptable change limit (ACL)
 155 are first observed. In sample types where statistical significance was not observed ($p \geq 0.05$), the storage times represent
 156 the time point, where the MPD first exceeds the ACL. This table includes only primary analytes. Lipoprotein particle
 157 subclasses are not included. Stability at -20°C and -80°C were not studied in heparin plasma. $n_{\text{serum}} = 7$, $n_{\text{EDTA}} = 7$, $n_{\text{HP}} =$
 158 4.
 159

160 RT: room temperature

161 *: $p < 0.05$

162 NO: no change observed

163 NE: not evaluable due to discrepancies in p-values and MPD vs ACL. Results in other sample types should be consulted
164 until further studies are conducted.

165 ^a ACL unknown, evaluated only as $p < 0.05$, time in heparin plasma represents the time, where a change in similar
166 magnitude as in serum and EDTA plasma is observed

167 ^b At 6 months of storage, MPD returned below the ACL and $p > 0.05$

168 **The effect of delayed plasma separation on biomarker stability**

169 We studied the effect of delayed plasma separation in 34 EDTA plasma samples, which had been stored as whole blood
170 in the refrigerator for 24 and 48 hours before separating plasma (Supplementary Table 5, table 3). Outlier removal
171 slightly affected result interpretation, thus the results were evaluated after outlier removal. Prolonged contact with red
172 blood cells (RBC) affected the concentration of many of the analytes. Significant changes ($p < 0.05$, $MPD > ACL$) after
173 one day's contact to RBCs in refrigerator temperature were observed for glucose and lactate. After two days' contact
174 with RBCs, significant changes were observed for citrate, amino acids alanine, histidine, isoleucine, leucine, valine,
175 phenylalanine and tyrosine, cholesterol, triglycerides, and lipoprotein particles and their constituents. Glutamine values
176 were also changed significantly ($p < 0.05$) at 48 hours of storage as whole blood, but MPD could not be evaluated
177 against the ACL due to the missing ACL.

178 Table 3. Critical storage times as whole blood for analytes sensitive to storage as whole blood.

Analyte	Storage time (p)
Glucose	24 hours (0.00)
Lactate	24 hours (0.00)
Citrate	48 hours (0.00)
Alanine	48 hours (0.00)
Glutamine	48 hours ^a (0.00)
Histidine	48 hours (0.00)
Isoleucine	48 hours (0.00)
Leucine	48 hours (0.01)
Valine	48 hours (0.00)
Phenylalanine	48 hours (0.00)

Tyrosine	48 hours (0.00)
HDL lipids	48 hours (0.03)
HDL particles	48 hours (0.00)
LDL diameter	48 hours (0.00)
VLDL diameter	48 hours (0.04)
Cholesterol	48 hours (0.02)
HDL cholesterol	48 hours (0.00)
Esterified cholesterol	48 hours (0.00)
Triglycerides	48 hours (0.00)
VLDL triglycerides	48 hours (0.00)

179 The storage time presented in the table represents the time point, where statistically significant changes (two-sided
180 Wilcoxon exact test $p < 0.05$) together with a mean percentage deviation (MPD) exceeding the acceptable change limit
181 (ACL) are first observed. $n = 34$. This table includes only primary analytes. Lipoprotein particle subclasses are not
182 included.

183
184 ^a ACL unknown

185

186 **Sample tube validation**

187 We studied the differences of seven different sample tubes and the variability between their two sample lots using
188 samples from 20 client-owned dogs (Supplementary Table 6). Outlier removal did not affect result interpretation, thus
189 results were evaluated without outlier removal.

190 Firstly, the different sample tubes were compared to the primary reference tube (Vacuette Lithium Heparin). Most
191 analytes showed comparable results in all tube types compared to the primary reference tube. Significant differences (p
192 < 0.05) with $MPD > ACL$ were noted for citrate, glucose, lactate and GlycA and significant differences ($p < 0.05$) with
193 $MPD < ACL$ were observed for glutamine, histidine, pyruvate and acetate. Glutamine showed significant differences (p
194 < 0.05) compared to the primary reference tube, but its ACL was not available. Most of this variability was observed
195 between analyte values in serum and plasma.

196 Secondly, analyte values in all serum tubes were compared to the reference serum tube (Vacuette Z Serum Clot
197 Activator). All serum tubes showed comparable results with the serum reference tube for all metabolites.

198 The only lot-to-lot variability was observed in MiniCollect Serum gel tubes for glutamine and GlycA. Results obtained
199 by both of these tube lots were still comparable to the reference serum tube (Vacuette Z Serum Clot Activator). In these

200 MiniCollect Serum gel tubes, the two lots represented tubes of old-type (lot A) and new-type (lot B) tubes, with the tube
201 having undergone changes in physical appearance, while sample tube constituents had remained the same.

202 **Interference studies**

203 We studied the interference of hemolysis, lipemia, and unconjugated and conjugated bilirubin using triplicate samples
204 of two different test pools for hemolysis and bilirubinemia and from one pool for lipemia (Supplementary Table 7).
205 Hemolysis interfered with the quantitation of albumin and certain lipids; lipoprotein particle cholesterol concentrations,
206 total triglycerides and certain lipoprotein triglyceride measures, multiple fatty acids, LDL and VLDL particle size,
207 certain lipoprotein particle concentrations, lipoprotein phospholipids and certain lipoprotein lipids. Bilirubinemia
208 interfered only with the quantitation of L-VLDL esterified cholesterol and S-HDL triglycerides. Lipemia affected the
209 quantitation of GlycA due to the lipid contribution to the GlycA signal. However, the concentration of GlycA remained
210 above the reference interval after the removal of lipids, indicating that lipemia was not the sole cause for raised GlycA
211 concentration in the lipemic pool.

212 **Method comparison**

213 Using 999 clinical canine samples, we evaluated, whether results of routine clinical chemistry analytes in the NMR
214 metabolomics platform are comparable with results obtained by conventional clinical chemistry methods
215 (Supplementary Table 8). The tested metabolites were glucose, lactate, creatinine, albumin, cholesterol and
216 triglycerides. The tested samples covered the clinical concentration ranges for all analytes. The comparison plots
217 showed satisfactory agreement between the two methods for all analytes. In some metabolites, logarithm transformation
218 showed better agreement than initial data, suggesting better agreement in the transformed data.

219

220 Glucose, lactate and albumin measurements reached the Total Allowable Error (TE_a) goals based on ASVCP Clinical
221 Decision limits⁴¹, indicating that the two methods can be used interchangeably without affecting clinical decision-
222 making. The bias percentage of creatinine, cholesterol and triglycerides was too high for the analytes to reach the TE_a
223 goals based on Clinical Decision limits, meaning that the two methods cannot be used interchangeably. Values obtained
224 by the conventional method can be evaluated against values obtained by the NMR method using the slope and intercept
225 calculated by Deming regression.

226 **Hyperglycemia associates with many metabolic changes**

227 We evaluated the practical relevance of the NMR metabolomics platform by comparing the metabolic profile of a
228 markedly hyperglycemic group ($n = 24$) to a normoglycemic group of routine laboratory diagnostic samples ($n = 781$).
229 Several biomarkers with previously reported changes in canine and human diabetes mellitus were significantly changed

230 (p < 0.05) in the hyperglycemic group, including the BCAAs isoleucine, leucine and valine, GlycA, acetate, fatty acids,
231 cholesterol, triglycerides, lipoprotein particle concentrations and composition, phenylalanine and lactate
232 (Supplementary Table 9).

233 Excellent fit in logistic regression modeling (smallest AICs and AUC values > 0.9), indicating good predictive value for
234 hyperglycemia, was observed for BCAAs and their ratios. Twenty metabolites had good fit (AUC < 0.8). 31 biomarkers
235 had a high proportion of missing values, reducing the comparability with other metabolites due to different sample
236 populations. There were 14 metabolites with good fit among these metabolites. Many of these metabolites are lipids,
237 that have very low concentrations in healthy dogs, causing a high proportion of automatically rejected very low values
238 by platform quality control.

239

240 **Discussion**

241 Metabolomics is a rapidly growing field holding great potential for numerous clinical and scientific applications.
242 Amongst various approaches, NMR spectroscopy is currently the most promising metabolomics method for clinical use
243 due to its' quantitative nature, high-throughput, accuracy and speed, all of which are currently not achievable with MS
244 based metabolomics. In this study, we describe a novel, cost-effective NMR metabolomics testing platform for dogs. A
245 similar approach has previously been demonstrated and widely utilized in human metabolomics⁴⁰. The testing platform
246 is capable of quantifying 123 biomarkers from a 100 µl sample of serum, heparin plasma, EDTA plasma or citrate
247 plasma. The throughput for one device is ~200 samples per 24 hours and around 70,000 samples per year. The
248 turnaround time is currently 5 days and can be markedly reduced. These characteristics make this comprehensive
249 metabolomics platform a new high-throughput method to facilitate veterinary research and clinical diagnostics with
250 significant implications on treatment, care and well-being of dogs.

251 Recent advances in metabolomics research are paving the way from diagnostics based on single biomarkers to a more
252 holistic approach. Especially the management of chronic metabolic diseases is thought to benefit from a methodology
253 capable of generating comprehensive information on the metabolic state of the individual, enabling a personalized
254 approach to management of the disease³⁴. Metabolomics is also proving itself valuable in disease risk prediction^{33,42-44}.
255 In contrast to genetics, metabolomics offers real-time information of the metabolic state of the animal, taking
256 environmental factors and treatment into account. This offers metabolomics the possibility to be utilized as both a
257 diagnostic tool as well as a follow-up tool. The advantage of the holistic nature of the NMR metabolomics platform was
258 demonstrated in the comparison of hyperglycemic samples to normoglycemic samples from a routine laboratory

259 diagnostic sample population. The NMR metabolomics platform was able to identify a plethora of biomarkers with
260 significant differences between the hyperglycemic group and the normoglycemic group, including BCAAs, GlycA,
261 acetate, fatty acids, cholesterol, triglycerides and lipoprotein particle concentrations and composition, phenylalanine and
262 lactate. Many of these changes have been associated with canine and human diabetes mellitus, and even diabetes risk
263 before disease onset^{16,25,34,42,44}. This suggests the possibility of the NMR metabolomics platform to enhance preventative
264 and precision veterinary medicine by providing new therapeutic targets and prognostic diagnostic markers.

265 In addition to a set of well-known clinical chemistry analytes, the established metabolomics platform includes a wide
266 range of previously clinically unquantified biomarkers. One of the most interesting novel biomarkers is GlycA, a
267 composite systemic inflammatory biomarker comprised of signals of acute-phase proteins α 1-acid glycoprotein,
268 haptoglobin, α 1-antitrypsin, α 1-antichymotrypsin, and transferrin, with a slight contribution of glycosylated
269 apolipoproteins⁴⁵. In humans, GlycA has been linked with systemic inflammation, cardiovascular disease risk, diabetes
270 mellitus, pregnancy, severe infection and all-cause mortality^{28,45-49}. GlycA was also changed in our hyperglycemic
271 group.

272 Formation of reference intervals is an important prerequisite for the clinical use of new laboratory method. We
273 established reference intervals for 123 analytes, most of which are previously unpublished. The reference intervals
274 differed between puppies, adult and senior dogs for many of the analytes, confirming substantial metabolic changes
275 occurring during the maturation of the individual⁵⁰⁻⁵³. The higher reference limit of GlycA in senior than adult dogs
276 suggests a higher prevalence of subclinical disease in older animals. The sample tube validation study demonstrated,
277 that reference intervals created for the same sample type should be used in interpretation of the results. It should also be
278 noted, that glycine and pyruvate cannot be quantified from EDTA plasma, and glutamine is not quantifiable from
279 oligofin oligomer gel tubes.

280 Clinical use of a new laboratory method requires excellent method precision. The precision study suggested, that the
281 precision of the NMR metabolomics platform is generally outstanding. For histidine, acetate, phenylalanine/tyrosine
282 and BCAA/tyrosine, further studies in diseased animals are needed to conclude, whether measurement imprecision
283 affects clinical utilization of these analytes.

284 The analytical method should be considered when evaluating laboratory results against treatment guidelines and
285 reference intervals³⁹. The method comparison study revealed a linear relationship between the conventional and NMR
286 method for all analytes, but all results cannot be evaluated interchangeably. Triglyceride measurements showed most
287 variability between the two methods, explained by differences in methodology, sample handling and biological

288 characteristics. Analysis with the conventional method was done immediately, whereas NMR aliquots were frozen and
289 one batch partly thawed during shipment. The methodology is also different, NMR measures triglycerides within
290 lipoprotein particles, whereas the conventional method is based on interaction with the triglyceride molecule. Lipemia
291 also affects the precision of triglyceride analyses, which causes more variability in high analyte concentrations.

292 Correct preanalytical measures are considered essential for the integrity of laboratory results^{54,55}. An appropriate sample
293 drawing technique is crucial for avoiding hemolysis, fasting before sample drawing for avoiding lipemia and prompt
294 RBC separation for avoidance of prolonged RBC metabolism. Hemolysis, bilirubinemia, lipemia and prolonged contact
295 with RBC all affected the quantification of certain metabolites. The profound impact of chylomicronemia on method
296 precision was caused by matrix heterogeneity due to the chylomicron cream layer formation, which could be reduced by
297 sample mixing immediately before analysis. Glycolysis-related metabolites are especially sensitive to contact with
298 RBC, causing plasma samples to have different concentrations of these analytes than serum samples. Prolonged contact
299 to RBCs affected also amino acids, which might explain why amino acid stability was better in our storage study using
300 plasma and serum samples, than in previous studies using whole blood^{56,57}.

301 It is critical to use sample storage and shipping conditions that preserve the integrity of laboratory results. All tested
302 analytes were stable at refrigerator temperature for one week, making refrigerated temperature the optimal sample
303 short-term storage and shipping temperature. Sample storage at room temperature should be minimized. Most amino
304 were more stable in EDTA-plasma samples than in serum samples in room temperature, which has also been reported in
305 previous studies⁵⁷. The gold standard for long-term storage of serum/plasma metabolomics samples is freezing the
306 samples immediately after serum/plasma separation and keeping them frozen at -80°C until analysis and avoiding
307 additional freeze-thaw cycles⁵⁸. All of the tested analytes remained stable during one year of storage at -80°C. Two
308 weeks of storage in -20°C was suitable for most of the analytes.

309 Limitations of this study include, that the precision and interference studies did not include samples at clinical decision
310 levels for all metabolites. Since the NMR platform is so extensive, acquiring these samples would have been extremely
311 difficult. The sample numbers especially in heparin plasma, were a limitation of the storage study. Limitations of the
312 method comparison study include lack of a reference method generating results that could be viewed as true values of
313 the analyte. No clinical data was available for the hyperglycemia study, thus the study should be repeated with samples
314 in confirmed diabetic animals. The identified analytes should also be studied for multicollinearity.

315 Owing to the advantages of quantitative results, high throughput and reproducibility, the NMR-based metabolomic
316 platform developed and validated here holds great potential for numerous clinical and research applications in

317 veterinary medicine. The performance of the NMR testing platform is generally outstanding and typical blood drawing
318 and processing guidelines ensure the integrity of the results. The diagnostic power of the established metabolomics
319 panel comes from the wide representation of markers from various molecular groups, including amino acids, fatty acids,
320 glycolysis-related metabolites and lipoproteins. This enables efficient monitoring of physiologic changes in health and
321 disease by the identification of condition-specific metabolic fingerprints.

322

323 **Materials and Methods**

324 **Proton NMR spectroscopy**

325 Metabolic profiling was conducted using a proton nuclear magnetic resonance (NMR) metabolomics technique similar
326 to what has been demonstrated for human samples in Soininen et al.⁴⁰. The fully automated process is capable of
327 quantifying 200 samples per 24 hours and the throughput for one device is around 70,000 samples per year. Usable
328 sample types are serum, EDTA-plasma, heparin plasma and citrate plasma. Since the concentration of the tested
329 molecule is in a linear relationship with NMR signal intensity, the linearity of NMR results is considered inherently
330 outstanding.

331 The NMR testing method was calibrated by firstly collecting 1008 canine EDTA-plasma samples and 120 serum
332 samples from client-owned dogs and running the samples with NMR. Only samples with plasma/serum separation done
333 within one day of sample collection were used for initial method calibration, resulting in 847 EDTA-plasma samples
334 and all 120 serum samples. The characteristic NMR signals of amino acids, glycolysis related metabolites, creatinine,
335 and GlycA are well-known, and the quantitative and linear relationship between the signal intensity and molecular
336 concentration is an inherent property of NMR. Thus, these biomarkers can be quantified using the established NMR
337 techniques. The identification of signals for triglycerides, cholesterol, and lipoprotein subclass particles and lipids was
338 verified by analyzing 200 of the previously collected EDTA-plasma samples with different lipid concentrations by high-
339 performance liquid chromatography⁵⁹. For these measures, the concentration range of the automated NMR spectral
340 analysis was also calibrated against these results.

341 Once the initial method calibration was completed, we collected a total of 4683 serum and 495 lithium heparin plasma
342 samples across Finland during fall 2017-fall 2018 for use in our canine NMR metabolomics project. Samples were
343 collected from dogs of all available breeds, with an emphasis of including dogs from genetically and morphologically
344 different breed groups. Samples were drawn by cephalic venipuncture from client-owned dogs. Serum samples were
345 collected into MiniCollect serum gel tubes and were allowed to clot for 30-45 minutes before centrifuging them at 3,000

346 x g for 10 minutes to separate the serum. Heparin plasma samples were collected into MiniCollect lithium heparin
347 plasma gel tubes and the samples were immediately centrifuged at 3,000 x g for 10 minutes to separate plasma. All
348 samples were then stored at -80°C before NMR analysis. All dog owners completed a history form, that included
349 signalment and details about the health status, diet, exercise, stress and reproductive state of the dog.

350 The fatty acid panel verification and calibration was done using 100 samples with different lipid concentrations,
351 collected for the canine NMR metabolomics project. Reference fatty acid concentrations were obtained using a
352 chromatographical method by a commercial laboratory (Vitas as, Norway).

353 **Reference intervals**

354 Determination of reference intervals (RI) of healthy dogs was performed according to the American College of
355 Veterinary Clinical Pathology (ASVCP) reference interval guidelines³⁹. We used the population-based nonparametric
356 method for RI calculation, and calculated 90% confidence intervals (CI) for the reference limits. A minimum of 120
357 samples is required for the use of this method. Inclusion in RI calculation was based on fasting duration (minimum of
358 12 hours), appropriate sample handling, and the lack of owner-reported biological confounding factors, including
359 diseases, severe anxiety/stress and strenuous exercise before blood collection.

360 A total of 865 samples collected for the canine NMR metabolomics project were included in the serum RI calculations.
361 The samples were chosen to include over 120 samples of puppies (under 1 year old), adults (1-7 years old) and senior
362 dogs (over 7 years old), to be able to calculate age-specific reference ranges. The 865 samples consisted of individuals
363 from 68 breeds, 347 males and 517 females, 152 puppies, 545 adult dogs and 168 senior dogs.

364 A total of 269 samples out of the initial 495 samples collected for the canine NMR metabolomics project were qualified
365 for lithium heparin plasma RI calculations. They consisted of individuals from 83 breeds, 155 males and 114 females,
366 29 puppies, 196 adult dogs and 44 senior dogs. Age-specific RI could be calculated for adult dogs.

367 Each metabolite was examined for outliers before RI calculation. We used Box Cox transformation to find a normal
368 transformation, and Horn's algorithm in the transformed data to identify the outliers. In Horn's algorithm, the criterion
369 for rejection of values is exceeding interquartile (IQ) fences set at $Q1 - 1.5 \cdot IQR$ and $Q3 + 1.5 \cdot IQR$ (IQR = interquartile
370 range; $IQR = IQ3 - IQ1$ where IQ1 and IQ3 are the 25th and 75th percentiles, respectively. Once we identified the
371 outliers, the sample and animal data were thoroughly reviewed for confounding analytical and preanalytical factors
372 before reaching a conclusion about sample exclusion. In metabolites, in which normal distribution was not found,
373 Horn's algorithm offered multiple outliers, which all were reviewed before reaching a conclusion about exclusion.

374 After calculating the RIs and the 90% CIs for lower and upper reference limits, we studied the relation between 90%
375 CIs and RI. It is recommended that CI should not exceed 0.2 times the width of the RI, since it may indicate, that the
376 sample number is insufficient.

377 Differences between RI for different age groups were evaluated by comparing the 90% CI of reference limits of puppies
378 and senior dogs to the 90% CI of reference limits of adult dogs. Differences in the reference limits were concluded to be
379 present, when there was no overlap between the 90% CI of reference limits between the two groups.

380 **Precision**

381 Precision of the NMR method was tested using three biologically different dogs (puppy, healthy adult, senior dog
382 suffering from hyperadrenocorticism). Blood was drawn by cephalic venipuncture into Vacuette 8ml Z Serum Sep Clot
383 Activator tubes. Samples were allowed to clot for 30-45 minutes and centrifuged at 2,000 x g for 10 minutes to separate
384 serum. Every sample was divided into 40 aliquots of 100 μ l. All samples were stored at -80°C before NMR analysis.

385 Two duplicate aliquots of each sample were analyzed each day during a twenty-day time period.

386 The sample from the senior dog suffering from hyperadrenocorticism had marked chylomicronemia with a chylomicron
387 coat forming on the top of the stored serum aliquots. The results of this sample were used to determine the effect of
388 chylomicronemia on test precision.

389 Total within-laboratory precision estimates were expressed both as coefficient of variation (CV%) and standard
390 deviation (S_T). These within-laboratory precision estimates were evaluated against laboratory precision goals. We also
391 calculated within-run precision, expressed as within-run standard deviation (S_r). All precision estimates were calculated
392 using methods described in CLSI EP5-A3⁶⁰. Outlier detection was based on comparison of the absolute difference of
393 the two duplicate aliquots analyzed in the same run to the mean absolute difference of all duplicates from the same
394 sample. A difference that exceeded the mean absolute difference fourfold was set as the rejection line. If an outlier was
395 detected, the duplicate was excluded from the calculations.

396 The currently most advisable hierarchy for performance specifications for human diagnostic laboratories is: 1.
397 biological outcomes, 2. biological variation and 3. state-of-the-art^{61,62}. Biological outcome-based performance goals
398 cannot be used in veterinary medicine, since they have not yet been published for animals. Canine CV% goals based on
399 biological variation are available for only a few analytes and can be so strict, that they are unachievable by clinical
400 laboratories and do not affect interpretation of the results⁴¹. State-of-the-art goals can lack accuracy and might not fit all
401 analytes. Due to the lack of specific performance goals for all analytes, we evaluated the precision results against a set
402 of performance goals.

403 In this study, ASVCP goals based on biological variation were set as the primary goals for CV%, since they are the only
404 canine-specific precision goals available⁴¹. The total accuracy of veterinary laboratory results is typically evaluated
405 using Observed Total Error (TE_{obs}) and evaluating it against against preset Total Allowable Error (TE_a) limits. TE_{obs} can
406 be calculated as TE_{obs} = Bias% + 2CV%. Bias was not applicable for this study as a deviation from a true value⁴¹. This
407 led us to use ½ of ASVCP TE_a limits based on Clinical Decision Limits as our secondary precision goals for CV%. If
408 ASVCP goals were not set, we used 20% as the goal for CV% (CV_{maxBG}), which is generally considered an acceptable
409 laboratory error in metabolomics^{63,64}. Since individual CV% max goals were not available for most of the analytes and
410 the 20% CV% goal might not be descriptive enough for metabolites that have a very wide or narrow reference range
411 due to inter-individual biological variation, we additionally compared the S_T to the S_{Tmax} goal of 1/8 of the width of the
412 reference range. This goal was based on the finding, that analytical variation over one quarter of the reference range is
413 considered to affect the clinical interpretation of the results⁶⁵. The S_{Tmax} goals based on 1/8 of the reference range width
414 were very descriptive of the effects of imprecision on clinical interpretation of the results, but have the drawback, that
415 imprecision affects the width of the reference range, itself⁶⁶.

416 **Sample storage study**

417 To define the effects of sample storage and shipping conditions on analyte results, we studied the effects of both long-
418 and short-term sample storage. Short-term storage was studied in room temperature and refrigerated temperature, and
419 long-term storage at -20°C and -80°C. To define storage effects in different sample matrices, we used three different
420 sample matrices: serum, EDTA-plasma and heparin plasma. The protocol of the sample storage study is presented in
421 Table 4.

422 To study storage effects in these above-mentioned sample matrices, we took serum and EDTA plasma samples of seven
423 client-owned dogs in a separate sampling situation, and heparin plasma samples of four dogs in another sampling
424 situation.

425 During the sample collection of EDTA plasma and serum samples for the storage study, blood was drawn by cephalic
426 venipuncture into Vacuette serum gel tubes and Vacuette EDTA K2 tubes. Samples were centrifuged and separated
427 according to the tube manufacturers' recommendations. EDTA-plasma samples were immediately centrifuged and
428 plasma separated. Serum samples were allowed to clot for 30-45 minutes before centrifuging to separate serum.

429 Separated plasma and serum were divided into 24 aliquots and stored in conditions outlined in Table 1 before
430 immediate analysis with proton NMR spectroscopy.

431 Heparin plasma samples for the storage study were collected by drawing blood by cephalic venipuncture into Vacuette
 432 Lithium Heparin tubes. Samples were immediately centrifuged and plasma separated. The samples were divided into 13
 433 aliquots and stored in conditions outlined in Table 4. After storage of samples in these test conditions, all samples were
 434 kept at -80°C for 2-3 weeks before NMR analysis.

435 Table 4. Sample storage study protocol.

Storage time (T_x)	Storage temperature			
	Sample transport simulation		Sample storage simulation	
	Room temperature	Refrigerator	-20°C	-80°C
0 hours (T_0)	S, EDTA, HP			
8 hours	S, EDTA, HP	S, EDTA, HP		
24 hours	S, EDTA, HP	S, EDTA, HP		
48 hours	S, EDTA, HP	S, EDTA, HP		
72 hours	S, EDTA, HP	S, EDTA, HP		
96 hours	S, EDTA, HP	S, EDTA, HP		
1 week	S, EDTA, HP	S, EDTA, HP	S, EDTA	S, EDTA
2 weeks			S, EDTA	S, EDTA
1 month			S, EDTA	S, EDTA
3 months			S, EDTA	S, EDTA
6 months			S, EDTA ^a	S, EDTA
12 months				S, EDTA ^a

436 S: serum

437 EDTA: EDTA plasma

438 HP: lithium heparin plasma

439

440 ^aThe EDTA plasma yield of two included dogs was slightly smaller than expected due to high hematocrit. For both
 441 dogs, one storage time point had to be excluded. The other sample was excluded from the 6 months storage of EDTA
 442 plasma at -20°C and the other one from 12 months storage of EDTA plasma in -80°C.

443

444 We used the two-sided Wilcoxon exact test to estimate the statistical significance of the changes in analyte values
 445 during sample storage. The test was performed between samples not subjected to storage (T_0) and samples subjected to
 446 different storage conditions at each storage time point (T_x) in each storage temperature (Table 4). Two-sided p-values of
 447 Wilcoxon test statistic less than 0.05 indicated a statistically significant difference between metabolite values in T_0 and
 448 T_x .

449 In addition to the Wilcoxon test, we calculated the mean percentage deviation (MPD)

450

$$\text{MPD} = \left(\frac{T_{x\text{mean}} - T_{0\text{mean}}}{T_{0\text{mean}}} \right) \times 100,$$

451 to indicate the stability/instability, as well as magnitude of change in metabolite values over a period of time⁶⁷. MPD

452 was compared to the acceptable change limit (ACL), according to ISO 5725-6⁶⁸:

$$\text{ACL} = 2,77 \text{ CV}_a$$

453 The CV_a was obtained from the precision study results presented in this study, calculated as the mean CV% of the non-

454 chylomicronemic samples. The factor 2,77 was derived from 1.96 x $\sqrt{2}$, where 1.96 represents the 95% of confidence

455 interval for bi-directional changes, and $\sqrt{2}$ was used as we compared two results with the same CV_a. A MPD higher

456 than the ACL represents a probable change in analyte concentration.

457

458 Outliers were detected by duplicate analysis of samples in storage time T_x and collection time T₀. Cut points to set the

459 sample as an outlier were

460

$$\overline{(T_x - T_0)} \pm 2 \cdot \text{STD}_{(T_x - T_0)},$$

461

462 where $\overline{T_x - T_0}$ is the mean and $\text{STD}_{(T_x - T_0)}$ the standard deviation between samples in storage time T_x and collection

463 time T₀.

464

465 If the difference T_{x_i} - T_{0_i} of the sample *i* exceeded or undercut these cut points, it was defined as an outlier. All

466 analyses were done as outliers included or not excluded in purpose to study the effect of outlier exclusion.

467 The criteria for clinically significant change was set so, that the change should be both statistically significant (p-value

468 of the two-sided Wilcoxon exact test <0.05) and the magnitude of the change, evaluated as MPD, was required to be

469 above the ACL, and the change was required to be consistent at the remaining time points. Sample stability was studied

470 only in primary analytes, analyte stability was not evaluated in ratio and percentage analytes.

471 **Delayed plasma separation study**

472 In the delayed plasma separation study, we defined, how delays of plasma separation time from the sample tube
473 manufacturer's recommendations affect metabolite values. Delayed plasma separation was studied by storing samples
474 as whole blood in refrigerated temperature for 24 and 48 hours before plasma separation.

475 Blood was drawn by cephalic venipuncture from 34 client-owned dogs into three Vacuette K2 EDTA tubes per dog.
476 One of the tubes was centrifuged and plasma separated according to sample tube manufacturer's recommendations
477 within one hour after blood sampling. The remaining two tubes were stored as whole blood in a refrigerator until
478 centrifugation and plasma separation after 24 and 48 hours of storage. After plasma separation, all samples were stored
479 at -80°C before NMR analysis. This experiment was conducted using an older version of the NMR testing platform,
480 thus the number of tested analytes was smaller, and fatty acids were not quantified.

481 For statistical analyses, we used similar methods as in the sample storage study. We used the two-sided Wilcoxon exact
482 test to assess the effect of storage time on the sample, and mean percentage deviation (MPD) to indicate
483 stability/instability and magnitude of change. Acceptable change limits (ACL) were the same as in the storage study.
484 Outlier detection followed the same procedure as in the storage study, and criteria for clinically significant change were
485 set similarly as $p < 0.05$ and $MPD > ACL$. T_0 in the delayed plasma study represents plasma separated according to
486 sample tube manufacturer's recommendations and T_x plasma separated after 24 and 48 hours.

487 **Sample tube validation**

488 To study, whether metabolite values differ between samples collected in different blood collection tubes (BCT), we
489 studied both differences between different blood collection tubes and lot-to-lot variability within these tubes. The
490 protocol was modified from the proposed blood collection tube validation process by Bowen and Adcock⁶⁹.

491 Thirty milliliters of blood was drawn from a cephalic iv-cannula from 20 dogs into seven different types of blood
492 collection tubes; MiniCollect Serum Separator Clot Activator tube, Vacuette Z Serum Separator Clot Activator tube,
493 Vacuette Z Serum Clot Activator tube, Vacuette K2 EDTA tube, MiniCollect Lithium Heparin Separator tube, Vacuette
494 Lithium Heparin tube and Vacuette Lithium Heparin Separator tube. Each tube type was tested with two tubes from
495 different lots. EDTA and lithium heparin samples were immediately centrifuged to separate plasma. Serum samples
496 were allowed to clot for 30-45 minutes before serum separation. The centrifugation conditions for samples collected
497 into MiniCollect tubes were 3,000 x g for 10 minutes, and 2,000 x g for 13 minutes for all other sample tube types. The
498 tested sample tubes and their specifications are presented in Table 5. All samples were stored at -80°C before analysis
499 with NMR spectroscopy. The sample tube validation was only conducted regarding primary analytes, not for ratio and
500 percentage analytes.

501 Table 5. Specifications of the tested blood collection tubes in the sample tube validation study.

Blood Collection tube	Sample type	Clotting activator/anticoagulant	Gel type	Lots
MiniCollect Serum Gel Tube 0,8ml	Serum	Clotting activator SiO ₂	Acrylic gel	A: 171010 B: A18023K3
Vacurette Z Serum Separator Clot Activator 3,5ml	Serum	Clotting activator SiO ₂	Olefin oligomer gel	A: A17103HT B: A18024EJ
Vacurette Z Serum Clot Activator tube 2ml	Serum	Clotting activator SiO ₂	-	A: A170837S B: A180644P
Vacurette K2 EDTA Tube 2ml	EDTA-plasma	Spray-dried K2 EDTA 1,2-2 mg (waterless)/ 1 ml blood	-	A: A17063FX B: A18043H3
MiniCollect Plasma Gel Tube 0,8ml	Lithium heparin plasma	Spray-dried lithium heparin 18 I.U./1 ml blood	Acrylic gel	A: A17114HE B: A18013N7
Vacurette Lithium Heparin Tube 2ml	Lithium heparin plasma	Spray-dried lithium heparin 18 I.U./1 ml blood	-	A: A17083HR B: A180247F
Vacurette Lithium Heparin Separator 3ml	Lithium heparin plasma	Spray-dried lithium heparin 18 I.U./1 ml blood	Olefin oligomer gel	A: A17123YY B: A18053D8

502

503 We used the two-sided Wilcoxon exact test and mean percentage deviation (MPD) (detailed in 2.4 Sample storage
504 study) to evaluate statistical significance of differences between the tubes.

505

506 Differences between different sample tubes were tested using this protocol:

507 1) Study, which tubes give comparable results with the primary reference tube

508 The Vacurette Lithium Heparin tube was set as the primary reference tube (T₀) and all other tubes (T_x) were compared
509 against it.

510 Since we hypothesized, that most of the differences between sample tubes would originate from differences between
511 serum and plasma, we secondly studied, which serum tubes give comparable results.

512 2) Study, which serum tubes give comparable results with the serum reference tube.

513 The Vacurette Z Serum Clot Activator tube was set as the serum reference tube (T₀) and all other serum tubes (T_x) were
514 compared against it.

515

516 The average of the analyte values in the two lots was used in these calculations.

517

518 Lot-to-lot variability was tested using this protocol:

519 1) Compare the two lots of each tube to another.

520 Tubes in lot A were set as a reference tubes (T_0) and tubes in lot B (T_x) were compared against them

521 2) If lot-to-lot variability was observed, compare results of each lot to reference tubes. This was done to determine,
522 whether samples from both lots give comparable results to the other tube types.

523 Lots A and B were compared separately to the Vacuette Lithium Heparin and Vacuette Z Serum Clot Activator tubes.

524

525 The same outlier detection protocol was used as in the sample storage study (2.4), except that we used coefficient 3 for
526 STD (instead of 2) in cut points: $(\overline{T_x - T_0}) \mp 3 \cdot \text{STD}_{(T_x - T_0)}$. We set broader cut off limits for outlier detection for the
527 sample tube validation than for the storage study, since the outlier test was more sensitive due to more observations in
528 the sample tube validation than in the storage study.

529 In addition to the tests above, we calculated bias as mean difference of T_x and T_0 with 95% confidence intervals, to
530 highlight the limits in which the differences of the samples in tubes (or lots) T_x and T_0 lie.

531 Although metabolite quantification is possible from citrate tubes, we did not include this tube type in this study, since
532 this tube type is not generally used in clinical applications in this field.

533 **Interference**

534 We tested our metabolites for interference of hemolysis, lipemia, and unconjugated and conjugated bilirubin. The
535 testing protocol was derived from CLSI-EP7-A2⁷⁰.

536 Hemolysis interference was studied using two client-owned dogs. 5 ml heparinized blood was collected by cephalic
537 venipuncture for preparation of the hemolysis stock solution. The samples were centrifuged and plasma separated and
538 discarded. The red blood cells were saved for the preparation of hemolysis stock solution. The hemolysis stock solution
539 was prepared by the osmotic shock procedure. The hemoglobin concentrations of the hemolysates were measured by
540 ADVIA 2120i hematology analyzer (Siemens)⁷¹. The base of the test and control samples were created by collecting
541 5ml heparinized blood by cephalic venipuncture from the same dogs that were used for the preparation of the
542 hemoglobin stock solutions. Test samples containing 500 mg/dl hemoglobin were produced by adding a measured
543 volume of the same dog's hemolysate to the same dog's plasma sample to produce the end concentration of 500mg/dl
544 hemoglobin. Test samples containing 250mg/dl hemoglobin were prepared by mixing an equal amount of the same

545 dog's hemoglobin 500mg/dl test and control samples. All samples were divided into three aliquots and run as triplicates
546 in the NMR analysis.

547 Bilirubin interference was studied by creating two sample pools (Pool 1 and Pool 2) from samples collected for the
548 canine NMR metabolomics project. The sample pools were created so, that as many metabolite levels as possible would
549 differ between the two pools. The baseline bilirubin concentrations of both pools were measured (540 nm) by a
550 modified version of the acid diazo coupling (Malloy-Evelyn) method (Bilirubin Total (NBD), Thermo Fisher Scientific
551 Oy, Finland), using a Konelab 60i chemistry analyzer⁷². The two pools were both divided into four pools, creating the
552 bases for test and control pools for unconjugated bilirubin, and test and control pools for conjugated bilirubin. The test
553 pools for conjugated bilirubin were prepared by adding bilirubin stock solution (201102 EMD Millipore) to the
554 conjugated bilirubin test pool samples. The control pools for conjugated bilirubin were prepared by adding the same
555 volume of water to the conjugated bilirubin control pool samples. The test pools for unconjugated bilirubin were
556 prepared by adding bilirubin stock solution (2011 EMD Millipore) to the unconjugated bilirubin test pools. The control
557 pools for unconjugated bilirubin were prepared by adding the same volume of NaOH to the unconjugated bilirubin
558 control pool samples. All pools were divided into three aliquots and run as triplicates in the NMR analysis.

559 Lipemia interference was studied by pooling lipemic serum samples collected for the canine NMR metabolomics
560 project. The test pool was created so, that the triglyceride concentration would be around 2mmol/l. The pool was
561 divided into a test and control pool. The lipids of the control pool were cleared by ultracentrifugation. Both samples
562 were divided into three aliquots and run as triplicates by NMR.

563 We used CLSI-EP7-A2⁷⁰ guidelines as the base for our statistical testing. Interference testing was only evaluated
564 regarding primary analytes, analyte stability was not evaluated in ratio and percentage analytes. Our testing was based
565 on the point estimate of the observed interference effect (d_{obs}) and the cut-off value (d_c) and the comparison of these
566 values.

567 The interference d_{obs} was defined as difference between the means of the test and control samples:

568 $d_{obs} = \bar{x}_{test} - \bar{x}_{control}$, where test and control samples indicate samples with and without the interfering subject,
569 respectively.

570

571 The cut-off value d_c was defined as $d_c = \frac{d_{null} + s z_{1-\alpha/2}}{\sqrt{n}}$, where d_{null} represented the assumed difference between the

572 means of test and control samples (set to 0), n was the number of replicates ($n=3$) and s was the mean within-laboratory

573 standard deviation (S_T) of non-lipemic samples obtained from the precision study results presented in this study. 95%
574 confidence intervals for the d_{obs} were calculated as

$$575 \quad d_{obs} = \bar{x}_{test} - \bar{x}_{control} \pm t_{0,975,n-1} S \sqrt{\frac{2}{n}}.$$

576 We established our decision of interference in the lower 95% limit of interference limit $d_{obs,low}$ - interference was
577 concluded to be present, if $d_{obs,low} > d_c$ in at least one primary sample/sample pool, since properties of the sample
578 matrix and/or analyte concentrations might affect the presence of interference. Hemolysis interference was concluded to
579 be present if $d_{obs,low} > d_c$ in at least the higher hemolysate concentration of 500mg/dl hemoglobin. Lipemia
580 interference was only evaluated for molecules other than lipids, since the lipid removal process inevitably decreases the
581 concentration of these molecules.

582 **Method comparison**

583 The method comparison study was designed to evaluate, whether results from the routine clinical chemistry analytes
584 included in the NMR metabolomics platform are comparable with results obtained by a conventional clinical chemistry
585 method. The evaluated analytes were glucose, lactate, creatinine, albumin, cholesterol and triglycerides. In this study,
586 bias was only evaluated as a measure of method interchangeability, since the values obtained by the conventional
587 method cannot be regarded as true values of the analyte.

588

589 Clinical canine samples ($n = 999$) were studied by both NMR and standard clinical chemistry analysis methods. The
590 samples included both heparin plasma and serum samples. The used samples were routine diagnostic sample material
591 submitted by veterinarians across Finland, sent by mail to a single laboratory provider (Movet Ltd, Kuopio, Finland).
592 Upon arrival, the samples were divided into two aliquots. One aliquot was immediately analyzed with conventional
593 laboratory methods. Triglycerides were analyzed by a photometric method, cholesterol by a colorimetric method,
594 albumin by a colorimetric method using bromcresol green, glucose by a photometric method using hexokinase,
595 creatinine by a colorimetric enzymatic method and lactate by an enzymatic photometric method. The other aliquot was
596 frozen and kept at -20°C for a maximum of 4 weeks before shipment to NMR analysis. The samples were sent in three
597 batches on cool packs to NMR analysis, shipping time 7-14 hours. After arrival at the NMR laboratory, the samples
598 were again stored at -80°C before NMR analysis. One of the sample shipments was sent over a 14h period on cool
599 packs and partly thawed during shipment. Samples that were sent with a 7h shipment duration remained frozen during
600 transportation. 404 of the samples were used for scaling the NMR albumin measurement, and thus excluded from the
601 method comparison of albumin.

602 Statistical analysis followed the ASVCP General Quality Control Guidelines⁷³. Agreement of the results was viewed
603 using a comparison plot with values the NMR method plotted on the y-axis and the conventional method on the x-axis.
604 Agreement plots were created for raw data and logarithm transformed data. Bland-Altman plots⁷⁴ were used to
605 visualize the distribution of differences.

606

607 We utilized Deming regression^{75,76} in calculation of slope and intercept for all analytes, and in addition, linear
608 regression for analytes with correlation over 0,99. Agreement of Deming transformed data was visualized by adding an
609 agreement plot for data in which conventional method data was transformed using Deming regression coefficients. To
610 determine the mean bias in the unit of the analyte at both ends of the reference intervals, bias was determined by slope
611 and intercept of the Deming regression for these points:

612

$$\text{Bias} = X_c(\text{slope} - 1) + \text{intercept},$$

614

615 where X_c is the corresponding end of the reference interval.

616

617 To determine the interchangeability of results obtained by the conventional and NMR method, TE_{obs} was evaluated
618 against ASVCP clinical chemistry guidelines based on Clinical Decision Limits⁴¹, which reflect the maximum values
619 that would affect clinical decision-making. Observed Total Error TE_{obs} was calculated using the formula⁴¹:

$$TE_{obs} = 2CVa + \text{Bias}\%,$$

620 where CVa was the mean CV% of the non-lipemic samples in the precision study and the mean bias percentage (Bias%)
621 was calculated using the formula⁴¹:

$$\text{Bias}\% = \frac{\text{Mean}_{\text{NMR}} - \text{Mean}_{\text{conventional}}}{\text{Mean}_{\text{conventional}}} \times 100.$$

622

623 **Hyperglycemia-associated biomarkers**

624 In order to demonstrate the utility of the developed method in an important biological question, we studied the
625 differences in metabolite profiles in a hyperglycemic and normoglycemic group. We used the results of the 999 clinical
626 canine samples utilized in the method comparison study as our base population for the study. No clinical data was
627 available for the samples.

628 Diabetes mellitus is defined as a disorder, where blood glucose continuously rises over the renal glucose reabsorption
629 threshold of 11mmol/l, causing glucosuria. The inclusion criteria to the case group (n = 24) was set as a NMR-measured
630 glucose concentration over 11mmol/l. Inclusion to the control group (n=781) was based on the NMR-measured glucose
631 being within the serum all dogs' reference interval (4.4-6.8mmol/l). Since the control group consisted of clinical
632 samples submitted to a clinical laboratory, the control group most likely includes animals suffering from diseases, that
633 do not cause hyperglycemia.

634 To define metabolic changes associated with hyperglycemia, we calculated the means and their 95% CI for the case and
635 control groups as well as the effect sizes of the observed changes. To visualize the differences in analyte concentrations
636 between the two groups, we plotted histograms of the analyte data of the two groups. The significance of the difference
637 between the metabolite values in the case and control group was studied using the two-sided Wilcoxon test's
638 Bonferroni-corrected p-values, with the cut-off for statistical significance set as $p < 0.05$.

639
640 To evaluate the metabolite's ability of predicting hyperglycemia, we used single-variable logistic regression, in which
641 case-control status served as the response variable and individual metabolites as dependent variables. Together with the
642 metabolite's maximum likelihood estimate's p-value (significant < 0.05), area under the curve (AUC) values calculated
643 for the model were used to determine the predictability of hyperglycemia, when the abnormal metabolite value is
644 present. The higher the AUC value is (≤ 1), the better the fit of the model in terms of predicting ability; 0.7 being the cut
645 point for fair, 0.8 for good and 0.9 for excellent model. Akaike information criterion (AIC) was performed in
646 comparison of the models using different metabolites. It has no cut-off point and a lower AIC indicates better model,
647 assuming that the test population in the models is the same. Glucose was excluded from this analysis, since it was used
648 to define the case and control groups.

649
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652

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654

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668 **Author Contributions**

669 HL and CO conceptualized the study. CO, JP, LV, KV and HL designed the study. HL, CO, JP and LV participated in
670 sample recruitment and collection. KV and CO conducted statistical analyses. HL supervised the study. CO drafted the
671 manuscript with the help from other authors. All authors read and approved the final manuscript.

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674 **Author Conflict of Interest Statement**

675 CO is an employee, KV a previous employee, and HL is an owner and the Chairman of the Board of PetBiomics Ltd.

676 **Ethical Approval**

677 All applicable international, national, and/or institutional guidelines for the care and use of animals were followed. All
678 procedures performed in studies involving animals were in accordance with the ethical standards of the institution or
679 practice at which the studies were conducted. Committee: Finnish national Animal Experiment Board, permit number:
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681 **Data availability**

682 The datasets generated during the current study are available from the corresponding authors upon request.

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