Comparison of a point-of-care serum amyloid A analyzer frequently used in equine practice with 2 turbidimetric immunoassays used in human and veterinary medicine

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Abstract. Rapid, accurate detection of serum amyloid A (SAA) is needed in equine practice. We validated a patient-side point-of-care (POC) assay (Stablelab; Zoetis) compared to the turbidimetric immunoassays LZ-SAA (TIA-Hum) and VET-SAA (TIA-Vet; both Eiken Chemical). Analytical performance was assessed at 3 different concentration ranges and with interferences. Inter-method comparison using 49 equine serum samples revealed a significant difference between median SAA results (p<0.0001), with the strongest bias between the POC and TIA-Vet (median 1,093 vs. 578 mg/L). The median SAA value obtained with the TIA-Hum method was 752 mg/L. Correlation between POC/TIA-Hum and between POC/TIA-Vet was −56.7%, −80.9%, and −28.2%, respectively. POC intra- and inter-assay CVs (16.1–30% and 19.8–35.5%) were higher than TIA CVs (generally <12%). Bilirubin and hemoglobin had a negative bias on POC and TIA-Vet results (−16.6 to −45.6%); addition of intralipid yielded a positive bias (35.9–77.4%). The POC had good linearity of SAA concentrations up to 10,312 mg/L (R² = 0.92). A hook effect was present at SAA >3,000 mg/L for the POC assay. Equine serum SAA was stable over a median period of 2.5 y when stored at −80°C. Overall, there was excellent-to-moderate correlation between tests, but imprecision and hook effect of the POC, as well as bias between the methods, must be considered.

Keywords: horses; method validation; point-of-care assay; serum amyloid A; turbidimetric immunoassay.

Serum amyloid A (SAA), the only major acute-phase protein in horses, is a highly sensitive and frequently used marker of inflammation in equine practice.16,23,35 Although systemic SAA in healthy horses is commonly reported as <0.5–20 mg/L,17,26,28 dramatic increases of up to 1,000-fold from baseline reflect inflammatory or infectious processes in diseased horses.4,5 In addition, its rapid increase over 6–12 h and short half-life of 30–120 min31 renders SAA a valuable biomarker to closely track disease severity.7 Different patient-side point-of-care (POC) assays22,29,30 and turbidimetric immunoassays (TIAs) for laboratory use17,18 have been validated for equine SAA measurement.

Although there is no gold standard for measurement of equine SAA,22 a TIA designed for measurement of human SAA concentrations (LZ-SAA, Eiken Chemical; hereafter TIA-Hum) has served as the reference method in several validation studies of equine SAA assays.17,22,29 Although TIA-Hum has proved reliable for equine SAA measurement,17 its drawback is the combination of monoclonal rat anti-human SAA1 antibody and polyclonal rabbit anti-human SAA1 antibody, which provide a high potential for batch-to-batch variation.17,18 Furthermore, TIA-Hum is linear only in a limited concentration range, making several dilutions and repeated analyses necessary to obtain a final SAA concentration.17,18

A novel TIA (VET-SAA, Eiken Chemical; hereafter TIA-Vet), designed for veterinary use and measurement of the wide SAA concentration ranges frequently encountered in horses, is available on the European market. TIA-Vet is based on the sole use of monoclonal rat anti-human SAA1 antibody, and has an acceptable reliability in measurement of extreme SAA concentrations.18 Contrary to TIAs, POC assays can be used stall-side, are user-friendly,22,29 and provide SAA results quickly and without the need for shipment of samples to a reference laboratory.30 To our knowledge, there are 3 POC assays available for equine SAA measurement22,29,30 based on the use of anti-human SAA antibody.22 A study describing the TIA-Hum (Eiken LZ-SAA) and POC assay (formerly Stablelab, Epona Biotech; now Stablelab, Zoetis) found the assay results unsuitable for comparison.20 There is no validation study comparing POC performance to
the TIA-Vet assay. Furthermore, studies on the impact of interferences on POC SAA results are lacking.

Our study aims were 1) to conduct an inter-method comparison between the POC Stablelab EQ1 handheld reader with Stablelab SAA test cartridges and both previously validated TIA assays (TIA-Hum and TIA-Vet) including determination of bias between methods and CVs to calculate the total observed error (TEobs), with the TIA-Hum assay serving as a reference method; and 2) to perform an intra-method comparison including determination of the impact of interferences, linearity, recovery, and hook effect, as well as the impact of storage.

Our hypotheses were, that 1) there is a significant bias between POC and TIA-Hum measurements and an even stronger bias between POC and TIA-Vet measurements, and 2) that equine serum SAA is stable over 2.5 y of storage at −80°C.

Materials and methods

Inter-method comparison

Study design. Study design was developed using the American Society for Veterinary Clinical Pathology (ASVCP) guidelines for quality assurance and method comparison.8 Our prospective study was performed March–December 2018 (validation POC assay vs. TIA-Hum) and January–February 2021 (validation POC assay vs. TIA-Vet). Our study was performed in accordance with the German Animal Welfare Act (Article 8). Ethical approval to use surplus sample volume from equine serum samples submitted for routine testing was given (Regierungspraesidium Giessen, Wetzlar, Germany, ethics committee: JLU_kTV_02_2021).

SAA measurement. Analyses were performed on equine serum samples submitted for SAA analyses to the Department of Veterinary Clinical Sciences, Clinical Pathology and Clinical Pathophysiology, Justus-Liebig-University Giessen, Germany. Samples were analyzed immediately (<1 h after blood collection) with the TIA-Hum test run on a clinical chemistry analyzer (Pentra 400; Horiba). The SAA results obtained during initial routine testing (subsequently TIA-Hum1) served as a reference value to assign the samples to 1 of 3 concentration ranges: SAA 0–300 mg/L (SAAlow), 301–1,000 mg/L (SAAmed), and 1,001–3,000 mg/L (SAAhigh). The 3 categories were only formed for validation purposes to assess the assay performance in different concentration ranges and do not directly correspond to the SAA RI for healthy horses, which is reported to be <20 mg/L.35

Samples were stored in aliquots ≥0.5 mL at −80°C until comparison with the POC assay (subsequently named POC1). TIA-Hum was performed a second time (subsequently TIA-Hum2) at the same time as POC1 and TIA-Vet results were obtained. The use of reagents, samples, and measurements of the POC assay were performed by a single trained person. The TIA assay was performed by trained laboratory technicians. If SAA concentrations exceeded the original working ranges of the TIA-Hum and TIA-Vet assays of 5–500 mg/L and 5–200 mg/L, respectively, automatic 1:6 preanalytical dilutions of 0.005 mL (TIA-Hum) and 0.002 mL (TIA-Vet) of samples were made according to the manufacturer’s instrument settings provided for the Pentra 400 analyzer. Given the extremely high SAA concentrations possible in horses, automatic sample dilution was followed by manual reflex 1:10 or 1:50 dilutions using 0.9% NaCl as required.

Internal quality control (QC) was performed once daily with QC material at 2 different concentrations for both TIA tests run on the Pentra 400 analyzer (TIA-Hum-SAA-QC-low and TIA-Hum-SAA-QC-high for TIA-Hum; VET-SAA-QC-low and VET-SAA-QC-high for TIA-Vet). Concentrations of QC material for TIA-Hum-SAA-QC-low and TIA-Hum-SAA-QC-high were 30.3 mg/L (range: 25.8–34.9 mg/L; 23.7% CV, 6.1 mg/L total bias) and 101.4 mg/L (range: 86.2–116.6 mg/L; 8.9% CV, 11.4 mg/L total bias), respectively. QC target values for VET-SAA-QC-low and VET-SAA-QC-high material were 10.2 mg/L (range: 7.7–12.8 mg/L; 8.3% CV, –1.9 mg/L total bias) and 52.3 mg/L (range: 39.2–65.4 mg/L; 7.1% CV, –6.9 mg/L total bias), respectively. Test calibration was performed using the calibration standards based on human recombinant SAA provided by the manufacturer (LZ-SAA standard Q for TIA-Hum, and VET-SAA calibrator set for TIA-Vet; Eiken Chemical). Although recommended for POC assays,8 POC QC was not possible because QC material was not provided by the manufacturer.

We processed 67 equine serum samples in the described fashion; 49 of 67 sera were used for determination of bias and correlation between methods. The remaining 18 of 67 sera were used for assessment of intra- and inter-assay CVs and the second part of the study, the intra-method comparison.

Inter-method comparison. We analyzed 49 equine serum samples spanning the concentration ranges SAAlow, SAAmed, and SAAhigh with the POC, TIA-Hum, and TIA-Vet tests. After a median storage of 2.5 y (0.3–4.8 y) following initial TIA-Hum1 measurements during routine testing, POC1 and TIA-Hum2 measurements were conducted simultaneously in March–December 2018. TIA-Vet measurements were performed in January–February 2021 (median 2.5 y [range: 2.0–3.8 y] later, when the species-specific TIA-Vet test kits were available). In addition to the mentioned storage time difference, SAA measurements also differ by an additional freeze–thaw cycle performed after the comparison between POC1 and TIA-Hum2.

For statistical analysis, statistical software programs (v.17.8.6, MedCalc Software; Prism 6, GraphPad Software) were used. A p-value ≤0.05 was considered significant. For the inter-method comparison, POC1, TIA-Hum2, and TIA-Vet results were used. Bland–Altman analysis was performed to assess the agreement between methods. Linear regression analysis was performed to assess the correlation between methods. Bias (between method) and precision (within method) was assessed using the mean difference method (MD) and coefficient of variation (CV). The total error (TEtotal) was calculated as the square root of the sum of squares of bias (BIAS) and precision (CV). The total observed error (TEobs) was calculated as the square root of the sum of squares of bias (BIAS) and precision (CV) in the presence of an interfering compound. The limits of agreement (LOA) were calculated as the mean ± 1.96 SE. The correlation coefficient (r) was calculated to assess the strength of the linear relationship between the methods. The significance level was set at p ≤0.05.
performed, and both mean percent bias and mean absolute bias were calculated. Calculation of Spearman rank correlation coefficient was performed, interpreting correlations as “excellent” for Spearman rho ($r_s$) = 0.93–0.99, “good” for $r_s$ = 0.80–0.92, “fair” for $r_s$ = 0.59–0.79, and “poor” for $r_s$ < 0.59, respectively.20,27 Additionally, Passing–Bablok analysis was performed. Before analysis, the data were visually inspected for adequate distribution over the investigated analyte range.

**Precision.** Intra- and inter-assay precision was assessed for each of the 3 assays at 3 SAA concentrations (i.e., SAA$\text{Low}$, SAA$\text{Med}$, and SAA$\text{High}$). Intra-assay variation was determined by analyzing 10 replicates of each concentration range on the same day; for calculation of the inter-assay variation, 1 replicate was run at each test level over 10 consecutive days, using a different single aliquot thawed every day. Intra- and inter-assay CVs, SAA mean (M), and SD were calculated for POC, TIA-Hum, and TIA-Vet assays from replicate measurements.

Imprecision was calculated based on M and SD for both intra- and inter-assay results:

$$\text{CV} \text[%] = \frac{\text{SD}}{\text{M}} \times 100\%$$

Given that no quality requirements are available on the desired CV for equine SAA assays, the observed (analytical) CV ($\text{CV}_a$) was compared to quality requirements derived from biologic variation (i.e., the intra-individual CV [$\text{CV}_i$] of 25% as described for human SAA measurement).

Quality requirements were fulfilled if:

$$\text{CV}_a \text[%] < 0.5 \times \text{CV}_i^{34} \text{; i.e., } \text{CV}_a < 12.5\%$$

**Total observed error.** TE$\text{obs}$ was calculated from intra-assay CV and percent bias (TE$\text{obs}$ = bias% + [2 × CV%]) for the comparisons POC1/TIA-Hum2, POC1/TIA-Vet, and TIA-Hum2/TIA-Vet. Given the absence of veterinary quality requirements for SAA, TE$\text{obs}$ was compared to the total allowable error (TE$\text{a}$) of 37% established for human SAA based on biologic variation.6

**Intra-method comparison**

**Interferences.** To investigate possible interferences, aliquots of a pooled equine serum sample with a mean SAA concentration of ~1,650 mg/L, as assessed with the TIA-Hum test, were used. For assessment of the effect of bilirubinemia, 20 mg of bilirubin (bilirubin ≥98% powder; MilliporeSigma) was dissolved in 1 mL of 0.1 M NaOH to obtain a stock solution of 20 g/L. Subsequently, 0.01 mL of the product was added to 0.24 mL of a non-spiked serum sample to achieve a bilirubin concentration of 0.8 g/L. To assess the effect of hemoglobinemia, a stock solution containing 100 g/L of hemoglobin (hemoglobin from bovine blood, lyophilized powder; MilliporeSigma) was prepared by diluting 30 mg of lyophilized bovine hemoglobin in 0.3 mL of 0.9% NaCl. Then, 0.01 mL of the solution was added to 0.24 mL of non-spiked serum sample resulting in a hemoglobin concentration of 4 g/L. To investigate a possible interference of lipemia, 0.01 mL of intralipid (Lipovenoes 20%; Fresenius Kabi) was added to 0.24 mL of serum sample to obtain a concentration of soybean oil of 8 g/L.

As reported previously,12 control samples were spiked with the same volume and carrier substance as the interferent to avoid a matrix effect resulting from different sample material. The spiked samples were analyzed in comparison to control samples, namely serum aliquots spiked with equal volumes of either 100 mM NaOH (bilirubin), 0.9% NaCl (hemoglobin), or pure double-distilled water (intralipid). All spiked sera and control samples were measured in triplicate with the POC and TIA-Vet tests and then analyzed statistically.

Percent bias between the mean of the test and the control samples was used to calculate the percent observed interference effect (d$\text{obs}$%):

$$d\text{obs}\% = \frac{\text{mean}_{\text{test}} - \text{mean}_{\text{control}}}{\text{mean}_{\text{control}}} \times 100\%$$

The d$\text{obs}\%$ between test and control sample was considered acceptable if it was within ±10%, as described for equine SAA assays.18

**Linearity.** Linearity was evaluated for the POC, TIA-Hum, and TIA-Vet assays by manual stepwise dilution of equine serum samples with increased SAA concentrations based on initial TIA-Hum results of SAA$\text{Low} \sim 190$ mg/L and SAA$\text{High} \sim 15,300$ mg/L for POC measurements, of SAA$\text{Low} \sim 190$ mg/L and SAA$\text{High} \sim 1,600$ mg/L for TIA-Hum measurements, and of SAA$\text{Low} \sim 180$ mg/L and SAA$\text{High} \sim 1,900$ mg/L for TIA-Vet measurements. We performed 2 experiments to assess linearity. First, we assessed linearity within the lower concentration range covering the reportable range of each assay; second, we evaluated linearity within the high concentration range to investigate if linearity is present at high SAA concentrations, for which dilution of an equine SAA sample would be required. Additionally, we chose 2 experiments to minimize dilution error in long dilution series. To assess POC SAA$\text{High}$ linearity, the manufacturer’s dilution protocol for SAA samples with a concentration >3,000 mg/L was used. To assess linearity of all 3 assays, serial dilution resulted in specimens with 1, 0.8, 0.6, 0.4, 0.2, and 0 of the original SAA concentration using the POC manufacturer’s buffer solution for investigation of POC linearity, and 0.9% NaCl for linearity assessment of TIA-Hum and TIA-Vet tests. All diluted aliquots were analyzed in triplicate.

**Recovery.** Linearity under dilution was investigated by visual inspection of the correlation of observed SAA values
plotted against a calculated (expected) SAA concentration. Percent recovery rate (RR) was calculated using the difference between actual and theoretical SAA concentration, as described for canine C-reactive protein (CRP)\textsuperscript{14}:

$$RR\% = \frac{\text{measured concentration}}{\text{expected concentration}} \times 100$$

Linear regression analysis was used to assess correlation between expected and measured results. The range of 80–120% was set as a quality goal for recovery after dilution, as described previously for immunoassay validation.\textsuperscript{1,21}

Hook effect. To investigate a possible hook (prozone) effect for the POC assay, a serum sample with a remarkably high SAA concentration of ~15,300 mg/L based on TIA-Hum analysis after dilution was used. In case of a hook effect, stepwise dilution of the sample was performed to assess the SAA concentration at which a correct detection of high values without a hook effect was possible. Potential dilution steps were 1, 0.8, 0.6, 0.4, 0.2, 0.1, and 0.05 of the original SAA concentration. Each dilution step was measured in triplicate.

The presence of a hook effect was investigated by visual inspection of the observed SAA values as a function of the calculated (expected) SAA concentration. Linear regression analysis was used to assess correlation between expected and measured results. Percent RR over all concentration steps was calculated as described above, and the range of 80–120% was set as the quality goal. \textsuperscript{1,21}

Storage. Given that the duration of storage ranged from 0.3 y at the beginning of the study in March 2018 to 7.3 y at the end of the study in January 2021, the impact of a variable length of storage at −80°C as well as of 2 freeze–thaw cycles on equine serum SAA were investigated at the time of TIA-Vet measurements. The same 10 equine serum samples were re-analyzed a second time with the POC method in January 2021 (subsequently POC2) and with the TIA-Hum a third time (subsequently TIA-Hum3) and compared to previous (TIA-Hum1, TIA-Hum2, and POC1) results. To assess the impact of the variable length of storage on the equine serum SAA samples, POC1, POC2, TIA-Hum1, TIA-Hum2, and TIA-Hum3 results were used. The same 10 equine SAA sera were compared at each time. A Wilcoxon signed-rank test was used to assess differences between POC1 and POC2 measurements. A nonparametric Friedmann test and a Dunn multiple comparison post-test were used for comparison of TIA-Hum1, TIA-Hum2, and TIA-Hum3 results.

Results

Inter-method comparison

There was a significant difference between median SAA measurements obtained with all analyzers ($p<0.0001$). Highest SAA results were obtained with the POC method; lowest SAA results were analyzed with the TIA-Vet assay. Median (minimum to maximum) values obtained were 1,093 mg/L (4–3,000 mg/L), 752 mg/L (0–2,682 mg/L), and 578 mg/L (0.5–2,255 mg/L) for the POC, TIA-Hum2, and TIA-Vet assays, respectively. The dataset used for method comparison covered the complete range of interest with homogeneous data distribution. The correlation between the POC and TIA-Hum assays (Fig. 1A1–C1) as well as between the POC and TIA-Vet assays (Fig. 1A2–C2) was fair, with $r_s=0.77$ and 0.69, respectively. Furthermore, Passing–Bablok regression analysis revealed both constant and proportional bias for the comparison of the POC and TIA-Hum assays as well as between the POC and TIA-Vet assays (Fig. 1A1, 1A2). Contrary to this bias, correlation between SAA measurements obtained with TIA-Hum and TIA-Vet tests was excellent, with $r_s=0.93$ and only a small constant and proportional bias. All 3 assays had a growing absolute bias with increasing SAA concentration (Fig. 1B1–B3). Bland–Altman analysis revealed a proportional bias of −56.7% and −80.9% between SAA measurements obtained with the POC assay and TIA-Hum and TIA-Vet assays, respectively (Fig. 1C1, 1C2). A bias of −28.2% was observed when both TIAs were compared (Fig. 1C3). Percent bias was highest when POC and TIA-Vet tests were compared (Fig. 1A2–C2), and lowest between both TIAs (Fig. 1A3–C3).

Precision. POC intra- and inter-assay CVs mainly exceeded quality requirements for human SAA, except for the intra-assay CV within the medium SAA concentration of 301–1,000 mg/L. TIA-Hum and TIA-Vet assays mainly fulfilled quality requirements (CV\textsubscript{a} < 12.5%) for all 3 concentrations; the only exception was the TIA-Hum inter-assay CV of the SAA\textsubscript{High} measurement. A broad range of intra-assay CVs was observed for the TIA-Hum test, which spanned from 3.1% in group SAA\textsubscript{Med} to 40.2% in group SAA\textsubscript{High} (Table 1).

Total observed error. $TE_{obs}$ of the assay comparisons POC1 versus TIA-Hum2 and POC1 versus TIA-Vet exceeded the $TE_a$ of 37% established for human SAA assays.\textsuperscript{6} $TE_{obs}$ was highest for the comparison of POC1 versus TIA-Vet for all investigated concentration ranges. The lowest $TE_{obs}$ were observed for the comparison TIA-Hum2 versus TIA-Vet, which met the quality requirements of <37% (Table 2).

Intra-method comparison

Interferences. The addition of 0.8 g/L bilirubin, 4 g/L hemoglobin, and 8 g/L intralipid resulted in detectable interferences on POC and TIA-Vet SAA results. The percent bias\textsubscript{obs} exceeded the published criteria of acceptability of ±10%\textsuperscript{18} in all cases. Hemoglobin and bilirubin had a negative bias on SAA results measured by both POC and TIA-Vet assays; intralipid had a positive bias on SAA in both cases. The strongest bias\textsubscript{obs} of 77.4% was seen when a serum sample spiked
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with intralipid was measured with the POC assay. A bias close to the limit of acceptance was detected for the TIA-Vet assay under hemoglobin influence (bias_{obs} = –10.6%; Table 3).

Linearity. The POC test was linear (R^2 = 0.87) when SAA_{Low} samples of 0–562 mg/L were analyzed (Table 4; Fig. 2A). POC linearity was superior (R^2 = 0.92) when SAA_{High} concentrations of 0–10,312 mg/L were measured (Table 4; Fig. 2B). POC SAA results were highly variable within SAA_{Low} and SAA_{High} triplicates (Table 4; Figs. 2A, 2B). For example, POC results were 4,904–10,312 mg/L when the concentration of 0.8 was measured (Fig. 2B). Both the TIA-Hum and TIA-Vet tests had excellent linearity (R^2 = 0.99 each) when SAA_{Low} samples of 0–198 mg/L and 1–115 mg/L, respectively, were measured (Table 4; Figs. 2C, 2E). TIA-Hum and TIA-Vet tests had good linearity (R^2 = 0.95 each) for measurement of SAA_{High} samples of 0–1,593 mg/L (TIA-Hum) and 0–969 mg/L (TIA-Vet; Table 4, Fig. 2D, 2F). When the SAA_{High} sample was measured with the TIA-Vet test and the dilution factors 0.4 and 0.6 were applied, no relevant difference in mean SAA results of 379 and 388 mg/L, respectively, was noted (Table 4; Fig. 2F).

Figure 1. Method comparison. The point-of-care analyzer was compared to 2 turbidimetric immunoassays run on an automated large bench top analyzer (Pentra 400; ABX Horiba). A1–3. Passing–Bablok regression analysis with 95% CIs of serum amyloid A (SAA) measurements performed with the 3 tests. B1–3. Bland–Altman difference plot demonstrating the absolute bias between SAA measurements obtained with the 3 assays. C1–3. Bland–Altman difference plot demonstrating mean percent bias_{obs} with its 95% CI and its 1.96-fold SD indicative of its limits of agreement. Sample stability proven to be >2.5 y. POC1 = point-of-care assay, measured in 2018 (Stablelab; Zoetis); TIA-Hum2 = turbidimetric immunoassay-human, measured in 2018 (LZ-SAA assay; Eiken); TIA-Vet = turbidimetric immunoassay-veterinary (VET-SAA assay; Eiken).
RR did not fulfill the quality criteria of 80–120% reported for immunoassays, especially regarding POC assay performance in both SAA Low and SAA High concentration ranges and almost every dilution step (Table 4). The RR of the TIA-Hum assay fulfilled quality criteria in almost all dilution steps of SAA Low and SAA High measurements, except when the dilutions 0.2 and 0.4 of the original SAA Low concentration were measured. The TIA-Vet test performed below the expected RR quality goals when dilution steps 0.2 and 0.6 of the SAA High sample were analyzed.

**Table 1.** Intra- and inter-assay CVs obtained from replicate serum amyloid A (SAA) measurements with POC, TIA-Hum, and TIA-Vet assays.

| SAA concentration range | Intra-assay CV | Inter-assay CV |
|-------------------------|----------------|---------------|
|                         | Mean (mg/L) | SD (mg/L) | CV (%) | Mean (mg/L) | SD (mg/L) | CV (%) |
| POC                     |              |            |        |              |            |        |
| 0–300 mg/L              | 118          | 19         | 16*    | 86           | 19         | 22*    |
| 301–1,000 mg/L          | 740          | 56         | 8      | 249          | 49         | 20*    |
| 1,001–3,000 mg/L        | 1,532        | 460        | 30*    | 1,534        | 550        | 36*    |
| TIA-Hum                 |              |            |        |              |            |        |
| 0–300 mg/L              | 34           | 1          | 5      | 74           | 6          | 9      |
| 301–1,000 mg/L          | 332          | 13         | 4      | 235          | 7          | 3      |
| 1,001–3,000 mg/L        | 1,073        | 17         | 2      | 2,136        | 860        | 40*    |
| TIA-Vet                 |              |            |        |              |            |        |
| 0–300 mg/L              | 94           | 3          | 3      | 42           | 2          | 5      |
| 301–1,000 mg/L          | 560          | 14         | 2      | 127          | 11         | 8      |
| 1,001–3,000 mg/L        | 865          | 31         | 3      | 1,206        | 140        | 12     |

POC = point-of-care assay (Stablelab; Zoetis); TIA-Hum = turbidimetric immunoassay-human (LZ-SAA assay; Eiken); TIA-Vet = turbidimetric immunoassay-veterinary (VET-SAA assay; Eiken).

* CVs exceeding the published quality requirements for human SAA of 12.5%.

**Table 2.** Total observed error (%) for comparison of the assays POC, TIA-Hum, and TIA-Vet in the 3 investigated concentration ranges.

| TE<sub>obs</sub> | TIA-Hum2 vs. POC1 | TIA-Vet vs. POC1 | TIA-Hum2 vs. TIA-Vet |
|------------------|-------------------|------------------|----------------------|
| SAA<sub>Low</sub> | 66.3*             | 87.5*            | 34.8                 |
| SAA<sub>Med</sub> | 64.3*             | 85.9*            | 33.2                 |
| SAA<sub>High</sub> | 59.9*            | 87.9*            | 35.2                 |

POC = point-of-care assay, measured in 2018 (Stablelab; Zoetis); SAA<sub>Low</sub> = group of low serum amyloid A (SAA) concentrations; SAA<sub>Med</sub> = group of intermediate SAA concentrations; SAA<sub>High</sub> = group of high serum amyloid A (SAA) concentrations; TE<sub>obs</sub> = total observed error; TIA-Hum2 = turbidimetric immunoassay-human, measured in 2018 (LZ-SAA assay; Eiken); TIA-Vet = turbidimetric immunoassay-veterinary (VET-SAA assay; Eiken).

* TE<sub>obs</sub> exceeding the published TE<sub>a</sub> for human SAA of 37%.

**Table 3.** Observed interference effects of bilirubin, hemoglobin, and intralipid on triplicate serum amyloid A (SAA) measurement with the POC and TIA-Vet assays.

| Interferent | POC | TIA-Vet |
|------------|-----|---------|
|            | Mean SAA<sub>control</sub> (mg/L ± SD) | Mean SAA<sub>est</sub> (mg/L ± SD) | Bias<sub>obs</sub> (%) | Mean SAA<sub>control</sub> (mg/L ± SD) | Mean SAA<sub>est</sub> (mg/L ± SD) | Bias<sub>obs</sub> (%) |
| Bilirubin, 0.8 g/L | 1,321 ± 504 | 1,106 ± 100 | −16* | 959 ± 55 | 710 ± 18 | −26* |
| Hemoglobin, 4 g/L | 2,224 ± 940 | 1,211 ± 281 | −46* | 730 ± 10 | 653 ± 12 | −11* |
| Intralipid, 8 g/L | 1,054 ± 179 | 1,870 ± 404 | 77* | 768 ± 18 | 1,043 ± 65 | 36* |

POC = point-of-care assay (Stablelab; Zoetis); TIA-Vet = turbidimetric immunoassay-veterinary (VET-SAA assay; Eiken).

* Results of the observed bias for the interfering substance exceeding 10%.

Recovery. RR did not fulfill the quality criteria of 80–120% reported for immunoassays, especially regarding POC assay performance in both SAA<sub>Low</sub> and SAA<sub>High</sub> concentration ranges and almost every dilution step (Table 4). The RR of the TIA-Hum assay fulfilled quality criteria in almost all dilution steps of SAA<sub>Low</sub> and SAA<sub>High</sub> measurements, except when the dilutions 0.2 and 0.4 of the original SAA<sub>Low</sub> concentration were measured. The TIA-Vet test performed below the expected RR quality goals when dilution steps 0.2 and 0.6 of the SAA<sub>High</sub> sample were analyzed.

Hook effect. A hook effect was observed at SAA concentrations >3,000 mg/L, which exceeded the original working range of the POC assay. A hook effect occurred from dilution step 0.4 of the original SAA concentration (Fig. 3). There was extensive variation of SAA results (minimum to
Table 4. Linearity and recovery rates of serum amyloid A (SAA) measurements performed with equine serum samples originally containing 190 mg/L (SAA\textsubscript{Low}/POC+TIA-Hum), 180 mg/L (SAA\textsubscript{Low}/TIA-Vet), 15,300 mg/L (SAA\textsubscript{High}/POC), 1,600 mg/L (SAA\textsubscript{High}/TIA-Hum), and 1,900 mg/L (SAA\textsubscript{High}/TIA-Vet) SAA as determined by the TIA-Hum method.

| DS | SAA\textsubscript{Low} |   |   |   |   |   |   |   |   |   | SAA\textsubscript{High} |   |   |   |   |   |   |   |   |   |
|----|-----------------|---|---|---|---|---|---|---|---|---|-----------------|---|---|---|---|---|---|---|---|---|
|    | POC            | TIA-Hum | TIA-Vet | POC            | TIA-Hum | TIA-Vet | POC            | TIA-Hum | TIA-Vet | POC            | TIA-Hum | TIA-Vet | POC            | TIA-Hum | TIA-Vet | POC            | TIA-Hum | TIA-Vet | POC            | TIA-Hum | TIA-Vet |
|    | RR (%) | Mean\textsubscript{exp} (mg/L) | Mean\textsubscript{meas} (mg/L) | RR (%) | Mean\textsubscript{exp} (mg/L) | Mean\textsubscript{meas} (mg/L) | RR (%) | Mean\textsubscript{exp} (mg/L) | Mean\textsubscript{meas} (mg/L) | RR (%) | Mean\textsubscript{exp} (mg/L) | Mean\textsubscript{meas} (mg/L) | RR (%) | Mean\textsubscript{exp} (mg/L) | Mean\textsubscript{meas} (mg/L) | RR (%) | Mean\textsubscript{exp} (mg/L) | Mean\textsubscript{meas} (mg/L) | RR (%) | Mean\textsubscript{exp} (mg/L) | Mean\textsubscript{meas} (mg/L) | RR (%) | Mean\textsubscript{exp} (mg/L) | Mean\textsubscript{meas} (mg/L) | RR (%) | Mean\textsubscript{exp} (mg/L) | Mean\textsubscript{meas} (mg/L) |
| 1  | 100  | 512  | 512  | 100  | 194  | 194  | 100  | 113  | 113  | 100  | 8,706 | 8,706 | 100  | 1,575 | 1,575 | 100  | 936  | 936  |
| 0.8 | 68* | 281  | 410  | 106  | 155  | 166  | 96   | 90   | 87   | 78*  | 6,965 | 7,960 | 93   | 1,260 | 1,174 | 102  | 749  | 765  |
| 0.6 | 79* | 245  | 307  | 111  | 116  | 130  | 93   | 68   | 63   | 110  | 5,223 | 4,953 | 116  | 945  | 1,097 | 69*  | 562  | 388  |
| 0.4 | 55* | 113  | 205  | 111  | 78   | 87   | 97   | 45   | 44   | 65*  | 3,482 | 3,433 | 53*  | 630  | 334  | 101  | 374  | 379  |
| 0.2 | 56* | 57   | 102  | 89   | 39   | 35   | 107  | 23   | 24   | 47*  | 1,741 | 1,746 | 63*  | 315  | 200  | 69*  | 187  | 131  |
| 0   | 100  | 0    | 0    | 100  | 0    | 2    | 100  | 0    | 0    | 100  | 0     | 0     | 100  | 0    | 0    | 100  | 0    | 0    |

DS = dilution step; mean\textsubscript{exp} = mean expected (calculated) SAA concentration; mean\textsubscript{meas} = mean measured SAA concentration; POC = point-of-care assay (Stablelab; Zoetis); RR = recovery rate; SAA\textsubscript{High} = group of high SAA concentrations; SAA\textsubscript{Low} = group of low SAA concentrations; TIA-Hum = turbidimetric immunoassay-human (LZ-SAA assay; Eiken); TIA-Vet = turbidimetric immunoassay-veterinary (VET-SAA assay; Eiken).

* RRs below the published quality criteria for immunoassays of 80–120%.1,10
Figure 2. Linearity of diluted equine serum samples measured with the POC, TIA-Hum, and TIA-Vet assays for A. POC measurement of a sample originally containing 190 mg/L serum amyloid A (SAA), B. POC measurement of a sample originally containing 15,300 mg/L SAA, C. TIA-Hum measurement of a sample originally containing 190 mg/L SAA, D. TIA-Hum measurement of a sample originally containing 1,600 mg/L SAA, E. TIA-Vet measurement of a sample originally containing 180 mg/L SAA, and F. TIA-Vet measurement of a sample originally containing 1,900 mg/L SAA. A serial dilution was performed in all cases to achieve 6 different SAA concentrations (i.e., 1, 0.8, 0.6, 0.4, 0.2, 0 parts of the original concentration). Original SAA concentrations were assessed with the TIA-Hum method. POC = point-of-care assay (Stablelab; Zoetis); TIA-Hum = turbidimetric immunoassay-human (LZ-SAA assay; Eiken); TIA-Vet = turbidimetric immunoassay-veterinary (VET-SAA assay; Eiken).
Figure 3. A hook effect was present in POC (point-of-care; Stablelab, Zoetis) assay measurements when serum amyloid A (SAA) values >3,000 mg/L exceeded the original working range of the assay. Linearity under dilution of an equine serum sample originally containing 15,300 mg/L SAA as determined by the TIA-Hum (turbidimetric immunoassay-human; LZ-SAA assay, Eiken) method is shown. The gray box marks the original concentration range of the assay. SAA values reported by the POC assay as “>3,000 mg/L” are replaced with 3,500 mg/L. A serial dilution was performed to achieve 7 different SAA concentrations (i.e., 1, 0.8, 0.6, 0.4, 0.2, 0.1, 0.05 parts of the original concentration).

Storage. The impact of storage on equine serum SAA results was measured with the POC and TIA-Hum tests. Median (minimum to maximum) SAA results obtained were 1,080 mg/L (11–2,430 mg/L) and 925 mg/L (0–3,000 mg/L) when POC1 and POC2 results were compared. As such, no significant impact of storage was observed ($p=0.19$; Fig. 4A). For TIA-Hum1, TIA-Hum2, and TIA-Hum3 measurements, median (minimum to maximum) values obtained were 778 mg/L (5–2,775 mg/L), 744 mg/L (1–2,682 mg/L), and 975 mg/L (9–2,637 mg/L), respectively. Although there was no significant difference between TIA-Hum1 and TIA-Hum2 values, TIA-Hum2 results were significantly lower than TIA-Hum3 results ($p=0.0034$; Fig. 4B).

Discussion

POC, TIA-Hum, and TIA-Vet results cannot be used interchangeably because of significant proportional bias of −56.7, −80.9, and −28.8% for comparison of POC/TIA-Hum, POC/TIA-Vet, and TIA-Hum/TIA-Vet results, respectively, with general overestimation of POC results compared to the reference method. Biases between different test methods and overestimation of POC results in comparison to TIA results have been described: an overestimation of Stablelab POC results in comparison to TIA-Hum results was reported, and a proportional bias of 19.7% was detected when POC and TIA methods for canine CRP measurement were compared.

Several factors contribute to biases between different assay types, such as user-dependent errors (dilution and pipetting errors), inter-batch imprecision, use of different analyzers, and “hardware errors” caused by biochemical assay background reactions. Given that trained personnel performed all measurements in our study, user-dependent errors were minimized. The Pentra 400 analyzer was used for all TIA measurements to exclude performance variations caused by the use of different analyzers. Furthermore, variation of antibody cross-reactivity can be observed between different assay types and between different batches of the same assay, which is a known problem in immunologic acute-phase protein analysis. POCTIA-Hum, and TIA-Vet assays use heterologous anti-human SAA1 antibody to detect equine SAA. However, human TIA antibody cross-reactivity might vary or even be insufficient for detection of animal blood analytes, as shown for measurement of canine CRP with a TIA based on anti-human CRP antibody. Additionally, heterologous calibration of the TIAs based on recombinant human SAA results in expression of equine SAA concentrations in human equivalents and hence different concentration read-outs. Affinity of anti-human SAA antibody toward equine SAA might vary between different assays, with higher affinity translating into a higher SAA concentration read from the calibration curve.

Our results call for the awareness of clinical pathologists and practitioners working with heterologous immunoassays. Even long-established assays may develop variances in cross-reactivity and in test performance and reliability, which is a well-known problem in veterinary laboratories. Especially in assays with known species-dependent batch-to-batch variances, brief validations of every assay batch are recommended. In assays generally known to give reliable results, as in our case, the laboratory may take assay validation studies including 20–40 samples into account if implausible results occur after a new batch has been obtained. Future research should be aimed at improvement of species-specific assays, as in our study.

Both TIA tests had excellent agreement ($r_s=0.93$), but the TIA-Vet method yielded 28.2% lower results compared to the TIA-Hum test in the method comparison study ($p≤0.01$). Even though TIA-Vet is designed for measurement of especially high SAA concentrations, 8% lower SAA results have been described for this method when compared to the TIA-Hum method. These findings can be the result of inter-batch and antibody cross-reactivity variations. Hence, follow-up examinations of the patient should be made based
Point-of-care assay validation for equine SAA on the same SAA assay type. Furthermore, the narrower reported working range of the TIA-Vet test (5–200 mg/L) compared to the TIA-Hum assay (5–500 mg/L) might be explained by the fact that the TIA-Vet assay exclusively uses monoclonal rat anti-human SAA1 antibody, whereas the TIA-Hum test is based on a combination of monoclonal rat anti-human SAA1 antibody and polyclonal rabbit anti-human SAA1 antibody.\(^{17,18}\)

Although POC intra-assay CVs were mainly >12.5%, except in the concentration group SAA\(_{\text{Med}}\) (7.6%), TIA-Hum and TIA-Vet intra-assay CVs (1.6–4.8% and 2.5–3.5%, respectively) fulfilled the quality specifications of CV\(_i\)<12.5% when compared to CV\(_i\) described for human SAA.\(^{21}\) Intra-assay CVs of 13–18% were published for the Stablelab POC assay for SAA concentrations, consistent with the low and medium concentrations of our study, and are comparable to the CVs that we obtained for the same SAA concentration range.\(^{29}\) We observed a markedly higher intra-assay CV of 30% for SAA concentrations in the high concentration range of the POC; however, a comparably high concentration was not assessed in the previous study.\(^{29}\)

Previously obtained TIA-Vet intra-assay CVs of 3–5.2%\(^{18}\) are in agreement with our results and confirm its good intra-assay repeatability in our study.

In agreement with our findings, a high inter-assay CV of up to 45.5% was reported previously for the POC assay and was considered to be the result of the use of different sample types (serum and plasma) and batch-to-batch differences.\(^{20}\) Impact of sample type and of batch, however, was excluded in our study because we used only serum samples and test cartridges of the same batch.

For high SAA concentrations (>500 mg/L), the TIA-Hum inter-assay CV of 40.2% markedly exceeded suggested quality requirements for human SAA. When SAA concentrations exceeded 500 mg/L, automatic dilution steps were performed by the analyzer during each analysis. Thus, errors caused by additional dilution steps contribute to imprecision and are considered the most likely reason for the high inter-assay CV observed for high SAA concentrations that exceed the original working range of the test.

In contrast to the POC, TIA-Vet inter-assay CVs were <12.5% and fulfilled human SAA quality requirements for measurement of high SAA concentrations, which is in agreement with the previous study.\(^{18}\)

Only TE\(_{\text{obs}}\) of the comparison of TIA-Hum2 versus TIA-Vet was less than the TE\(_{\text{s}}\) of 37% based on biologic variation established for human SAA.\(^{6}\) To date, there are no published data on analytical quality requirements for equine SAA,\(^{20}\) hence comparison to TE\(_{\text{s}}\) for human SAA, as well as comparison of CV\(_i\) to human CV\(_i\), can only serve as a rough guideline.

The addition of the interferents hemoglobin, bilirubin, and intralipid resulted in a bias on POC and TIA-Vet results exceeding the accepted deviation of ±10%.\(^{18}\) Investigation of the impact of interferences was warranted given that the POC methodology relies on a color change, and conditions such as hemolysis, hyperbilirubinemia, and lipemia can affect results.\(^{20}\) Although there was no interference effect on TIA-Vet results under the influence of 10 g/L hemoglobin and 5 g/L intralipid in a previous study,\(^{10}\) the addition of 4 g/L hemoglobin, 8 g/L intralipid, and 0.8 g/L bilirubin had a positive percent bias\(_{\text{obs}}\) on POC and TIA-Vet results in the case of

![Figure 4](https://example.com/figure4.png)

**Figure 4.** Impact of storage on equine serum amyloid A (SAA) samples demonstrated with POC assay results and TIA-Hum results obtained at different times after storage. TIA-Hum1 represents the original measurement during routine testing in 2013–2017; TIA-Hum2 and POC1 the measurement in 2018 after a median storage of 2.5 y (range: 0.3–4.8 y); TIA-Hum3 and POC2 results were obtained January 2021 (i.e., a median of 2.5 y after the previous measurement). Box-and-whisker diagram demonstrates median range of the SAA measurements obtained with the 2 analyzers at different times. The horizontal line in the boxes is the median, the whiskers indicate the range, and the box represents the 25th–75th percentile. The asterisks indicate the strength of the p-value: **p≤0.01. A. TIA-Hum results at 3 different times (1–3) were compared. B. POC results at 2 different times (1–2) were compared. POC1–POC2 = point-of-care assay, measured in 2018 and 2021, respectively (Stablelab; Zoetis); TIA-Hum1–TIA-Hum3 = turbidimetric immunoassay-human (LZ-SAA assay; Eiken), measured in 2013–2017, 2018, and 2021, respectively.
intralipid, and a negative percent bias \( \text{obs} \) in the cases of hemoglobin and bilirubin. The different results of our study compared to the reference study may be explained by different spike protocols used by the researchers. Impact of the interferents hemoglobin, bilirubin, and intralipid on equine serum SAA results must be considered when analyzing samples of horses with relevant clinical conditions.

POC linearity was good \( (R^2 = 0.92) \) from 0–10,312 mg/L, and inferior \( (R^2 = 0.87) \) from 0–562 mg/L, with fair agreement between measured and expected results. This is in contrast to a previous publication that describes an acceptable POC linearity up to 1,000 mg/L and less precise results >1,000 mg/L, with POC overestimation of samples >3,000 mg/L compared to TIA-Hum (Eiken LZ-SAA) results.29 The dilution protocol provided by the manufacturer of the POC was considered inadequate to obtain linear results by the authors of the previous method validation study.29 In contrast, the manufacturer’s dilution protocol for the POC led to good linearity up to the highest measured SAA concentrations in our study. The manufacturer’s dilution protocols might have been revised since publication of the previous study.

TIA-Hum and TIA-Vet assays were linear up to SAA concentrations of 1,593 and 969 mg/L, respectively, with overall good agreement of measured and expected results. Minor signs of inaccuracy consisted of slight underestimation of TIA-Hum and TIA-Vet results in the SAAHigh group. A possible explanation for this observed underestimation is the heterologous calibration of SAA assays reporting equine SAA results as human equivalents.18

When the SAAHigh sample was measured with the TIA-Vet test in dilution steps 0.4 and 0.6, respectively, no difference between expected and measured mean SAA results was noted despite the different degree of dilution. The fact that no difference between expected and measured mean SAA results was noted could be explained by a dilution error during preparation of dilution step 0.6 given that all other results closely met the expected values. Clustering of original TIA-Hum results of ~400 mg/L was observed in a previous study when dilutions recommended for human SAA measurement were applied29; a 1:6 dilution was suggested for TIA-Hum SAA measurements >250 mg/L.17,29 In accordance with the previous recommendation, clustering of SAA values was not observed for both TIA-Hum and TIA-Vet in our study in which an automated TIA-Hum 1:6 dilution was made for SAA exceeding the upper measuring range of the test, suggesting appropriate dilution protocols for measurement of equine SAA on the Pentra 400.

RR for the POC, TIA-Hum, and TIA-Vet tests were 47–110%, 53–116%, and 69–107%, respectively, and therefore partially diverged from the quality goals of 80–120% published for immunoassays.1,21 Low RR occurred especially within the higher concentration range of the linearity experiment; pipetting errors could have been introduced by the need for extensive dilution.10

A hook effect was observed at SAA concentrations >3,000 mg/L, exceeding the original working range of the POC assay. This finding is in accordance with the manufacturer’s information. SAA concentrations >3,000 mg/L are frequently encountered in horses.18,33 Relevant changes at very high SAA concentrations cannot be tracked reliably with the present POC assay. However, good linearity \( (R^2 = 0.92) \) was achieved for the same SAAHigh sample when the manufacturer’s dilution protocol was applied in the linearity study. Based on our results, it is advisable to perform repeat SAA measurements using the manufacturer’s dilution protocol when clinical appearance of a horse and SAA results do not correlate and hence may represent falsely low concentrations secondary to a possible hook effect.

A major limitation of our study is that it was conducted over the course of several years, hence making 2 freeze–thaw cycles necessary to obtain separate aliquots. However, the median sample age difference of 2.5 y had no impact on equine serum SAA stability in our study. Although previous studies show that equine serum SAA is stable over 17 d when stored at room temperature and refrigerated at 4°C,11 our study suggests that equine serum SAA is stable for much longer when stored at −80°C. TIA-Hum3 results that were obtained after a median of 2.5 y after TIA-Hum2 results were significantly higher than TIA-Hum2 results \( (p \leq 0.01) \). This is in contrast to other studies that report a general decrease of stored blood analytes.25,32 Furthermore, it has been reported that samples may be degraded by repeated freeze–thaw cycles.5 Given that different TIA-Hum batches were used for TIA-Hum2 and TIA-Hum3 measurements, it is most likely that the spurious SAA increase after storage and repeated freeze–thaw cycles was the result of an inter-batch imprecision with different antibody cross-reactivity of TIA-Hum anti-human SAA antibody to equine SAA, as reported previously.17,18

Acknowledgments

We thank Walter Gilbert and Sabine Zielinsky for their contribution to sample measurements on the benchtop analyzer.

Declaration of conflicting interests

The authors declared no potential conflicts of interest with respect to the research, authorship, and/or publication of this article.

Funding

Stablelab (Epona Biotech) provided part of the POC test kits and the SAA reader used in our study. We received no further financial support for the research, authorship, and/or publication of this article.

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References

1. Andreasson U, et al. A practical guide to immunoassay method validation. Front Neurol 2015;6:179.
2. Arnold JE, et al. ASVCP guidelines: principles of quality assurance and standards for veterinary clinical pathology (version 3.0); developed by the American Society for Veterinary Clinical Pathology’s (ASVCP) Quality Assurance and Laboratory Standards (QALS) Committee. Vet Clin Pathol 2019;48:542–618.
3. Cerón JJ, et al. A seven-point plan for acute phase protein interpretation in companion animals. Vet J 2008;177:6–7.
4. Christensen M, et al. Evaluation of an automated assay based on monoclonal anti-human serum amyloid A (SAA) antibodies for measurement of canine, feline, and equine SAA. Vet J 2012;194:332–337.
5. Coutinho da Silva MA, et al. Serum amyloid A concentration in healthy periparturient mares and mares with ascending placentitis. Equine Vet J 2013;45:619–624.
6. d’Eril GM, et al. Biological variation of serum amyloid A in healthy subjects. Clin Chem 2001;47:1498–1499.
7. Eckersall PD, Bell R. Acute phase proteins: biomarkers of infection and inflammation in veterinary medicine. Vet J 2010;185:23–27.
8. Flatland B, et al. ASVCP guidelines: quality assurance for point-of-care testing in veterinary medicine. Vet Clin Pathol 2013;42:405–423.
9. Fransson BA, et al. Assessment of three automated assays for C-reactive protein determination in dogs. Am J Vet Res 2007;68:1281–1286.
10. Harr KE, et al. ASVCP guidelines: allowable total error guidelines for biochemistry. Vet Clin Pathol 2013;42:424–436.
11. Hillström A, et al. Evaluation of an in-clinic serum amyloid A (SAA) assay and assessment of the effects of storage on SAA samples. Acta Vet Scand 2010;52:8.
12. Hillström A, et al. Validation and application of a canine-specific automated high-sensitivity C-reactive protein assay. J Vet Diagn Invest 2015;27:182–190.
13. Hindenberg S, et al. Evaluation of a species-specific C-reactive protein assay for the dog on the ABX Pentra 400 clinical chemistry analyzer. BMC Vet Res 2017;13:146.
14. Hindenberg S, et al. Evaluation of a novel quantitative canine species-specific point-of-care assay for C-reactive protein. BMC Vet Res 2018;14:99.
15. Ishioka K, Hayakawa N. Serum amyloid A concentrations in cats measured using a newly developed feline-specific latex agglutination immunoassay. Jpn J Vet Res 2019;67:145–150.
16. Jacobsen S, Andersen PH. The acute phase protein serum amyloid A (SAA) as a marker of inflammation in horses. Equine Vet Educ 2007;19:38–46.
17. Jacobsen S, et al. Evaluation of a commercially available human serum amyloid A (SAA) turbidometric immunoassay for determination of equine SAA concentrations. Vet J 2006;172:315–319.
18. Jacobsen S, et al. Validation of an equine serum amyloid A assay with an unusually broad working range. BMC Vet Res 2019;15:462.
19. Jasensky A-K, et al. Evaluation of three different point-of-care tests for quantitative measurement of canine C-reactive protein. Vet Clin Pathol 2015;44:205–214.
20. Jensen AL, Kjelgaard-Hansen M. Method comparison in the clinical laboratory. Vet Clin Pathol 2006;35:276–286.
21. Jensen AL, Kjelgaard-Hansen M. Diagnostic test validation. In: Weiss DJ, Wardrop KJ, eds. Schalm’s Veterinary Hematology. 6th ed. Wiley Blackwell, 2010:1027–1033.
22. Karam B, et al. Whole-blood validation of a new point-of-care equine serum amyloid A assay. J Equine Vet Sci 2020;94:103222.
23. Kjelgaard-Hansen M, Jacobsen S. Assay validation and diagnostic applications of major acute-phase protein testing in companion animals. Clin Lab Med 2011;31:51–70.
24. Klenner S, et al. Evaluation of three automated human immunoturbidimetric assays for the detection of C-reactive protein in dogs. J Vet Diagn Invest 2010;22:544–552.
25. Marjani A. Effect of storage time and temperature on serum analytes. Am J Appl Sci 2008;5:1047–1051.
26. Nunokawa Y, et al. Evaluation of serum amyloid A protein as an acute-phase reactive protein in horses. J Vet Med Sci 1993;55:1011–1016.
27. Papasouliotis K, et al. Comparison of white blood cell differential percentages determined by the in-house LaserCyte hematology analyzer and a manual method. Vet Clin Pathol 2006;35:295–302.
28. Pepys MB, et al. Serum amyloid A protein (SAA) in horses: objective measurement of the acute phase response. Equine Vet J 1989;21:106–109.
29. Schwartz D, et al. Analytical validation of a new point-of-care assay for serum amyloid A in horses. Equine Vet J 2018;50:678–683.
30. Stack JD, et al. Comparison of serum amyloid A measurements in equine synovial fluid with routine diagnostic methods to detect synovial infection in a clinical environment. Front Vet Sci 2019;6:325.
31. Tape C, Kisilevsky R. Apolipoprotein A-I and apolipoprotein SAA half-lives during acute inflammation and amyloidogenesis. Biochim Biophys Acta 1990;1043:295–300.
32. Verma M, et al. Effect of blood storage on complete biochemistry tests for quantitative measurement of canine C-reactive protein. Vet Clin Pathol 2015;65:1011–1016.
33. Westgard JO, et al. Criteria for judging precision and accuracy in method development and evaluation. Clin Chem 1974;20:825–833.
34. Wińkowska-Pilaszwicz OD, et al. Serum amyloid A in equine health and disease. Equine Vet J 2019;51:293–298.