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UPLC-MS/MS method for quantitative determination of the advanced glycation endproducts N^{ε} -(carboxymethyl)lysine and N^{ε} -(carboxyethyl)lysine†

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During blood storage, red blood cells (RBCs) undergo physical, chemical, and metabolic changes that may contribute to post-transfusion complications. Due to the hyperglycemic environment of typical solutions used for RBC storage, the formation of advanced glycation endproducts (AGEs) on the stored RBCs has been implicated as a detrimental chemical change during storage. Unfortunately, there are limited studies involving quantitative determination and differentiation of carboxymethyl-lysine (CML) and carboxyethyl-lysine (CEL), two commonly formed AGEs, and no reported studies comparing these AGEs in experimental storage solutions. In this study, CML and CEL were identified and quantified on freshly drawn blood samples in two types of storage solutions, standard additive solution 1 (AS-1) and a normoglycemic version of AS-1 (AS-1N). To facilitate detection of the AGEs, a novel method was developed to reliably extract AGEs from RBCs, provide Food and Drug Administration (FDA) bioanalytical quidance criteria, and enable acceptable selectivity for these analytes. Ultra-performance liquid chromatography with tandem mass spectrometry (UPLC-MS/MS) was utilized to identify and quantify the AGEs. Results show this method is accurate, precise, has minimal interferences or matrix effects, and overcomes the issue of detecting AGE byproducts. Importantly, AGEs can be detected and quantified in both types of blood storage solutions (AS-1 and AS-1N), thereby enabling long-term (6 weeks) blood storage related studies.

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Introduction

Transfusion medicine is a critical component of modern healthcare, evident by the 10.7 million units of red blood cells (RBCs) transfused annually to patients. The collection of whole blood and subsequent storage of various blood components (such as the RBCs or plasma) into units ready for transfusion is relatively simple. Briefly, the process involves collection of $\sim\!\!450$ mL of whole blood from donors, centrifugation to separate the RBCs from the plasma and leukocytes, followed by storage in separate bags at 4 °C for various lengths of time depending on the component (plasma or RBC) and country regulations. 2,3

A key feature of current protocols for RBC storage is the collection solution into which the whole blood is drawn, and

the solution into which the separated RBCs are stored (the latter also known as additive solution). The most popular collection solution is citrate-phosphate-dextrose (CPD), which contains citrate, phosphate, and dextrose (glucose) at concentrations shown in Table 1. Following centrifugation, the RBCs are then stored in one of multiple available additive solutions (*e.g.*, AS-1, AS-3, AS-5, or AS-7). The contents of AS-1 are also shown in Table 1. While the current blood storage procedure has been in place since the 1970s, there are many reports showing adverse

Table 1 Proposed normoglycemic storage solution constituents compare to FDA approved AS-1 solution

| CPD | CPD-N | AS-1 | AS-1N |
|------|-----------------------------|--|---|
| 129 | 5.5 | 111 | 5.5 |
| 89.4 | 89.4 | | |
| 16.1 | 16.1 | | |
| | | | |
| 15.6 | 15.6 | | |
| | | 154 | 154 |
| | | 2.0 | 2.0 |
| | | 41 | 41 |
| 5.6 | 5.6 | 5.8 | 5.8 |
| | 129 89.4 16.1 15.6 | 129 5.5 89.4 89.4 16.1 16.1 15.6 15.6 | 129 5.5 111 89.4 89.4 16.1 16.1 15.6 15.6 15.4 2.0 41 |

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effects of storage over time on the RBC's chemical and physical properties. These adverse effects, collectively known as the RBC storage lesion, involve chemical, physical, and metabolic changes, as well as functional changes, to the RBC while in storage. Unfortunately, the exact origins of the storage lesion are not known, nor is the mechanism leading to the various changes to the stored RBC well understood.

An interesting feature of the collection and additive solutions used in RBC storage is the high level of glucose in the CPD and the AS-1. Specifically, typical blood glucose concentration ranges from 4-6 mM in a healthy person. 16 Currently, approved versions of CPD and AS-1 have glucose concentrations that exceed 110 mM; even after the RBCs are added, and the AS-1 is diluted due to mixing of the RBCs with the AS-1, the concentration of the glucose in the RBC/AS-1 solution is still in excess of 40 mM, a value much higher than that of healthy humans and humans with diabetes. 10,17 It is noteworthy that after transfusion of the ~280 mL of RBCs into a human, the glucose concentration in the storage bag will not affect the glucose levels in the human transfusion recipient (due to dilution of the 280 mL into a human who typically has a total blood volume of \sim 5 L); rather, the concerning feature of the high glucose is the effect on the RBC properties during storage.

In continuance, people with high blood glucose levels, such as people with diabetes, have RBCs with increased levels of advanced glycation endproducts (AGEs), which are thought to be a negative determinant in overall cell health. 18,19 Past work involving AGEs on the RBCs in the hyperglycemic bloodstream of people with diabetes provided the rationale to investigate the possible formation of AGEs on the RBCs in AS-1. A previous report²⁰ suggests the formation of AGEs later in storage and provides motivation to (1) quantitatively determine the concentrations of the AGEs being formed and (2) evaluate these concentrations of RBC-bound AGEs from the beginning (day 1) to the end of storage duration (>42 days). 15,20,21 Such time-based studies of AGE formation on stored RBCs are without precedent. Here, we describe novel mass spectrometric determination of two AGEs, N^{ε} -carboxymethyl-lysine (CML) and N^{ε} -carboxyethyllysine (CEL).22 The glyoxal mechanism produces CML, the first AGE discovered and the most widely studied AGE.23 Another important AGE, CEL, is associated with diabetes-related complications and derived from the methylglyoxal pathway.24 Measurement of the formation of these two AGEs (CML and CEL) on the stored RBC is the focus of this study to subsequently serve as a tool for determination of improved storage products in transfusion medicine.

Experimental methods and materials

RBC collection

Blood draw followed a protocol approved by the Institutional Review Board of Michigan State University. Blood was obtained from healthy humans and informed consent was obtained from all donors. All record keeping complied with Health Insurance Portability and Accountability Act regulations. Whole blood was collected *via* venipuncture from healthy consenting donors (140 mL total whole blood) into multiple 10 mL uncoated

collection tubes (Thermo Fisher Scientific, Waltham, MA) that were prepared to contain either 1 mL of CPD or 1 mL of a normoglycemic version (CPD-N) (see Table 1). Each blood draw was divided into CPD or CPD-N solution tubes to maintain consistency between storage solution results; that is, each donor had 5 tubes of blood collected in CPD and 5 tubes collected in CPD-N. All whole blood tubes were centrifuged at 2000g for 10 minutes, the plasma and buffy coat (containing the white cells or leukocytes) were removed by aspiration, and packed RBCs (pRBCs) were kept in the tubes.

Glucose concentration and hematocrit percentage

The pRBCs were added to either AS-1 or AS-1N in a 2:1 volume ratio and mixed and stored at 4 °C for one hour before initial glucose concentration and percent hematocrit (the percentage of volume occupied by the RBCs) readings. The glucose concentration was determined with an Aimstrip Plus Blood Glucose Meter (VWR, Radnor, PA) using a 22-gauge needle and 1 mL syringe to collect and transfer a drop of RBC sample onto an Aimstrip Plus Blood Glucose Test Strip (VWR). The glucose reading was repeated, and the average glucose concentration (mg dL⁻¹) was converted to a mM value. The glucose was adjusted to 5.5 mM after one hour for the AS-1N sample by adding an appropriate volume (typically between 50-400 µL) of a 100 mM glucose solution in 0.9% saline. The same volume of 0.9% saline was added to the AS-1 sample to maintain similarity in handling between the AS-1 and AS-1N. The RBC sample hematocrit was determined using a StatSpin MP microhematocrit centrifuge (Beckman Coulter, Brea, CA) and a hematocrit reader (StatSpin CritSpin). The remaining pRBCs were used for analysis either the same day or stored at 4 °C. Periodic feeding of the RBCs stored in AS-1N was achieved using a closed and automated feeding system. Stored samples were removed from bags on day 1, 8, 15, 22, 29, 36, and 43 for UPLC-MS/MS analysis following sample preparation techniques described below.

Protein precipitation for free lysine quantification

To promote cell lysis, packed RBCs (100 μ L) were frozen at $-20\,^{\circ}$ C for one hour and then thawed for 10 minutes at room temperature before mixing with 300 μ L of HPLC grade acetonitrile. The sample was centrifuged at 12 000g for 10 minutes at room temperature, and the supernatant was removed and stored at $-80\,^{\circ}$ C until dried using a SpeedVac (Savant SpeedVac Concentrator, Thermo Fisher Scientific) with an acid vapor trap (Savant Refrigerated Vapor Trap, Thermo Fisher Scientific). The dried sample pellet was stored at $-80\,^{\circ}$ C until prepared for measurement.

Preparation of RBC samples for UPLC-MS/MS

RBC samples (100 μ L) for acid hydrolysis were frozen for one hour at -20 °C and thawed for 10 minutes at room temperature before centrifugation at $10\,000g$ for 10 minutes at 4 °C. The supernatant was removed, and pRBCs were collected for acid hydrolysis. The pRBCs were diluted to make a 1% RBC solution in constant boiling sequencing grade 6 M HCl (Thermo Fisher

Scientific) in glass tubes and transferred to 10 mm, 6 mL vacuum hydrolysis tubes (Thermo Fisher Scientific). The samples were hydrolyzed at 110 °C for 16 hours using a 120 V Digital Dry Bath/Block Heater and Dry Bath Block Insert (Thermo Fisher Scientific). The samples were then removed using customized glassware (MSU Chemistry glass shop) designed to attach a 2 mL Pasteur rubber pipette bulb (Sigma Aldrich, St. Louis, MO) and extract the sample (see Fig. S1 in the ESI†). The glassware was designed using current Pasteur pipette models to extract a sample that would otherwise be unable to be removed cohesively. Using the glassware, the sample was placed into 1.7 mL tubes and frozen overnight at $-80\ ^{\circ}$ C. The following day, the samples were dried to completion using a SpeedVac at 75 °C for 4–5 hours and then kept at $-80\ ^{\circ}$ C until ready for measurement.

Reagent materials and preparation

Immediately prior to measurement, the dried RBC sample used to detect free lysine sample pellet was reconstituted in 300 μL of 10 mM perfluoroheptanoic acid (PFHA) in water, and centrifuged at 13 000g for 10 minutes. The dried, hydrolyzed additive solution RBC samples were reconstituted in 500 µL of 10 mM PFHA in water, and centrifuged at 13 000g for 10 minutes. All supernatant samples were removed and used for measurement or stored at -80 °C. The free lysine sample was diluted 1:100 using 10 mM PFHA in water. Each additive sample supernatant was divided into two categories: CML/CEL detection and lysine detection. The lysine detection samples were diluted 1:40 000 in 10 mM PFHA, and the CML/CEL detection samples were diluted 1:40 in 10 mM PFHA. The sample supernatants were mixed 1:1 (v/v) with an internal standard (IS) mixture containing: 0.1 μ M N^{ϵ} -(1-carboxymethyl)-L-lysine-d₃ (Cayman Chemical, Ann Arbor, MI), 0.1 µM carboxyethyl-L-lysine-d4 (Toronto Research Chemicals, Toronto, ON), and 0.1 μ M 13 C₆, $^{15}\mathrm{N}_2$ labelled L-Lysine (Sigma Aldrich). N^{ϵ} -(1-carboxymethyl)-Llysine (Cayman Chemical), N^e-(1-carboxyethyl)-L-lysine (Cayman Chemical), and ¹³C₆, ¹⁵N₂-lysine (Sigma Aldrich) are used to prepare calibrator samples by dissolving the lyophilized samples in water, diluting with 10 mM PFHA for a ten-point calibration curve, and mixing with the same internal standard mixture as above. The ten-point calibration curve (0, 4, 8, 16, 64, 128, 512, 1500, 2048, 5000 nM) was used to quantify CML, CEL, and lysine by adjusting analyte peak area relative to associated internal standards to illicit an overall response. During initial sample preparation, AS-1 and AS-1N solutions were also hydrolyzed to determine if diluting the samples in reconstituted

sample solution would decrease matrix effects. Quality control (QC) samples were prepared using a stock of RBCs, from a healthy consenting donor, in AS-1 or AS-1N following the procedure above, preparing aliquots of the hydrolyzed sample stock into 1.7 mL vials, drying, and freezing at $-80\,^{\circ}\text{C}$ until they were reconstituted for measurement.

Chromatography and MRM mass spectrometry

To determine if CML, CEL, and lysine were present, the reconstituted hydrolyzed RBC samples were analyzed by UPLC-MS/ MS using a Waters Xevo TQ-S Micro QqQ system interfaced with a Waters Acquity UPLC. A 10 µL aliquot was injected onto a reverse phase column (High Strength Silica (HSS) T3 2.1×100 mm) and the compounds were separated using ion-pair chromatography. Mobile phase A was 10 mM PFHA in water and mobile phase B consisted of 10 mM ammonium formate in a solution consisting of 45% acetonitrile, 45% methanol, and 10% water. The analytes were eluted using the following gradient: initial conditions were 90% mobile phase A and 10% mobile phase B for 5 minutes and then changed to 35% mobile phase A and 65% mobile phase B before switching to 1% mobile phase A and 99% mobile phase B between 5.01 and 6 minutes. Finally, at 6.01 minutes, mobile phase A was 90% and mobile phase B was 10%, and these conditions were held until the 8 minute run was complete. The column was maintained at 40 $^{\circ}\mathrm{C}$ and a flow rate of 0.3 mL min⁻¹. Prior to sample injection, a needle wash was utilized containing 80/20 isopropyl alcohol (IPA)/water with 10 mM ammonium formate. For multiple reaction monitoring mass spectrometry (MRM MS) of 9 channels, the sample was ionized by electrospray operating in positive ion mode (ES+), with a capillary voltage of 1.00 kV (Table 2).

The source temperature was 150 °C, desolvation temperature was 350 °C, desolvation gas flow was 800 L h⁻¹, and cone gas flow was 40 L h⁻¹. CML, CEL, and lysine concentrations were calculated based on the integrated areas relative to the internal standard peak areas. The total AGE concentrations were calculated from measured CML and CEL and adding them together for each sample. The total protein-bound lysine helps evaluate total protein concentration over time, and thus the number of lysines that can be glycated.

Carry-over

For each analyte, there was significant carry-over of analyte signal (>20%), but less than 5% of IS signal. Therefore, after

Table 2 Mass spectrometry for analyte and internal standard compounds

| Analyte compound | Precursor ion mass | Product ion mass | Cone voltage (V) | Collision energy (eV) |
|---|--------------------|------------------|------------------|-----------------------|
| ¹³ C ₆ , ¹⁵ N ₂ lysine | 147.1 | 84.0 | 19.0 | 14.0 |
| N-CML | 205.0 | 84.0 | 15.0 | 22.0 |
| N-CEL | 219.0 | 84.0 | 15.0 | 28.0 |
| ¹³ C ₆ , ¹⁵ N ₂ labelled L-lysine | 155.1 | 90.1 | 19.0 | 14.0 |
| CML-d ₃ | 208.0 | 87.0 | 15.0 | 28.0 |
| $\mathrm{CEL}	ext{-}\mathrm{d}_4$ | 223.0 | 134.0 | 15.0 | 15.0 |

each standard curve, at least 5 blanks (water only) were run before continuing sample analysis. After analysis, 5 blanks were sufficient to remove any remaining analyte eluting (<0.1%). Also, a minimum of two blank samples were run in between each analyte sample (see Fig. S2†).

Data analysis

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Calibration plots of analyte/IS peak area ratio versus CML, CEL, and lysine concentrations were constructed, and a linear regression was used for all analytes. The peak area ratio of sample versus associated IS was used to produce a response to determine the concentrations from the calibration line. SigmaPlot (Systat Software Inc.) was used to plot results and R software (Rstudio version 4.2.2) was used to perform statistical testing in the "rstatix" package (0v.7.2; Kassambara, A. 2023). One-way ANOVA tests for each analyte were conducted to examine differences between storage solution types over 43 days. Days were treated as a repeated measure independent variable and the storage solutions were treated as a betweensubjects dependent variable. Prior to conducting the ANOVA test, the assumptions of normality and homogeneity of variances were assessed using Shapiro-Wilk and Levene's tests, respectively. If these assumptions were met, the ANOVA test was performed. In the case of a significant ANOVA result, post hoc Bonferroni tests were conducted to determine which specific solution types and/or days in storage exhibited statistically significant differences on the 4 analytes.

Results

Intra- and inter-assay accuracy and precision

Intra- and inter-assay variations were assessed utilizing quality control (QC) reconstituted hydrolyzed RBC samples spiked with known concentrations of CML, CEL, or lysine for each batch analysis. The assay variation was determined by reading the plate samples (n = 6) for over 20 hours kept at 4 °C in the sample manager of the autoinjector. Inter-assay variation was determined by measuring the analytes on 4 different samples (n = 4)over a 2 day period at 4 °C in the sample manager. Both intraand inter-assay variations were prepared for the analytes and the results are reported in Table 3.

As seen in eqn (1), the stock QC sample expected concentration (spike concentration) was determined and either diluted

1:40 for CML/CEL or 1:40 000 for lysine. The two spiked concentrations (concentration of spiking solution) relate to a low (10 times the exogenous concentration), and high (40 times the exogenous concentration) concentration of the analytes. Percent recovery (% of target) in eqn (2) was used to determine the accuracy and subsequent percent relative standard deviation (RSD%) in eqn (3) for precision.

$$\left\langle \frac{\text{spike concentration} \times \text{volume of sample}}{\text{concentration of spiking solution}} \right\rangle \tag{1}$$

Percent recovery =
$$\left(\frac{C_{\text{spiked}}[\text{measured}] - C_{\text{nonspiked}}}{C_{\text{spiked}}}\right) \times 100$$
 (2)

Percent relative standard deviation =
$$\left(\frac{\text{std dev}}{\text{average}}\right) \times 100$$
 (3)

The sample accuracy and precision were evaluated at two different concentrations of spiked analyte (low and high) as reported in Table 3. Intra-assay accuracy ranged between 113% and 116% for CML, 93.7% and 97.4% for CEL, and 94.2% and 102% for lysine. All analyte intra-assay accuracy results are in the 80-120% range, which is acceptable for EMA and US FDA bioanalytical guidelines. Each analyte precision was below the EMA and US FDA threshold of 15%. Inter-assay accuracy ranged between 92.8% and 102% for CML, 93.5% and 94.4% for CEL, and 93.4% and 98.8% for lysine. These intra-assay accuracy results are also in the acceptable range for EMA and US FDA bioanalytical guidelines, as is the inter-assay precision.

Linearity, detection limit, and quantification limit

To assess linearity of the developed method, nine calibration standards and a zero standard were measured for each analyte in three experiments; specifically, each standard was diluted using either 10 mM PFHA in water, AS-1, or AS-1N reconstituted supernatant. To test the effect of acid hydrolysis on the reconstitution solution, AS-1 and AS-1N solution containing no sample were hydrolyzed and dried under vacuum. After initial analysis, it was determined that the hydrolyzed sample solutions did not statistically alter the peak shape, retention time, or matrix effects. Therefore, to simplify sample processing, all standards were simply diluted in 10 mM PFHA in water. The

Table 3 Intra- and inter-assay method validation

| | Intra-assay $(n=6)^a$ | | | | Inter-assay $(n=4)^b$ | | | |
|---------|-----------------------|--------------|--------------|-------|-----------------------|--------------|-------------|-------|
| | Accuracy (9 | % of target) | Precision (1 | RSD%) | Accuracy (| % of target) | Precision (| RSD%) |
| Analyte | Low | High | Low | High | Low | High | Low | High |
| CML | 113 | 116 | 4.26 | 4.47 | 92.8 | 102 | 12.69 | 12.2 |
| CEL | 97.4 | 93.7 | 14.3 | 1.87 | 93.5 | 94.4 | 5.25 | 3.58 |
| Lysine | 94.2 | 102 | 3.36 | 1.27 | 93.4 | 98.8 | 8.14 | 7.06 |

^a Intra-assay (six repeated analyses within one experiment during 20 h with samples stored in sample manager). ^b Inter-assay (three independent experiments within 2 days) validation results. Concentration low: ten-times endogenous levels, high: forty-times endogenous levels for all analytes.

Table 4 Linearity, LOD, LOQ, and matrix effects percentage⁶

| Analyte | Linearity (R^2) | LOD (nM) | LOQ (nM) | Matrix effects (%) |
|---------|-------------------|----------|----------|--------------------|
| CML | 0.999 | 1.07 | 3.58 | 7.48 |
| CEL | 0.999 | 0.494 | 1.65 | 8.99 |
| Lysine | 0.999 | 0.345 | 1.15 | 6.33 |

^a Validated method utilized (n=6). CML: N^{ε} -carboxymethyl-lysine (CML); CEL: N^{ε} -carboxyethyl-lysine (CEL). LOQ: limit of quantification (10 × standard deviation_{blankresponse}/slope). LOD: limit of detection (3 × standard deviation_{blankresponse}/slope). Matrix effects calculated from ratio of ((IS area in matrix/IS area in blank sample) -1) × 100. IS: internal standard.

correlation coefficient for standard curves using 10 mM PFHA in water (R^2) are reported in Table 4.

The limit of detection (LOD) and limit of quantification (LOQ) were calculated based on the signal-to-noise ratio of the blank signal and the sample signal as reported in Table 3 for each analyte. The calibration curve of all three analytes using stable IS was measured and indicated good linearity within the concentration range selected (see Table 4). The relative peak area *versus* injected relative concentration to IS was found to be linear with a regression coefficient $R^2 = 0.999$ for all three analytes. The LOD and LOQ were determined using standard deviation of the blank response divided by the slope of the calibration curve multiplied by 3 or 10, respectively.

Matrix effects

There are no commercially available RBC samples in AS-1/AS-1N that are free of the three analytes, so it is impossible to obtain analyte-free biological samples for method validation. However, to adhere to the bioanalytical method guidelines, the hydrolyzed RBC samples were prepared and analyzed to determine the overall extent of suppression or enhancement of signal. The matrix effects were determined using only the IS mix in either sample or blank (water). The matrix effects were calculated using eqn (4).

Matrix effects (%) =
$$\left(\frac{\text{IS peak area [analyte]}}{\text{IS peak area [water]}} - 1\right) \times 100$$
 (4)

According to bioanalytical method validation, less than 13% matrix effects are acceptable criteria. Due to presence of endogenous concentrations of the three analytes tested, the matrix effects were evaluated using IS peak areas. Matrix effects were found to be in all analytes tested (Table 4). All matrix effects were below 13%, which shows there is no significant enhancement or suppression of chromatogram signal due to the blood component matrix. Therefore, the chromatograms can be used to quantify analytes during further analysis.

Absolute quantification of CML, CEL, and lysine

The AGEs under investigation, CML and CEL, were successfully detectable in all blood samples from five healthy controls assayed in biological duplicate, as shown in Fig. 1. While there

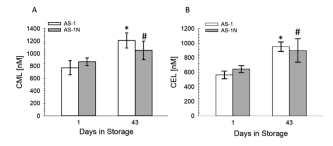


Fig. 1 CML and CEL quantification in AS-1 and AS-1N on day 1 and day 43 in storage. (A) There is no statistical difference between the two solutions (AS-1 = white bars, AS-1N = gray bars) on either day 1 or day 43. However, there is a statistical difference between days 1 and 43 for both solutions for CML levels (**p < 0.01, n = 4–5, error = SEM). (B) There is no statistical difference between the two solutions solutions (AS-1 = white bars, AS-1N = gray bars) on either day 1 or day 43. However, there is a statistical difference between days 1 and 43 for both solutions for CEL levels (**p < 0.01, p = 4–5, error = SEM).

appeared to be no statistically significant differences in AGE formation based on storage solution, there was a significant difference in AGE formation as a function of storage duration for both CML and CEL. The CML concentration in AS-1 on day 1 of storage (774 \pm 113 nM) was not statistically different when stored in AS-1N (869 \pm 61 nM). There was also no statistically significant difference for CML at day 43 of storage between AS-1 (1210 \pm 120 nM) and AS-1N (1052 \pm 149 nM). However, there was a statistically significant increase in AGE formation for both storage solutions across the 43 day storage period (see analysis in Fig. 1 caption). In fact, there was a 53% increase in CML concentration for the RBCs stored in AS-1, a value much higher than the 21% increase in CML formation on RBCs stored in AS-1N. Similar results were measure for the CML, which had

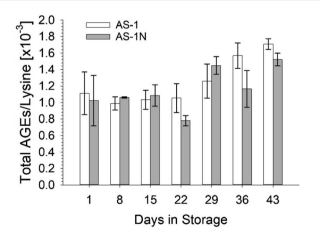


Fig. 2 Total AGEs/lysine [$\times 10^{-3}$] for AS-1 and AS-1N storage solutions over 43 days. Total AGEs (combined CML and CEL) normalized to total protein-bound lysine for AS-1 (white bars) and AS-1N (gray bars) do not show statistically significant differences between the solution types over time. There is a noticeable upward trend from day 1 to day 43 with increasing levels of total AGEs/lysine concentration and show statistically significant differences between day 1 vs. 43 (p < 0.05), day 8 vs. 43 (p < 0.05), day 15 vs. 43 (p < 0.05), day 22 vs. 43 (p < 0.01). n = 4–5, error = SEM.

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a concentration in AS-1 of 565 \pm 53 nM on day 1 and a concentration of 953 \pm 65 nM on day 43. The CML formation on the RBCs stored in AS-1N increased from 643 \pm 49 nM on day 1 to 901 \pm 161 nM on day 43. The percent increase for the CEL in AS-1 was higher (69%) than the AS-1N stored cells (40%).

The data in Fig. 2 represent the total increase in AGEs for each storage solution during the entire storage duration (43 days) relative to the lysine concentration. Similar to the individual determinations of CML and CEL, the storage solutions did not seem to have an effect on the AGE formation. Furthermore, although the total AGEs clearly increased as a function of storage duration, there was not a large difference in AGE formation from day 1 to day 43 (total AGE change was 54% for AS-1 and 49% for AS-1N). However, an increase in AGEs (relative to day 1 of storage) were measured after 3 weeks of storage.

Discussion

This study includes a new protocol with high accuracy and precision for the quantification of two types of AGEs using UPLC-MS/MS. In the present study, UPLC-MS/MS with MRM was highly useful for the measurement of AGEs in blood samples. Specifically, we were able to report the concentrations of CML, CEL, and lysine in two different blood samples on day 1 through day 43 of storage. This study focused on CML and CEL without the addition of common byproducts because these are the most abundant and widely studied AGEs related to complications associated with diabetes. Due to the hyperglycemic storage conditions used in RBC storage for transfusion medicine, it was anticipated that the stored RBCs would also exhibit AGE formation. There is limited research for detecting and quantifying both CML and CEL, and almost no present research involving stored blood. Current commercial kit assays are not selective enough for only CML or CEL without other glycation mechanism byproducts. Current methods of detecting and quantifying AGEs include nonselective enzyme linked immunosorbent assay (ELISAs), which use antibodies and are often limited to reliable types of antibodies and decreased sensitivity.22 Other types of analytical methods include time-consuming immunohistochemical detection, fluorescence spectroscopy and size exclusion chromatography with fluorescence detection.25-27 Thus, we developed a reliable, sensitive, specific (see ESI Fig. S3†), and reproducible method to detect CML, CEL and lysine in human blood samples.

Other challenges when developing a robust method for detection and quantification of AGEs, include the ten-thousand-fold difference between the concentration of protein-lysine on RBC membranes *versus* modified lysine adducts (CML and CEL); only 1% of RBC proteins that contain lysine are glycated and form AGEs.^{23,28} Acid hydrolysis is a well-known technique that can be used to extract proteins and amino acids on the RBC membrane to adequately prepare various biological samples (urine, pRBCs, plasma) for high throughput analysis such as UPLC-MS/MS.^{29,30} There are many drawbacks with previous methodology reports, such as biological interferences and matrix effects enhancing or suppressing chromatographic signal.³¹⁻³³ There are several methods that report CML, CEL,

and/or lysine detection for AGEs studies utilizing various analytical techniques.^{34–36}

The bioanalytical method validation by the European Medicines Agency (EMA) provides guidance and recommendations for bioanalytical assays, which can be seen as the "gold standard" in other types of method development.37 By using the FDA approved guideline M10 by the International Council for Harmonization of Technical Requirements for Pharmaceuticals for Human Use (ICH), this study used sample analysis recommended chemical, biological, and metabolite drug guidelines to explore various analytical parameters, such as LOD, LOQ, matrix effects, intra- and inter-assay accuracy and precision, and matrix interferences. The EMA bioanalytical method validation guidance provides clear acceptance and reliability for biological assays and analytical results. Through this process, our results showed acceptable linearity, LOD, and LOQ. The matrix effects did not contribute to chromatographic signals, which is a major concern for biological samples. There was clear specificity, requisite peak resolution, and distinguishable IS correlation to the analyte detected. The intra- and inter-assay results showed EMA and US FDA bioanalytical guideline acceptable accuracy and precision. Overall, the results showed this methodology to detect and quantify CML, CEL, and lysine in stored blood solutions can be used in other analyses involving suspected AGE formation on blood components.

To date, most reports reporting AGE formation have primarily focused on AGEs in food, biological plasma or serum, or human tissues (retina, kidney, endothelial and smooth muscle cells).24,27,38-45 Increased protein glycation has been reported in clinical studies involving people with diabetes, and it is linked to various complications associated with increased AGEs and oxidative stress. 19,46-48 Glycation involves the Maillard reaction, a non-enzymatic process where the carbonyl group of a reducing sugar reacting with the amino group of a protein, lipid or nucleic acid, generates Schiff bases to produce Amadori products.22 These Amadori products, as well as other byproducts and intermediates during glucose oxidation, lead to the formation of various AGEs, such CML and CEL.22 The mechanism that produces CML was the first discovered and is the most widely studied AGE.23 Another important AGE, CEL, is associated with diabetic complications and derived from the methylglyoxal pathway (see ESI Fig. S4†).24 Although glycation occurs on most cell types, AGE detection of bloodstream such bloodstream components as proteins and intact cells may provide insight into increased pathologic conditions.28,49 For example, glycated albumin in the bloodstream has been shown to influence delivery of biologically active peptides to healthy RBCs, while that same delivery was reduced in RBCs from people with diabetes. 50,51

Here, there were no statistically significant differences in the two blood storage solutions on a given day, although there were some interesting features of the collected data. For example, while there was no difference in AGE formation between the AS-1 and AS-1N storage solutions on any particular day, there was a significant increase in AGE formation on the RBCs during storage duration for both storage solution strategies. Another interesting feature of the data shown in Fig. 2 is the increase in total AGE formation after 2–3 weeks in storage. While the

concept of "fresh" stored RBCs do not perform any better than "older" stored RBCs is debatable, it is clear in the literature that fresh RBCs are preferred in certain situations (e.g., infant transfusions). 52,53 Furthermore, some reports suggest a reduction in post-transfusion complications when using RBCs stored for less than 2 weeks. 54,55 The data in Fig. 2 show that the covalent AGE formation experiences an increase in formation rate around this 14–21 day period. These results correspond closely to our previous results showing irreversible damage to the RBCs stored in AS-1 after \sim 2 weeks. 56

Conclusions

AGEs are known to be related to the pathomechanism of diabetes and other degenerative disorders. This study involved utilizing blood from healthy donors to explore methodology that can reliably and accurately detect and measure AGEs, while maintaining sensitivity and bioanalytical merits to other reports using different instruments or assays. We were able to confirm the detection and quantification of CML, CEL, and lysine present on stored RBCs, without significant issues of matrix effects and with analyte recovery. The purpose of this small initial study was to validate the robustness of the biomarkers and provide a pilot study for future blood banking AGE research. The data encourages further investigation of the accumulation of AGEs.

Author contributions

The idea of the experiment belongs to M. G. and L. A. S. S.; M. G. and L. A. S. S. planned and organized the work. L. D. S. performed the blood collection and L. D. S. and L. A. S. S. performed blood preparation; L. A. S. S. and M. G. developed TQ-S Micro sample preparation methods and protocol; L. A. S. S. conducted the experiments, developed the quantitative method, and interpreted the UPLC-MS/MS data; L. A. S. S. performed calculations and revisions; L. A. S. S., L. D. S., M. G., A. S., and D. S., helped in the writing of the manuscript.

Conflicts of interest

There are no conflicts to declare.

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