ABSTRACT

Purpose. An improved HPLC method with fluorescence detection was developed and validated for determination of glucosamine in human and rat biological samples. Method. Aliquot of 0.1 mL plasma was spiked with mannosamine HCl as the internal standard (IS); proteins were precipitated with acetonitrile; the clear layer was derivatized with 9-fluorenylmethyl chloroformate (8 mM/acetonitrile) in presence of borate 0.2 M buffer at 30°C for 30 min. The excess derivatizing agent was removed with 1-amino adamantane HCl (300 mM in acetonitrile-water 1:1). Chromatographic separation was achieved on a C18 (100mm X 4.6 mm, id 3μm) reversed phase column using 0.1% acetic acid/acetonitrile gradient mobile phase at 1 mL/min flow rate. Glucosamine was determined in the plasma of a human and rats and also in rat urine. Results. The analytes were detected at excitation and emission wavelengths of 263 and 315 nm, respectively. The assay was linear over the range of 0.05-20 μg/mL with a typical correlation coefficient of 0.999 and intra-day and inter-day coefficient of variation of <15%. The lowest limit of quantification was set at 50 ng/mL. The recovery for glucosamine and mannosamine was 98 and 96%, respectively. Conclusion. We were able to improve glucosamine assay suitable to quantify glucosamine in both human and rat plasma and rat urine.

INTRODUCTION

As an amino sugar the compound lacks chromophores, hence, is undetectable in the ultraviolet-visible ranges. Early studies on glucosamine pharmacokinetic have been carried out using radio labeled compound which could not distinguish between the parent compound and its biosynthetic or degraded products (7). The first reported high performance liquid chromatography (HPLC) methods for determination of glucosamine in plasma were based on derivatization of glucosamine with phenylisothiocyanate (8) or 1-naphthyisothiocyanate (9) to form a compound that can be detected in the UV region. Those methods, however, lack sufficient sensitivity in the presence of human plasma (> 1.25 μg/mL). More recently, sensitive liquid chromatography methods have been reported that involve detection of intact or derivatized glucosamine product by electrospray
ionization with mass spectroscopy (10-15). Suitable instruments for these assays, however, are not commonly available. In 2006, Zhang et al reported an HPLC method with fluorescence detection for determination of glucosamine in human plasma that involved derivatization with 9-fluorenlymethoxycarbonyl chloride (FMOC-CL) without using an internal standard (Figure 1) (16). The derivatization yields two anomers (stereoisomers of a cyclic sugars that differ in their configuration at the anomeric carbon) peaks (16). This method was later modified by Huang et al who added vertilmicin sulfate as an internal standard (17). The minimum quantifiable concentration of this method was 100 ng/mL. Our attempts to utilize this method, however, failed due the presence of broad interfering peaks and elevation of the HPLC pump pressure following repeated injections. The goal of this work was to improve and optimize the reported method for determination of glucosamine in the rat and human biological fluids.

EXPERIMENTAL

Material and Reagents
Glucosamine HCl, mannosamine HCl, amantadine HCl (1-aminoadmantane HCl, ADAM) and FMOC-CL were purchased from Sigma-Aldrich Canada, LTD, (Oakville, ON, Canada), HPLC grade acetonitrile and water were obtained from Caledon Laboratories Ltd, (ON, Canada).

Solutions and standards
Glucosamine stock solution was prepared by dissolving 60 mg glucosamine HCl in 100 mL water to yield a 0.5 mg/mL solution of glucosamine free base. The stock solution was further diluted with water to give standard solution containing 0.05, 0.1, 0.25, 0.5, 1, 2.5, 5, 10 and 20 µg/mL. Mannosamine was used as the internal standard (IS) for which a stock solution was prepared by dissolving the compound in water to give 10 µg/mL. The derivatizing reagent solution (8 mM) was prepared freshly by dissolving 10.3 mg FMOC-CL in 5 mL acetonitrile. Borate buffer 0.2 M was prepared by dissolving 6.18 g boric acid in 425 mL water followed by pH adjustment to 8.5 using 10 M NaOH. A 300 mM solution of ADAM was prepared by dissolving 280 mg in 5 mL 1:1 (v/v) acetonitrile-water.

Sample preparation and derivatization
Aliquot of 0.1 mL human or rat plasma was spiked with 50 µL of 10 µg/mL IS. Plasma proteins were precipitated by addition of 200 µL acetonitrile followed by 1 min vortex-mixing and centrifugation for 3 min at 10,000 g into clean test tubes; 100 µL of the supernatant were transferred and 50 µL of borate buffer were added followed by addition of 50 µL of freshly prepared FMOC-CL. This was followed by 1 min vortex-mixing and incubation in a water bath at 30°C for 30 min. After incubation, 50 µL of ADAM was added to the test tubes to react with the excess derivatizing agent. The samples were then diluted with 1 mL acetonitrile-water (1:1) and 5 µL injected into the HPLC.

Chromatographic conditions
A Shimadzu prominence HPLC system (Mandel Scientific, Guelph, ON Canada) consisting of a DGU-20A5 degasser, a LC-20AT pump, a SIL-20A autosampler, a CTO-20AC column oven, a RF-10AXL fluorescence detector and a CBM-20A communication bus module was used. The integration was performed using Shimadzu Class-VP 7.4 software. Chromatographic separation was achieved on Phenomenex C18 (100mm X 4.6 mm, id 3µm) reversed phase column guarded with Phenomenex Security Guard Cartridge C18 (4mm x 3 mm) column both purchased from Phenomenex.
(Torrance, CA USA). The mobile phase consisted of 0.1% glacial acetic acid in HPLC-grade water (A) and acetonitrile (B). A gradient elution was programmed to commence with 23% B for 13 min post-injection followed by gradual increase in 2 min of B to 90%. The composition was maintained for 8 min when was gradually decreased back to 23% of B in 2 min. The flow rate was 1mL/min and the column oven temperature was set at 40°C. The detection was carried out at excitation and emission wavelength of 256 nm and 315 nm, respectively. The sample run time was 35 min. The peak area was used on all calculations.

Validation
Calibration curves were prepared by spiking aliquots of 0.1 mL plasma with glucosamine to yield standard samples containing 0.05, 0.5, 5 and 20 µg/mL of the compound. The standard curves were constructed by plotting glucosamine:IS peak area ratio versus the added concentration of glucosamine. Three calibration curves were constructed in the same day to determine intraday variability. The method was repeated on three different days to determine the interday variability. The accuracy was determined from % error = (mean observed concentration – added concentration) x100/ added concentration. The coefficient of variation (CV%) was used to estimate the assay precision.

Recovery
Solutions of 0.05, 0.5, 5 and 20 µg/mL glucosamine were prepared in plasma and in water and analyzed as mentioned before, each in triplicate. The % recovery of glucosamine from plasma was estimated from % recovery = (glucosamine peak area in plasma sample /glucosamine peak area in water sample) x 100. The recovery of mannosamine from plasma was also determined using the same approach.

Stability
Short term stability. Stability of the samples during analysis was tested using 4 standard samples of glucosamine in plasma (0.05, 0.5, 5 and 20 µg/mL). Samples were prepared and derivatized as mentioned before. They were analyzed at 0, 4, 8 and 24 h after derivatization. The % accuracy and % CV were calculated.

Freeze and thaw stability. Four standard samples of glucosamine in plasma (0.05, 0.5, 5 and 20 µg/mL) were prepared and kept in the freezer at -20°C for 24 h. Samples were then removed from the freezer and allowed to thaw at room temperature. They were refrozen again for another 24 h. Aliquots of the standard samples were derivatized and analyzed after each freeze-thaw cycle and the % accuracy and CV were determined.

Application
The method was used to detect glucosamine in human plasma separated from blood collected from arms vain of a male adult 0, 1 and 6 h after a single 1500 mg oral dose of glucosamine sulfate (Webber Naturals, Coquitlam BC, Canada, Lot # 567521), and in plasma of 5 rats separated from the jugular veins blood collected through surgically inserted catheters at 0, 0.25, 0.5, 0.45, 1, 2.5, 3, 4, 6 and 8 h after single oral doses of 200 mg/kg glucosamine HCl. We also collected the total urine output of the rats during 0-6 h post-dose.

RESULTS
The validation data generated using the rat and human were identical. Herein, therefore, the human data is presented. As expected (13, 14) both glucosamine and IS appeared as equal size pairs of resolved anomers peaks (Figures 2 and 3). The retention times were 10.6 and 13.1 min for mannosamine and 11.7 and 14.6 min for glucosamine peaks. There was no interfering peak in either spiked or post-dose samples except for an endogenous compound that appeared too close to glucosamine’s first peak. Our attempt to resolve this interference resulted in broad peaks and elongation of the elution time. Hence, since the anomeric peaks of each of the two compounds were identical in size, we used the second peak of each compound for quantification.

The assay was found linear over the examined range of 0.05-20 µg/mL in human plasma with a typical regression equation of y = 0.307x + 0.003 and a correlation coefficient of 0.999. We set the lowest limit of quantification at 50 ng/mL. The inter- and intra-day variations were less than 20% for the 50 ng/mL samples and less than 10% for other concentrations. The accuracy ranged from -1.72 to 2.46% (Table 1).
Figure 2. HPLC chromatogram of blank human plasma (a), blank human plasma spiked with 500 ng/mL glucosamine (b) and plasma obtained from healthy volunteer 1 h after oral administration of 3X500 mg glucosamine sulfate showing 180 ng/mL glucosamine (c). Keys: M, mannosamine; G, glucosamine.

Figure 3. HPLC chromatogram of blank rat plasma (a), blank rat plasma spiked with 500 ng/mL glucosamine (b) and plasma obtained from rat 1 h after oral administration of 200 mg/kg glucosamine (c). Keys: M, mannosamine; G, glucosamine.
Table 1. Precision and Accuracy of glucosamine assay

| Concentration, µg/mL | CV | Accuracy Percent |
|----------------------|----|-----------------|
| Added                | Observed |           |
| **Intra-day**        |           |               |
| 0.05                 | 0.051 ± 0.001 | 2.49 | 1.21 |
| 0.5                  | 0.49 ± 0.02  | 4.87 | -1.71 |
| 5                    | 4.92 ± 0.20  | 4.12 | -1.62 |
| 20                   | 20.49 ± 0.48 | 2.33 | 2.46 |
| **Inter-day**        |           |               |
| 0.05                 | 0.049 ± 0.002 | 4.86 | -1.72 |
| 0.5                  | 0.500 ± 0.008 | 1.63 | -0.03 |
| 5                    | 5.01 ± 0.085 | 1.70 | 0.14 |
| 20                   | 20.30 ± 0.25 | 1.24 | 1.49 |

The percent recovery of glucosamine from plasma in the lowest, medium and highest concentration were determined to be 119±9.1, 100±4.1, 97.2±2.1 and 98.2±1.8% for 0.05, 0.5, 5 and 20 µg/mL, respectively. For mannosamine, the % recovery from plasma was 95.9±2.

Derivatized glucosamine was found stable during analysis in all plasma samples (Table 2). It was also stable in plasma after three freeze and thaw cycles (Table 2).

Table 2. Stability of glucosamine in plasma

| Concentration, µg/mL | CV | Accuracy Percent |
|----------------------|----|-----------------|
| Added                | Observed |           |
| **Short term, 24 h** |           |               |
| 0.05                 | 0.058 ± 0.01 | 11.2 | 12.7 |
| 0.5                  | 0.48 ± 0.01  | 1.41 | 0.5 |
| 5                    | 4.91 ± 0.08  | 1.54 | -1.7 |
| 20                   | 20.7 ± 0.54  | 2.63 | -1.3 |
| **Freeze-thaw stability, 3 cycles freezing at -20ºC** |           |               |
| 0.05                 | 0.051 ± 0.01 | 9.10 | 10.78 |
| 0.5                  | 0.52 ± 0.03  | 6.55 | -3.08 |
| 5                    | 4.90 ± 0.06  | 1.21 | -1.32 |
| 20                   | 20.2 ± 0.05  | 0.25 | 0.28 |

In human plasma, the baseline glucosamine was below the detection limit. A single 1500 mg oral dose of glucosamine yielded plasma concentrations of 184 and 42 ng/mL at 1 and 6 h post-dose (Figure 2). In the rat plasma, glucosamine concentration over the 8 h post-dose period ranged from 7.43 to 0.10 µg/mL that was within the sensitivity of the method (Figure 4).

We did not carry out detailed validation tests for glucosamine in urine. However, the calibration curves generated using spiked rat urine samples were identical to those observed when the compound was tested in water. In the rat urine, after 10 times dilution of the samples, glucosamine concentration ranged from 13-27 µg/ml which account for 1.2 ± 0.42% of the administered dose excreted unchanged in the urine in 6 h post-dose. The anomers were resolved except for the first peak of glucosamine that, as observed in the presence of plasma, eluted along with an interfering peak.

Figure 4. Mean plasma concentration of glucosamine in rat (n=5) after oral administration of 200 mg/kg dose.

DISCUSSION

As a derivatizing agent, FMOC-CL has been used since 1980s for HPLC analysis of amino acids (18, 19). The advantage of this compound is its ability to rapidly react in alkaline media with primary and secondary amines and form a highly fluorescence derivative. In our attempts to conduct a detailed pharmacokinetic study on glucosamine, we used a previously reported method that involved formation of an FMOC-glucosamine derivative (16).

Application of this method gave us a crowded chromatogram in which glucosamine peaks were trapped between two large peaks of the endogenous compounds. Upon repeated injections, the large peaks overlapped with glucosamine. This was accompanied by an elevation in the column pressure. Moreover, the recovery of glucosamine from plasma samples was only 38%. Attempts to
improve the recovery by increasing the amount of the derivatizing agent resulted in more interfering peaks. In the reported methods for determination of amino acids after derivatization with FMOC-CL, the aqueous part of the mobile phase has generally been an acetate buffer rendering the acidic analytes (amino acids) less polar, hence, allowing longer retentions on the reversed phase column. Our replacement of the acetate buffer with 0.1% acetic acid in HPLC water resulted in cleaner and more reproducible chromatograms (Figures 2 and 3). In the meantime, we decreased the percentage of acetonitrile in the first 13 min from 30% to 23% to allow maximum separation of the peaks.

Huang et al reported that FMOC-CL precipitated when acetonitrile concentration dropped below 30%. To avoid precipitation we injected aliquots of only 5 µL of samples after dilution with 1:1 acetonitrile-water.

Under our conditions, most of the plasma interfering peaks eluted after the appearance of mannosamine HCl that we used as the internal standard. Mannosamine has a chemical structure and physical properties similar to glucosamine except in the amino group at the position 2. This group is equatorial in glucosamine and axial in mannose, hence, provides a different three dimensional orientation and facilitate efficient HPLC separation of the two compounds (20). Reaction with FMOC-CL adds a large molecule at position 2 and obtains a steric hindrance that prevents the possibility of inter-conversion between glucosamine and mannose.

For the amino acid analysis, the excess derivatizing agent has been generally removed by extraction with pentane or by reaction with ADAM to form a hydrophobic complex that elutes towards the end of the run time (21). Since glucosamine is a weak base and FMOC-CL and FMOC-glucosamine peaks are eluted at different times, Zhang et al (16) and Huang et al (17) did not need to apply the sample cleaning step. However, under our experimental condition, we noticed a small peak that interfered with that of glucosamine. This interfering peak was eliminated when we added ADAM. In addition, the sample cleaning rendered the baseline more stable even after doubling the amount of FMOC-CL (from 4 mM to 8 mM) that enable us to achieve a lowest limit of quantification at 50 ng/mL and a percent recovery of 98% and 96% for glucosamine and mannose, respectively.

The method is both accurate and reproducible. It also is more sensitive than some of the reported methods [e.g., (9)] and provides more facile procedure than others (10-12, 14, 17). The application of the assay in pharmacokinetic studies of glucosamine in the rat allowed analysis of plasma concentrations up to 8 h post-dose. In our previous report we had analyzed pharmacokinetics of glucosamine only up to 4 h due to the lack of sensitivity (22). The previously reported methods including ours (9) did not detect baseline glucosamine concentrations in the rat due to the inherit lack of sensitivity. In the present study, on the other hand, we did observed baseline glucosamine levels but were generally low (see the 8 h value in Figure 4) but as high as 250 ng/mL in a single rat. Such an example is presented in Figure 3. This may render the reliability of the post 4 h concentration questionable. We did not correct for the baseline values as we were not certain of the consistency of the endogenous baseline levels during the 8 h experiment. This is inevitable with sensitive assays developed for the determination of endogenous compounds. For glucosamine, the post 4 h concentrations are rather low, hence, their contribution to the overall area under the curve should be negligible; however, they may introduce error in calculation of the compound’s terminal half-life. The assay is also suitable for the measurement of glucosamine in human plasma. We detected no baseline glucosamine in the pre-dose human sample. Jackson et al (23) and Zhong et al (11) also did not detect baseline glucosamine concentrations in their subjects. Roda et al (12), on the other hand, have reported a mean of 64±47 ng/mL endogenous glucosamine. It is worthy of mentioning that Roda et al’s reported overall glucosamine concentrations (12) are substantially greater that those reposted by Jackson et al (23), and Zhong et al (11). Nevertheless, similar to what we have observed in the rat (Figure 3), the possibility of endogenous glucosamine must also be considered for interpretation of human plasma samples containing low concentrations of the compound.

We did not completely validate glucosamine assay in urine. Nevertheless, this is the first reported HPLC method for determination of unchanged glucosamine in urine samples. The total
glucosamine excreted in human urine in 3 h has previously been determined by ion exchange chromatography with amperometric detection to be less than 1% of the administered dose (24). Our results suggest 1.2% cumulative urinary excretion in the rat over 6 h following oral doses to rats. In this context, it is important to consider the relative bioavailability of oral glucosamine doses (9).

Glucosamine concentration in human plasma appears to be dependent on the administered product. The reported values following 1500 mg glucosamine as hydrochloride (23) or sulfate (25) are approximately three-fold different from each other. We found 184 and 42 ng/mL of glucosamine in plasma of a man following a single 1500 mg oral dose of the sulfate salt. The 6 h sample was below the set sensitivity of the assay, i.e., 50 ng/mL. However, we injected only 5 µL of the final solution to prevent pump pressure build-up. An increased volume of injection of only those samples with lower concentration, however, is not expected to cause pressure build-up but will improve the assay sensitivity. Our observed concentrations in human plasma samples are in agreement with those reported by Jackson et al (23). Nevertheless, the method offers sensitivity within the reported concentration ranges.

CONCLUSION

Herein we present an improved assay of glucosamine in biological fluids suitable for pharmacokinetic studies. The assay has a reliable sensitivity for concentrations above 50 ng/mL based on 5 µL of injection volume onto HPLC. The assay sensitivity, however, can be improved using larger injection volumes.

ACKNOWLEDGEMENT

AI is a recipient of an Egyptian government scholarship. This work was supported by a grant from the Canadian Foundation for Innovation.

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