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LC-MS/MS method for the determination of carbamathione in human plasma

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ABSTRACT

Liquid chromatography-tandem mass spectrometry methodology is described for the determination of S-(*N*,*N*-diethylcarbamoyl)glutathione (carbamathione) in human plasma samples. Sample preparation consisted of a straightforward perchloric acid medicated protein precipitation, with the resulting supernatant containing the carbamathione (recovery ~98%). For optimized chromatography/mass spec detection a carbamathione analog, S-(*N*,*N*-di-*i*-propylcarbamoyl)glutathione, was synthesized and used as the internal standard. Carbamathione was found to be stable over the pH 1–8 region over the timeframe necessary for the various operations of the analytical method. Separation was accomplished via reversed-phase gradient elution chromatography with analyte elution and re-equilibration accomplished within 8 min. Calibration was established and validated over the concentration range of 0.5–50 nM, which is adequate to support clinical investigations. Intra- and inter-day accuracy and precision determined and found to be <4% and <10%, respectively. The methodology was utilized to demonstrate the carbamathione plasma-time profile of a human volunteer dosed with disulfiram (250 mg/d). Interestingly, an unknown but apparently related metabolite was observed with each human plasma sample analyzed.

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1. Introduction

Disulfiram has been used for the treatment of alcohol dependence since its accidental discovery approximately 60 years ago [1]. The pharmacological basis and rationale for the use of disulfiram as a deterrent treatment for alcohol dependence is its inhibition of liver low *K_m* mitochondrial aldehyde dehydrogenase (ALDH₂). The inhibition of ALDH₂ results in an increase in acetaldehyde which is believed to be responsible for the adverse effects known as the disulfiram-ethanol reaction. Disulfiram is a pro-drug that requires bioactivation to an active metabolite which is responsible for the inhibition of ALDH₂ [2–4]. The metabolite responsible for the inhibition of ALDH₂ has been identified as DETC-MeSO [5] and the cytochrome P450 enzymes required in this bioactivation process has been delineated [6,7]. DETC-MeSO is oxidized to S-methyl N,N-diethylthiocarbamate sulfone (DETC-MeSO₂) [8] and subsequently to S-(*N*,*N*-diethylcarbamoyl)glutathione(carbamathione) [9] (Fig. 1).

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Carbamathione is a partial NMDA glutamate antagonist [9] but does not inhibit liver ALDH₂ [10]. The clinical significance of carbamathione and its possible neurochemical activity [9] was not considered until studies showed that disulfiram was a promising therapeutic agent for treating cocaine dependence [11–15]. Although the mechanism of action of disulfiram's effectiveness in cocaine dependence is unclear, a working hypothesis has emerged that carbamathione may be the responsible metabolite because of its NMDA glutamate receptor activity [9]. Early metabolite distribution studies with radio labeled ³⁵S [16,17] suggested a possible disulfiram metabolite in rat brain would no longer carry a disulfiram derived sulfur but rather a metabolite with only a N.Ndiethylcarbamoyl moiety. In rat studies carbamathione has since been found in rat urine, bile [18], plasma and brain [19]. While carbamathione has been detected in a variety of rat fluids, the suggestion that carbamathione may have a neurological action in humans must be supported by detection in humans.

In a previous report [19], methodology was described for the determination of carbamathione present in rat brain and plasma where sampling and sample preparation was accomplished by microdialysis. The samples further subjected to reversed-phase gradient elution liquid chromatography (LC) separation followed by detection and quantification by tandem mass spectrometry (MS/MS). While allowing for determination at the 1 nM level, the

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Fig. 1. Metabolites of disulfiram referred to in this work.

combination of internal standard (IS) retention and chromatographic re-equilibration resulted in a sample throughput time of 50 min.

The goal of the present investigation was the development of methodology for the determination of carbamathione in human plasma samples suitable for support of human clinical investigations with improved sample throughput. Issues to be addressed include plasma sample preparation and the identification of an internal standard that is optimized with respect to the LC retention window and provides for sensitive and selective MS detection. While plasma sample preparation proved to be straightforward using acid mediated protein precipitation, the identification of an internal standard meeting the previously noted criteria involved the synthesis and evaluation of several glutathione analogs in order to meet the desired criteria. Ultimately the developed methodology described herein was applied to human plasma samples obtained from disulfiram-dosed healthy volunteers, and to our knowledge, appears to be the first report of the presence of carbamathione as a disulfiram metabolite in humans.

2. Experimental

2.1. Chemicals and reagents

Carbamathione was synthesized using methods previously developed [18] and obtained from Kaul et al. [19]. S-*n*-butylglutathione, tris hydrochloride, ammonium bicarbonate, di-*i*-propylcarbamoylchloride, and propyl-, butyl- and pentyliso-cyanate were purchased from Sigma-Aldrich (St. Louis, MO, USA). Ammonium acetate, sodium phosphate, formic acid, perchloric acid (70%) and HPLC grade methanol were obtained from Fisher (Fairlawn, NJ, USA). Nanopure water was prepared by a Water Pro Plus purification system (Labconco, Kansas City, MO, USA). Blank pooled human plasma was purchased from Innovative Research (Novi, MI, USA).

2.2. Synthesis of internal standard candidates

2.2.1. General procedure for S(N-alkylcarbamoyl)glutathione derivatives

Carbamathione analogs were synthesized by reacting propyl-, butyl- or pentyl isocyanate with glutathione according to the method of Han et al. [20]. The structures of the products obtained were confirmed by 1 H NMR and mass spectrometry.

2.2.1.1. *S*-(*N*-*n*-propylcarbamoyl)glutathione. Exact mass determination of $[M+H]^+$ C₁₄H₂₅N₄O₇S was 393.1451±0.0015 (*n*=3) which deviated by 1.8 ppm from the expected mass. The proton NMR chemical shifts were δ 0.82 t 3H (CH₃CH₂CH₂NH–), δ 1.41 h 2H (CH₃CH₂*CH₂NH–), δ 1.91 q 2H (Glu- β , β'), δ 2.3 m 2H (Glu- γ , γ'), δ 3.05 q 2H (CH₃CH₂CH₂*NH–), δ 3.32 m 2H (Cys- β , Glu- α), δ 3.68 d 2H (Gly- α , α'), δ 4.34 t, 1H (Cys- α) (*indicates specific protons).

2.2.1.2. *S*-(*N*-*n*-butylcarbamoyl)glutathione. Exact mass determination of $[M+H]^+$ C₁₅H₂₇N₄O₇S was 407.1590±0.0008 (*n*=3) which deviated by -2.5 ppm from the expected mass. The proton NMR chemical shifts were δ 0.85 t 3H (CH₃CH₂CH₂CH₂NH–), δ 1.25 h 2H (CH₃CH₂*CH₂CH₂NH–), δ 1.38 q 2H (CH₃CH₂CH₂*CH₂NH–) δ 1.91 m 2H (Glu- β , β'), δ 2.29 q 2H (Glu- γ , γ'), δ 3.08 m 2H (CH₃CH₂CH₂CH₂*NH–), δ 3.32 m 2H (Cys- β , Glu- α), δ 3.68 dd 2H (Gly- α , α'), δ 4.33 dt, 1H (Cys- α) (*indicates specific protons).

2.2.1.4. S-(N,N-di-i-propylcarbamoyl)glutathione. A solution of diisopropylcarbamoylchloride (1.2 mmol in 7 ml pyridine) was added to a solution of glutathione (0.6 mmol in 3 ml water) and stirred over night at room temperature. The resulting mixture was evaporated to dryness and re-dissolved in 6 ml water. Care must be taken that the reaction mixture is not heated excessively (30 °C max) during reaction or solvent evaporation as excessive heating results in an oily orange byproduct, which is difficult to remove. The crude product was subjected to C18 semi-preparatory HPLC (70% H₂O, 30% methanol and 0.1% formic acid) and dried in a lyophilizer, resulting in a white powder. The structure was confirmed by ¹H

NMR and mass spectrometry. Exact mass determination of $[M+H]^+$ $C_{17}H_{31}N_4O_7S$ was 435.1901 \pm 0.0006 (n = 3) which was -2.8 ppm from the expected mass. The proton NMR chemical shifts were δ 1.21 d 12H ((CH₃)₂CH)₂N-), δ 1.91 q 2H (Glu- β , β'), δ 2.29 m 2H (Glu- γ , γ'), δ 3.33 m 5H (((CH₃)₂CH^{*})₂N-, Cys- β , Glu- α), δ 3.69 d 2H (Gly- α , α'), δ 4.35 t, 1H (Cys- α) (*indicates specific protons).

2.3. Methods

2.3.1. LC method for MS detection

Solvents were delivered by a Waters Acquity UPLC system. Chromatographic separation was performed using a Phenomenex Kinetex column (phase C18; diameter, $2.1 \text{ mm} \times 50 \text{ mm}$; particles size, 1.7 µm; pore size, 100 Å), which was protected by a Phenomenex KrudKatcher Ultra filter (0.5 µm pores, 316 stainless steel) followed by a Waters Vanguard C8 pre-column. Chromatographic solvents consisted of (A) 99% H₂O, 1% methanol and 0.1% formic acid and (B) 1% H₂O, 99% methanol and 0.1% formic acid delivered at a flow rate of 400 µl/min. The sample was injected (50 µl volume) onto a column that was pre-equilibrated at 5% B. After 0.2 min a linear gradient of 11% B/min for 4 min was implemented. The column was then washed with 80% B for 2 min and subsequently re-equilibrated at 5% B for 1.5 min, resulting in a total run-time of 8 min. For the first two min, chromatographic effluent is diverted to waste, preventing salts and polar endogenous compounds from entering the mass spectrometer.

2.3.2. LC method for UV detection

The HPLC system consisted of two Shimadzu LC6A pumps and a Shimadzu SIL-6B auto injector (50 μ l injection utilized) controlled by a Shimadzu SCL-6B system controller. Detection was accomplished using a Kratos UV/Vis chromatographic detector set at 215 nm with TurboChrom V4 used for data collection. Chromatographic separation was conducted using a Phenomenex Inertsil ODS-3 column (150 mm × 4.6 mm, 5 μ particles, 100 Å) protected with a Supelcosil LC-8 guard column (2 mm × 2.1 mm, 5 μ m particles). An isocratic solvent program consisting of 70% A (95% H₂O, 5% methanol and 0.1% formic acid) and 30% B (5% H₂O, 95% methanol and 0.1% formic acid) with a flow rate of 1 ml/min, was utilized for the chromatography.

2.3.3. Mass spectrometry parameters

Mass spectrometry was performed on a Quattro Ultima "triple" quadrupole instrument (Micromass Ltd., Manchester, UK). The mass spectrometer was run in positive ion mode using an ESI ionization source. The mass spectrometer source block was set at 100 °C and the desolvation gas temperature was set at 300 °C. Argon collision gas was set to attenuate the beam by 10-20% (10^{-3} mbar) . Quadrupoles 1 and 3 were tuned to a resolution of 0.9 amu FWHH. Collision energy and cone voltage settings are optimized for each compound. Data processing was performed using MassLynx 4.1 and Graphpad Prism 5. Fragmentation patterns of S-n-butyl-glutathione, S-(N-n-propylcarbamoyl)glutathione, S-(*N*-*n*-butylcarbamoyl)-glutathione, S-(*N*-*n*-pentylcarbamoyl)-S-(*N*,*N*-di-*i*-propylcarbamoyl)glutathione glutathione, and carbamathione and were determined by infusing an aqueous solution of the respective compound at 400 µl/min 30% solvent B (HPLC-MS method) for source optimization. Product ion scans were acquired at varying collision energies, 10-45 V. One MRM transition unique to the substance was chosen for each compound and the collision energies optimized. The most abundant MRM transitions were chosen for the secondary carbamoyl substances and S-*n*-alkyl conjugates. However, for the tertiary carbamoyl substances a carbamoyl moiety related fragment was selected as most suitable reporter transition (Fig. 2). The mass transitions and optimum settings for all compounds are summarized in Table 1.

Subsequently a mixture of all five internal standard candidates (100 nM solution of each) was run on the HPLC-MS system to determine relative retention times with respect to carbamathione (Fig. 3).

2.3.4. Sample pretreatment

Plasma, 100 μ l in an Eppendorf tube, was spiked with 10 μ l of 100 nM S-(*N*,*N*-di-*i*-propylcarbamoyl)glutathione (internal standard) and for calibration curves, 10 μ l of a carbamathione solution. Water, 45 μ l, was then added and the mixture vortexed for 5 s. In the case of blank or patient samples, 55 μ l water was added instead of 45 μ l to compensate for the volume. Proteins were precipitated by the addition of 35 μ l perchloric acid and the mixture vortexed for 5 s and then placed on ice for 5 min. The precipitated sample was subsequently centrifuged for 5 min at 11,000 times gravity (13,000 rpm at 60 mm radius). The supernatant was recovered and transferred to an autosampler vial for subsequent determination.

2.3.5. Stability

Carbamathione plasma stability was tested in pooled human plasma. Samples (n = 5) enriched with 150 μ M carbamathione were allowed to incubate at room temperature for 2, 4, 6 and 24 h prior to acid mediated precipitation. Analyte stability in the plasma extract was assayed in the fashion, but substituting plasma for the acid plasma extract. A limited freeze-thaw stability experiment was performed by analyzing carbamathione enriched plasma samples (n = 5) at day 1, 3 and 7 of a seven day period. Carbamathione stability was checked against a five point calibration curve ranging from 50 to 250 μ M carbamathione in plasma.

2.4. Validation

The sample pretreatment method was applied (n=5) to five spiked plasma concentrations of 50.0, 10.0, 5.0, 1.0, and 0.5 nM and the samples analyzed according to the described methodology. This experiment was repeated for 4 consecutive days to determine intra and inter-day stability of the method. Data was processed for linearity, precision, accuracy and to determine the limit of quantification. An infusion of carbamathione was utilized to obtain a qualitative assessment of ionization suppression caused by the human plasma matrix. An infusion pump was used to deliver a constant amount of carbamathione post-column into the LC stream entering the mass spectrometer. The mass spectrometer was set in SRM mode to monitor the analyte signal stability and a blank plasma sample was injected onto the column. Carbamathione stability was assessed at various pH's and at elevated temperature. Solutions of carbamathione (100 μ M) were prepared in five 100 mM buffer solutions of perchloric acid (pH 1), formic acid (pH 2.5), acetate (pH 4), tris (pH 6) and phosphate (pH 8). Sample vials were kept at 50 °C and samples taken every 24 h for 5 consecutive days to determine stability employing the HPLC-UV method. Samples were checked against a control sample that was stored at -20 °C and thawed immediately prior to analysis.

2.5. Human studies

A healthy 52-year-old female Caucasian volunteer with no alcohol or other substance use disorders was recruited. Voluntary, written, informed consent was obtained, and approval obtained from the University of California at San Francisco Committee on Human Research. The patient volunteer weighed 69 kg and was 160 cm in height. Disulfiram 250 mg/d was administered with staff observation for 3 days. On the fourth day, after an 8 h overnight fast, an antecubital venipuncture was undertaken and 7 ml of blood was drawn into a heparinized tube to obtain trough disulfiram concentration in the plasma. A urine sample was also collected prior to



Fig. 2. Potential internal standards and their CID fragmentation: **A**, S-*n*-butylglutathione; **B**, S-(*N*-*n*-propylcarbamoyl)glutathione; **C**, S-(*N*-*n*-butylcarbamoyl)glutathione; **D**, S-(*N*-*n*-pentylcarbamoyl)glutathione; **E**, S-(*N*.*N*-di-*n*-propylcarbamoyl)glutathione; **F**, carbamathione. Structures are annotated with the mass in the spectrum, backbone fragments have an additional proton to that suggested by the bars.

disulfiram dosing to test for recent use of any illicit substances. A 250 mg dose of disulfiram was then administered and blood samples drawn at 1, 2, 4, 6, 8, and 10 h after disulfiram dosing. Each blood sample was immediately centrifuged and the plasma separated and frozen at -70 °C until the time of determination of carbamathione.

3. Results and discussion

3.1. Chromatography

As noted previously, human plasma assay methodology for carbamathione must possess the necessary sensitivity and selectivity to support continued clinical investigation, but from an analytical perspective, should also be robust and enable high sample throughput. In order to achieve high efficiency rapid separations one frequently resorts to the use of high linear velocities (>2 mm/s), small particles (<3 μ m) or a combination of these parameters, which can result in the need for much higher operating pressures as compared to those available to many common LC systems currently in use. An alternative approach is the use of solid core particles featuring a porous shell that results in efficient kinetic performance without the need for substantially higher operating pressures. Based on these considerations chromatographic method development was based on the use of a reverse-phase Phenomenex Kinetex column (1.7 μ m porous shell particles), which reduces

Table	1
Parent	t

Parent ion and SRM fragm	ent for quantitation ar	nd their optimum con	e voltage and collision	i energy settings.
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	Compound	Parent ion	Fragment	Cone voltage	Collision energy (V)
А	S-butylglutathione	364.0	218.0	35.0	15.0
В	S-(N-propylcarbamoyl)glutathione	393.0	264.0	35.0	14.0
С	S-(N-butylcarbamoyl)glutathione	407.0	278.0	35.0	14.0
D	S-(N-pentylcarbamoyl)glutathione	421.0	292.0	35.0	14.0
E	Diisopropylcarbamathione	435.0	128.0	35.0	30.0
F	Carbamathione	407.1	100.0	35.0	30.0

diffusion distance in the retention process and thus mimics the chromatographic performance of smaller particles (e.g., 1 μ m particles) without the associated increase in backpressure needed to achieve similar linear velocities with the smaller diameter particles.

Other aspects addressed include the development of an efficient chromatographic gradient elution program. For high sensitivity bioanalysis, one is often interested in using a relatively large injection volume with respect to the dimensions of the chromatographic column in order to minimize dilution. To this end, when possible it is desirable to use an initially weak mobile phase composition with respect to the analyte and/or injection solvent composition to allow for larger volume injections, thus providing for the achievement of trace enrichment, which translates into the potential to achieve high sensitivity detection.

With these criteria in mind, the gradient profile implemented consisted of an analyte focusing step (initial mobile phase composition was \sim 5% MeOH), a relatively steep gradient elution program (11%/min linear ramp of solvent B, solvent B \sim 99% MeOH) and a washing step (80% solvent B), followed by return to the initial conditions with only 1.5 min required to achieve column re-equilibrium. The total injection to re-equilibration time was achieved in 8 min, representing a substantial throughput enhancement as compared to the previously reported methodology [19]. The adopted solvent program (details presented in the experimental section) allowed the loading of a relatively large volume injection (50 µl applied to a 2.1 mm i.d. column), and when followed by the previously noted steep gradient program, resulted in excellent sensitivity (details provided in a latter section) without the need for pre-concentration during sample preparation. A typical chromatogram of the analyte present in aqueous solution is provided in Fig. 3A, where the elution time is noted as \sim 2.7 min.



Fig. 3. (A) HPLC/SRM chromatograms of carbamathione and (B) five potential internal standards: **A**, S-*n*-butylglutathione; **B**, S-(*N*-*n*-propylcarbamoyl)glutathione; **C**, S-(*N*-*n*-butylcarbamoyl)glutathione; **D**, S-(*N*-*n*-pentylcarbamoyl)glutathione, **E**, S-(*N*,*n*-di-i-propylcarbamoyl)glutathione. SRM transitions are in Table 1, 100 nM solutions of each substance were injected.

3.2. Internal standard selection

The incorporation of an internal standard (IS) generally adds to the quantitative robustness of bioanalytical methods. With regard to physical-chemical properties, an ideal situation occurs when the IS exhibits similar solubility, chemical reactivity, ionization constants, possesses a closely matched chemical structure as compared to the analyte of interest. The closest match occurs with the use of a stable isotope modification of the analyte, which is especially useful for LC-MS/MS based methodology. In the present case, stable isotope labeled carbamathione is not readily available, nor is any (isotope labeled) precursors or building blocks that would allow for a straight-forward synthesis of this molecule. Accordingly, efforts were directed to obtaining five IS candidates offering appropriate chromatographic and MS fragmentation characteristics.

As with carbamathione, all of these substances can be viewed as straightforward chemical elaborations of glutathione, and can be categorized into three different categories based on their functional groups present. They are seen to be an S-*n*-alkyl-glutathione derivative (Fig. 2A) that deviates the most from carbamathione, having an alkyl moiety bonded to the cysteine sulfur and thus lacking the carbamoyl linkage, three S-(*N*-mono-*n*-alkylcarbamoyl)glutathiones (Fig. 2B–D) each possessing the carbamoyl linkage of carbamathione, but with only a single alkyl group present on the nitrogen and an S-(*N*,*N*-di-*n*-alkylcarbamoyl)glutathione (Fig. 2E), that possesses a tertiary carbamoyl moiety (two alkyl substituents on the carbamoyl nitrogen) that closely mimics the structure of carbamathione.

In order to determine the optimal internal standard for the quantitation of carbamathione, the properties of the five IS candidates were investigated with respect to their LC retention times and fragmentation during collision induced dissociation (CID). As expected, the various potential internal standards showed significant differences in retention time (Fig. 3B) and bracketed the elution window of carbamathione (Fig. 3A). As regards MS/MS detection criteria, the CID fragmentation patterns of each substance are shown in Fig. 3. From these results, the most sensitive mass transition for each IS candidate was selected for evaluation against that of carbamathione (Table 1). Based only on retention time, S-(N-n-butylcarbamoyl)glutathione (Fig. 2C) would seem to be the preferred internal standard as it co-elutes with carbamathione (Fig. 3). However, this substance is isobaric with carbamathione and as a result this internal standard candidate would only be suitable if it offered a unique CID fragment (compare Fig. 2C-F). Several unique transitions are present, however, crosstalk between carbamathione and S-(N-n-butylcarbamoyl)glutathione SRM channels could not be avoided (note the low intensity $364 \rightarrow 278$ transition, which is the major carbamathione transition) regardless of which transition was selected. The difficulty of avoiding the cross talk, presumably with low abundance glutathione backbone fragments, required a reevaluation of internal standard selection criteria. Note that the CID spectra of S-alkyl- and S-(N-alkylcarbamoyl) glutathiones are dominated by backbone fragments (Fig. 2A-D) and have common low mass fragments below m/z 200.

Moving to other candidates exhibiting a different retention times as compared to carbamathione, alleviates LC co-elution and

Table 2 Stability of carbamathione at room temperature (21 °C).

^a Expected concentration for full recovery is $150 \,\mu$ M.

provides unique fragmentation products amenable to quantitation. Comparing the S-(*N*-mono-n-alkyl) secondary carbamoyl derivatives (Fig. 2B–D) to the tertiary carbamoyl group of carbamathione (Fig. 2F) suggests the tertiary carbamoyl group stabilizes a side chain only fragment. This motivated the synthesis of S-(*N*,*N*-di*i*-propylcarbamoyl)glutathione (Fig. 2E). As expected, the **E** CID spectrum has abundant carbamoyl related fragments and at optimum CE the m/z 128 fragment dominates the CID spectra, which is similar to the dominant carbamoyl related m/z 100 fragment of carbamathione. Based on these results, substance **E** (Fig. 2) was selected as the IS for the plasma assay of carbamathione.

3.3. Sample preparation

The polar analyte glutathione can be extracted efficiently out of human plasma by various acid mediated protein precipitation procedures, followed by centrifugation [21,22]. The resulting aqueous solution is compatible with a reversed phase HPLC separation, resulting in a convenient sample preparation strategy. Since the targeted analyte and IS are each simple chemical elaborations of glutathione, acid mediated protein precipitation was evaluated as an approach to sample deproteinization. Accordingly, aqueous and plasma samples containing carbamathione were prepared and subjected to perchloric acid (PCA) mediated protein precipitation (n=5). After recovery of the resulting supernatant, analyte determination revealed recovery of $98.1 \pm 2.0\%$ for plasma with respect to the aqueous standards. Due to this result other acids were not investigated However, to determine if unknown plasma components present in the deproteinized plasma supernatant resulted in an altered MS ionization process, a post-column infusion of carbamathione and the IS was conducted. The results obtained revealed that in the elution window of interest there was no endogenous compound interference or ion suppression. An observation that was further reinforced by the quantitative recovery of carbamathione.

With this information in place, there was still the need to evaluate the stability of the analyte in the media of various pH values expected to be encountered during the envisioned assay operations. During assay development of carbamathione present in rat brain and plasma microdialysis samples, Kaul et al. [19] found the substance to be stable in Ringer's solution at room temperature, frozen and when subjected to several freeze-thaw cycles over the analytical timeframe. However in the present work, there was need to establish solution stability over a wider pH range and in an attempt to force degradation, solutions of carbamathione were prepared at pH 1.0–8.0 and heated to 50 °C. These solutions were evaluated every 24 h for a week with no observed degradation, thus establishing that carbamathione is stable at 50 °C for a week over the pH 1.0–8.0 range, clearly sufficient for the various analytical operations needed during plasma sample determination.

The stability of carbamathione within a plasma matrix was also evaluated (Table 2). Carbamathione does demonstrate some plasma instability at room temperature, with 7.8% degradation observed after 2 h of incubation. The post-extraction sample was determined to be stable for at least 24 h, which is a sufficient timeframe for LC-MS analysis (Table 2). Since carbamathione is stable

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Freeze-thaw stability of carbamathione.

	Carbamathione	Carbamathione concentration (μ M)			
	Nominal	Measured	RSD (%)		
Day 1	100	107.2	5.9		
	150	158.7	4.4		
	200	202.0	1.7		
Day 3	100	119.5	6.7		
	150	154.4	2.1		
	200	189.1	4.4		
Day 7	100	110.8	3.2		
	150	139.4	8.3		
	200	192.7	12.2		

in both dialysates and plasma extracts, the plasma instability of carbamathione is indicative of an enzymatic degradation pathway. Therefore it is important that plasma samples are processed within 10 min of thawing. Since sample processing consists out of a simple protein precipitation step, this is not problematic. The freeze-thaw stability of carbamathione was analyzed by subjecting enriched samples to three freeze-thaw cycles over seven days (Table 3). No significant degradation was observed over this time frame.

3.4. Method linearity, accuracy, and precision

The linearity, accuracy and precision of the method were determined by the analysis of calibration plots ranging from 0.5 nM to 50.0 nM spiked plasma concentration (n = 5 per concentration) for four consecutive days. Intra- and inter-run accuracy and precision were typically within 10% of the target value. A typical calibration curve was described by the following linear equation: y = 1.978x - 0.039 (y = dimension less number obtained by normalization of the carbamathione response to the internal standard; x = concentration of carbamathione in nM). The lower limit of quantification (LLOQ) with acceptable accuracy and precision (<10%) was determined to be at 0.5 nM in plasma. While FDA guidelines state that LLOQ precisions of <20% are acceptable, the LLOQ was not challenged further as 0.5 nM was determined to be adequate for the current investigation. The limit of detection was not challenged below 0.1 nM in plasma, while the 0.1 nM plasma sample still revealed S/N > 5. The method exhibits good intra- and inter-run linearity with $R^2 = 0.9972$ and $R^2 = 0.9976$, respectively. Accuracy values were within 5% of expectation and sample precision was within 10% over the validated range (Table 4).

3.5. Human plasma levels

The developed carbamathione plasma assay methodology was successfully applied to determination of samples obtained from a disulfiram-dosed volunteer. Fig. 4 depicts a typical chromatogram of the results obtained, in this case that of a sample obtained 1 h after dosing where a chromatographic peak is observed that represents carbamathione present in a concentration of 2.1 nM. Interestingly, the various samples show an unidentified peak (Fig. 4), which was detected in every sample containing carba-

Table 4

Summary of validation data showing intra and inter-run accuracy and precision.

Carbamathione concentration (nM)	e Intra-run (n = 5) (nM)			Inter-run (<i>n</i> =20)		
	Mean observed concentration (nM)	Precision (RSD %)	Mean accuracy of target value (%)	Mean observed concentration (nM)	Precision (RSD %)	Mean accuracy of target value (%)
0.5	0.5	7.9	100.6	0.5	9.0	103.8
1.0	1.0	4.9	99.3	1.0	4.9	99.8
5.0	5.2	3.2	103.0	4.9	5.4	98.8
10.0	9.9	3.9	99.3	9.9	6.0	98.6
50.0	49.9	4.8	99.8	50.2	4.0	100.4



Fig. 4. HPLC SRM of a plasma sample obtained from a female volunteer 1 h after oral administration of disulfiram, 250 mg (see Section 2.5).



Fig. 5. Plasma profile obtained from a female volunteer after oral administration of 250 mg of disulfiram (see Section 2.5).

mathione. This peak results from the MRM transition used to detect carbamathione and also matches the carbamathione profile in plasma. It is highly probable that this unknown substance results from a disulfiram metabolite possessing the carbamoyl (100 m/z) fragment. The plasma concentration profile obtained as a function of time (1, 2, 4, 6, 8, and 10 h) subsequent to oral dosing of a healthy volunteer is shown in Fig. 5.

4. Conclusion

A selective and sensitive method for the detection and quantification of carbamathione in human plasma has been developed. The method exhibits accuracy and precision of 5% and \leq 10%, respectively, and achieves good chromatographic throughput (8 min injection to injection with a methanol:water gradient). Detection was accomplished by tandem mass spectrometry in positive ion mode, using SRM and a structural homolog, S-(*N*,*N*-di-*i*propylcarbamoyl)glutathione, Fig. 2E, as the internal standard. In human plasma samples, the limit of quantification and detection were 0.5 nM and 0.1 nM, respectively. The rapid chromatographic cycle and simple sample pretreatment allow for a high sample throughput, which is very desirable for bioanalytical methodology required to support any clinical investigation. Furthermore, the method was successfully applied to biological samples and used to determine the plasma profile of carbamathione in a single human volunteer. The analysis of disulfiram-dosed volunteer plasma samples resulted in the discovery of a compound peak having a similar plasma-time profile as carbamathione. The observed compound contains a 100 m/z fragment, which is highly suggestive that it is related to the carbamoyl fragment observed for carbamathione.

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