Glutathione transferases (GSTs) are enzymes devoted to the protection of cells against many different toxins. In erythrocytes, the isoenzyme (e-GST) mainly present is GSTP1-1, which is overexpressed in humans in case of increased blood toxicity, as it occurs in nephroptic patients or in healthy subjects living in polluted areas. The present study explores the possibility that e-GST may be used as an innovative and highly sensitive biomarker of blood toxicity also for other mammals. All distinct e-GSTs from humans, Bos taurus (cow), Sus scrofa (pig), Capra hircus (goat), Equus caballus (horse), Equus asinus (donkey) and Ovis aries (sheep), show very similar amino acid sequences, identical kinetics and stability properties. Reference values for e-GST in all these mammals reared in controlled farms span from 3.5 ± 0.2 U/gHb in the pig to 17.0 ± 0.9 U/gHb in goat; such activity levels can easily be determined with high precision using only a few microliters of whole blood and a simple spectrophotometric assay. Possibly disturbing factors have been examined to avoid artifact determinations. This study provides the basis for future screening studies to verify if animals have been exposed to toxicologic insults. Preliminary data on cows reared in polluted areas show increased expression of e-GST, which parallels the results found for humans.

RESULTS

Molecular properties of e-GSTs from different mammalian species

The amino acid sequences of e-GSTs from swine, goat, cow, sheep and two equine species display extraordinary similarity among them, and also with the human enzyme. Most of these GSTs show >85% of sequence identity with the human isofrom (Supplementary Figure S1 and Supplementary Table S1). Of interest is the strict conservation of the four cysteines (Cys14, Cys47, Cys101 and Cys169) that confer peculiar redox sensitivity to this enzyme. Both in human and horse e-GSTs, many oxidizing chemicals may induce the formation of an intra-chain disulfide involving the two highly reactive cysteines, that is, Cys47 and Cys101.10,11 These oxidized forms are completely inactive, but they can be reactivated under reducing treatment with DTT at alkaline pH values. The presence of a detectable inactive oxidized form of GSTP1-1 has been recently found in salivary samples of healthy human subjects.11 The possible presence of such oxidized forms in the blood of different mammals will be explored below.

Kinetics and stability properties of e-GSTs from different mammalian species

The similarity of the primary structures of these distinct e-GSTs suggests almost identical kinetics properties. As expected, $K_m$ values for GSH and CDNB are very similar (Table 1) and $k_{cat}$ values also range between 75–85/s for all mammalian species selected.
for this study (Table 1). Thus, all these mammalian isoforms of e-GST appear kinetically very similar.

Essential for a possible use of these enzymes in screening analysis is their stability upon storage. In intact erythrocytes, almost all these transfersases display very high stability for many days when stored at 4°C (no statistically significant decrease in activity after 7 days). The exception is the pig e-GST that is stable only for 2 days, displaying a loss of 25% of its original activity after 7 days (Supplementary Table S2). Storage of blood samples at −20°C must be avoided as it causes partial inactivation of e-GSTs from all species (data not shown).

Activity of e-GST in bovine erythrocytes

e-GST activity has been measured in 40 pregnant cattle reared in controlled farms. Pregnant cattle display 11.2 ± 0.7 U/g Hb, a value higher than that found in humans (5.8 ± 0.4 U/g Hb) (Figures 1a and b). For these animals, we tested if the e-GST level could be influenced by different physiological conditions, that is, pregnancy, and during lactating phase, 1 month (10.0 ± 0.7 U/g Hb) and 4 months (11.0 ± 0.8 U/g Hb) after partum. In Supplementary Table S3 is reported the statistical significance among e-GST activities from the three different physiological conditions.

Table 1. A comparison of the kinetics parameters of mammalian e-GSTs*

| Species             | K_m (mM) | K_m (mM) | k_cat (s⁻¹) |
|---------------------|----------|----------|-------------|
| Homo sapiens        | 0.11 ± 0.01 | 1.0 ± 0.1 | 79 ± 5      |
| Bos taurus          | 0.12 ± 0.02 | 0.8 ± 0.2 | 83 ± 7      |
| Capra hircus         | 0.14 ± 0.02 | 0.9 ± 0.1 | 85 ± 6      |
| Ovis aries           | 0.10 ± 0.01 | 0.8 ± 0.2 | 77 ± 8      |
| Equus caballus       | 0.10 ± 0.02 | 0.8 ± 0.2 | 82 ± 6      |
| Sus scrofa           | 0.10 ± 0.02 | 0.9 ± 0.1 | 75 ± 7      |

*Data shown are the mean ± S.E.M. from three distinct experiments.

Oxidized e-GST in bovine erythrocyte: reality or artifact?

As mentioned above, the human GSTP1-1 may form an oxidized enzyme involving the two more reactive cysteines (i.e., Cys47 and Cys101).9 The resulting oxidized enzyme is fully inactive.9 The presence of oxidized GSTP1-1 has recently been discovered in human saliva,11 but it has never been observed in mammalian blood. After incubation of hemolyzed total blood with DTT, we observed a conjugation rate of GSH with CDNB higher than that found in the absence of reducing treatment, suggesting the presence of a significant, not spurious amount of oxidized e-GST (Figure 2a). Surprisingly, this additional activity was not recovered in isolated erythrocytes, but only in serum (Figure 2a). In a recent reinvestigation of the possible presence of GSTP1-1 in human serum, we concluded that it was absent or below the detection limit of the usual spectrophotometric assay,12 but we did not verify the possible occurrence of oxidized GSTP1-1. So far, no data are available in the literature concerning the presence of oxidized GSTP1-1 in bovine blood. Further experiments to clarify this point gave surprising results. In fact, the addition of NBDHEX, a strong inhibitor of e-GST and other GST isoenzymes belonging to the alpha, pi and mu classes,13 did not suppress this additional GST activity (Figure 2b), and curiously the activity was also found in the absence of GSH (Figure 2a). Thus, the additional activity observed after reduction with DTT cannot be explained by the presence of oxidized GST in the samples.

Reduced serum albumin simulates a pseudo-GST activity

The above findings indicate that an unknown sulfhydryl compound present in blood and formed during DTT treatment is able to react with CDNB at a high rate. Low molecular mass disulfides can be excluded, because serum depleted of high mass components by filtration does not show any reactivity with CDNB (Figure 2b). Thus, our attention instead focused on proteins and BSA was a likely candidate. This protein is highly concentrated in serum and displays 17 disulfides that can be reduced under DTT conditions, whereas human serum albumin does not show any reactivity with CDNB (data not shown).

Figure 1. e-GST and e-CAT in selected mammalian species. (a) Bovine e-GST in different physiological conditions. Values are the mean of three sets of measurements performed on 40 cows during pregnancy, during lactating phase at 1 month (lactation 0) and at 4 months postpartum (lactation 1). (b) e-GST in humans and in selected mammalian species. (c) e-CAT in humans and in selected mammalian species. Error bars are the S.E.M.
treatment. Purified BSA, reduced as described under Materials and Methods section, used at same serum concentration reacts with CDNB with a velocity identical to that observed using the bovine serum (Figure 2a). This suggests that reduced BSA is responsible for the apparent extra GST activity detected in our samples.

The absence of oxidized e-GST in normal bovine blood samples does not exclude its presence after severe oxidative stress. Thus, the levels of GSH, GSSG and GSP1-1 were determined after incubation of intact erythrocytes with t-BOOH, a well-known trigger of oxidative stress. Although GSH is rapidly oxidized to GSSG, the activity of e-GST remained unchanged, indicating a strong resistance to oxidants (Figure 3). In addition, the enzymatic cell protection system (glutathione reductase) is rapidly involved and causes a restoration of the original concentration of GSH in < 2 h.

e-GST levels in cows raised in highly polluted areas

Preliminary data obtained analyzing blood samples from 10 cows reared in farms located in a well-known polluted area (River Sacco valley, located in the Frosinone district, Lazio, Italy) showed that e-GST is overexpressed, displaying an average value of 18 ± 3 U/g Hb, much higher than that found in the samples obtained from controlled farms located in unpolluted areas (10.7 ± 0.4 U/g Hb) (Figure 1a). This behavior closely parallels the one found for humans living in the same area. 7

**DISCUSSION**

This study shows a first comparison of kinetics and molecular properties of different e-GSTs from mammalian species. An examination of sequence data and kinetics properties of the e-GSTs from pig, goat, cow, sheep and horse demonstrated that all these enzymes are very similar and almost identical to the human GSTP1-1 (Table 1). It is interesting that the basic level of e-GST activity does not seem to be connected with the general level of oxidative stress in the animal; in fact a comparison between e-GST and e-CAT activities shows completely different species profiles (Figure 1).

All these e-GSTs display an extraordinary stability at 4 °C except for the pig e-GST that began to lose activity after 2 days (Supplementary Table S2). As expected, the expression of e-GST...
activity is species-specific; the lowest levels were found in humans and in the pig, whereas the highest activity was observed in the goat. Physiological phases like lactation or pregnancy gave no statistically significant differences (Figure 1a) (Supplementary Table S3). The present study also explores the possibility that the oxidized inactive form of e-GST may be present in blood, given that two critical cysteines, Cys47 and Cys101 (the latter corresponding to Cys103 in pig e-GST) are strictly conserved in all mammals tested, and that they form a disulfide both in vivo and in vitro as proved for the human enzyme.\textsuperscript{13,14} Our data indicate that oxidized e-GST, if present, is below the detection limit of the assay. Even strong oxidative stress does not produce appreciable amounts of oxidized e-GST. An artifact GST-like activity is observed in serum after DTT reduction, mainly due to hyper-reactivity of a few cysteines of serum albumin. This phenomenon must be seriously considered for a correct determination of e-GST in blood. Preliminary results on cows reared in farms residing in a highly polluted area confirm that the e-GST activity is a highly sensitive parameter for detecting increased toxicity levels. As observed in humans,\textsuperscript{7} the overexpression of e-GST in animals is likely a defense response to an increased blood toxicity, and this behavior resembles the increased production of white blood cells in case of bacterial infections. An increased e-GST level in animals is therefore an alarm signal that must be followed up by more accurate investigation to assess the chemical nature of the contaminants.

All distinct e-GSTs from humans, Bos taurus (cow), Sus scrofa (pig), Capra hircus (goat), Equus caballus (horse), Equus asinus (donkey) and Ovis aries (sheep) show very similar amino acid sequences, identical kinetics and stability properties. Reference values for e-GST activity in all these mammals have been determined. Possibly disturbing factors have been examined to avoid artifact determinations. Preliminary data on cows reared in polluted areas showed increased expression of e-GST that parallels the results found for humans. This study provides the basis for future screening studies to verify if animals have been exposed to toxicologic insults.

**MATERIALS AND METHODS**

**Ethics statements**

70 healthy volunteers were recruited for blood recovery after written consent. The study conformed to the standards set by the Declaration of Helsinki. The study was approved by local Ethics Committee of University of Rome Tor Vergata. Animal blood samples have been recovered according to EU Directive 2010/63/EU for animal experiments. The study was carried out in five private dairy farms located on private land, with permission from the owners. This study has been performed employing the blood samples collected as part of farm routine health controls. Specific permissions from Italian Ministry of Health (Ordinance n. 153/2001-A) for animal care and use, and from the Ethics Committee of the University of Rome Tor Vergata, were obtained for these locations/activities and study. As the study has been carried out on dairy cattle and farm animals, the field study did not involve endangered or protected species. None of the animals were sacrificed for this study.

**Blood samples**

All animals tested were randomly chosen from controlled farms of the Latium and Tuscany countrysides. We examined in particular: Bos taurus (n = 40), Sus scrofa (n = 18), Capra hircus (n = 20), Equus caballus (n = 14), Equus asinus (n = 14) and Ovis aries (n = 15). For the Bos taurus species, 40 animals were tested during pregnancy and during lactation, after 1 month and after 4 months postpartum, respectively. As no relevant changes in e-GST levels were observed for bovine species (see Results section), all other mammalian species were tested without considering these physiological phases. Peripheral blood samples were collected from the jugular vein in tubes with tripotassium EDTA in adult subjects. The samples were collected by a veterinarian using the proper procedure of blood collection, and stored at 4 °C for no more than 2 days. Furthermore, a complete blood cell count (hemogram) was performed within 24 h using an automated counter Cell-Dyn 3700. 12 parameters (Abbott Laboratories, Abbott Park, IL, USA) to assess animal health status, mainly the hemoglobin value.

For human blood, 70 healthy volunteers with normal renal function and no history of diabetes mellitus served as healthy controls. Blood samples were collected from the antecubital vein and stored into tripotassium EDTA tubes at 4 °C.\textsuperscript{5}

**Materials**

Reduced glutathione (GSH), oxidized glutathione (GSSG), 1-chloro-2,4-dinitrobenzene (CDNB), dithiothreitol (DTT), chemicals for Lowry solutions, S-hexylglutathione sepharose 6B, tert-buty1 hydroperoxide (t-BOOH), EDTA, N-ethylmaleimide, 4-chloro-7-nitrobenzofurazan, 6-mercaptop-1-hexanol, 3-bromopyruvate, bovine serum albumin (BSA) and all other reagents were purchased from Sigma-Aldrich (St. Louis, MO, USA). The compound 6-(NBD-4-ylthio)-hexanol (NBDHEX) was synthesized as described previously.\textsuperscript{13} Human e-GST was expressed in E.coli and purified as described,\textsuperscript{14} whereas GSTA1-1 and GSTM2-2 were expressed and purified according to a previous study.\textsuperscript{15}

**e-GST activity**

e-GST activity was determined with a spectrophotometric assay at 340 nm (37 °C). 40 μl of whole blood were diluted in 1 ml of bi-distilled water causing erythrocyte hemolysis. After 2 min, 0.1 ml samples were diluted to a final volume of 1 ml containing 1 mM GSH, 1 mM CDNB in 0.1 M potassium phosphate buffer, pH 6.5, according to the standard procedure of Habig et al.\textsuperscript{16} Results were expressed as enzyme units per gram of Hb (U/g Hb).\textsuperscript{15,16} 1 unit represents the amount of enzyme that catalyzes the conjugation of 1 micromole of GSH to CDNB in 1 min at 37 °C. e-GST activity in isolated erythrocyte was determined as above after hemolysis of collected erythrocytes contained in 40 μl of total blood.

**Kinetics parameters**

The $K_{m}$ value for GSH was obtained by reacting about 0.5 μg of purified mammalian e-GSTs with variable amounts of GSH (from 0.02 to 2 mM) in the presence of 1 mM CDNB, in 0.1 M potassium phosphate buffer, pH 6.5 (25 °C). The $K_{m}$ value for CDNB was obtained by reacting 0.5 μg of purified e-GSTs of mammals with variable amounts of CDNB (from 0.05 to 2 mM) in the presence of 1 mM GSH, in 0.1 M potassium phosphate buffer, pH 6.5 (25 °C). From these experimental data, $k_{cat}$ values (at saturating CDNB and GSH) were also calculated.

**Erythrocyte catalase activity**

Erythrocyte catalase (e-CAT) activity was determined with a spectrophotometric assay at 240 nm (25 °C). 5 μl of hemolyzed blood was diluted in 1 ml of potassium phosphate buffer (0.05 M, pH 7.0) with EDTA 0.1 mM, and finally in 10 μl of H$_2$O$_2$ (1 M) according to the standard procedure of Beers and Sizer.\textsuperscript{17} Results were expressed as enzyme units per gram of Hb (U/g Hb).\textsuperscript{15} 1 unit represents the amount of enzyme that catalyzes the decomposition of 1 micromole of H$_2$O$_2$ in 1 min at 25 °C.

**Reactivation procedure for oxidized e-GST**

Different hemolyzed blood samples (total blood, isolated erythrocytes or serum) and purified BSA approximately at the same concentration present in hemolyzed blood (23 μM), were all incubated for 60 min at 37 °C in a dry block thermostat with 1 mM final concentration of DTT and 0.01 M potassium phosphate buffer (pH 8.0).

**Erythrocyte GSH and GSSG determination**

Whole blood (0.5 ml) were reacted with 0.05 ml of 310 mM N-ethylmaleimide, and then GSH and GSSG were determined by following the procedure described previously.\textsuperscript{18}

**Sequence analysis**

Amino acid sequences were selected from NCBI and UniProtKB databases: Homo sapiens (code: P09211), Bos taurus (code: NP_803482.1), Capra hircus (code: XP:218236), Equus caballus (code: XP:001498156.1), Ovis aries (code: XP_004019771.1) and Sus scrofa (code: M3V836). Sequence alignment and percentage of similarities were derived using ClustalW v2.1\textsuperscript{19} available on the website of EMBL.
Purification of e-GST

Purification of e-GST from hemolyzed erythrocytes of all species tested, was performed with a single-step affinity chromatography method using S-hexylglutathione sepharose 6B. Protein concentration was determined using the procedure described by Lowry et al.\textsuperscript{21}

Statistical analysis

All data were expressed as mean±S.E.M. P-values reported in the text and in tables have been estimated on the basis of the mean±S.D. All continuous variables were checked for normality using the Levene's test. One-way analysis of variance (ANOVA) was employed to compare the data between different species. To compare the e-GST activity values of Bos taurus, in different physiological conditions, One-way analysis of variance (ANOVA) was used. A value of P < 0.05 was considered statistically significant. Data were processed using the statistical software MedCalc (Mariakerke, Belgium).

ABBREVIATIONS

DTT, dithiothreitol; GSH, glutathione; GSSG, oxidized glutathione; GST, glutathione transferase; e-CAT, erythrocyte catalase; e-GST, erythrocyte glutathione transferase.

AUTHOR CONTRIBUTIONS

GR planned the experiments. AB and RF performed all the experiments reported in the study and planned the human healthy volunteers recruiting. OL, LA and CR planned the animal blood sample recruiting. GR, along with all other authors, contributed to the writing of the paper.

COMPETING INTERESTS

The authors declare no conflict of interest.

REFERENCES

1. Hayes JD, Fulford DJ. The glutathione S-transferase supergene family: regulation of GST and the contribution of the isoenzymes to cancer chemoprotection and drug resistance. Crit Rev Biochem Mol Biol 1995; 30: 445–600.
2. Armstrong RN. Structure, catalytic mechanism and evolution of the glutathione transferases. Chem Res Toxicol 1997; 10: 2–18.
3. Bocedi A, Fabrini R, Farrotti A, Stella L, Ketterman AJ, Pedersen JZ et al. The impact of nitric oxide toxicity on the evolution of the glutathione transferase superfamily: a proposal for an evolutionary driving force. J Biol Chem 2013; 288: 24936–24947.
4. Adler V, Yin Z, Fuchs SY, Ben Ezra M, Rosario L, Tew KD et al. Regulation of JNK signaling by GSTP. EMBO J 1999; 18: 1321–1334.
5. Dessi M, Noce A, Dawood KF, Galli F, Taccone-Gallucci M, Fabrini R et al. Erythrocyte glutathione transferase: a potential new biomarker in chronic kidney diseases which correlates with plasma homocysteine. Amino Acids 2012; 43: 347–354.
6. Noce A, Ferrannini M, Fabrini R, Bocedi A, Dessi M, Galli F et al. Erythrocyte glutathione transferase: a new biomarker for hemodialysis adequacy, overcoming the Kt/V(urea) dogma? Cell Death Dis 2012; 3: e377.
7. Fabrini R, Bocedi A, Del Grosso E, Morici L, Federici G, Palleschi A et al. Erythrocyte glutathione transferase: a novel biomarker to check environmental pollution hazardous for humans. Biochem Biophys Res Commun 2012; 426: 71–75.
8. Noce A, Fabrini R, Dessi M, Bocedi A, Santini S, Rovella V et al. Erythrocyte glutathione transferase activity: a possible early biomarker for blood toxicity in uremic diabetic patients. Acta Diabetol 2014; 51: 219–224.
9. Ricci G, Del Boccio G, Pennelli A, Lo Bello M, Petruzelli R, Caccuri AM et al. Redox forms of human placenta glutathione transferase. J Biol Chem 1991; 266: 21409–21415.
10. Ricci G, Del Boccio G, Pennelli A, Aceto A, Whitehead EP, Federici G. Nonequivalence of the two subunits of horse erythrocyte glutathione transferase in their reaction with sulfhydryl reagents. J Biol Chem 1989; 264: 5462–5467.
11. Fabrini R, Bocedi A, Camerini S, Fusietti M, Ottaviani F, Passali FM et al. Inactivation of human salivary glutathione transferase P1-1 by hypotiocyanite: a post-translational control system in search of a role. PLoS One 2014; 9: e112797.
12. Fabrini R, Bocedi A, Massoud R, Federici G, Ricci G. Spectrophotometric assay for serum glutathione transferase: a re-examination. Clin Biochem 2012; 45: 668–671.
13. Ricci G, De Maria F, Antonini G, Turella P, Bullo A, Stella L et al. 7-nitro-2,1,3-benzoazadiazole derivatives, a new class of suicide inhibitors for glutathione-S-transferases. Mechanism of action of potential anticancer drugs. J Biol Chem 2005; 280: 26397–26405.
14. Lo Bello M, Battistoni A, Mazzetti AP, Board PG, Muramatsu M, Federici G et al. Site-directed mutagenesis of human glutathione transferase P1-1. J Biol Chem 1995; 270: 1249–1253.
15. De Maria F, Pedersen JZ, Caccuri AM, Antonini G, Turella P, Stella L et al. The specific interaction of dinitrosyl-diglutathionyl-iron complex, a natural NO carrier, with the glutathione transferase superfamily suggestion for an evolutionary pressure in the direction of the storage of nitric oxide. J Biol Chem 2003; 278: 42283–42293.
16. Habig WH, Pabst MJ, Jakoby WB. Glutathione S-transferases. The first enzymatic step in mercapturic acid formation. J Biol Chem 1974; 249: 7130–7139.
17. Beers RF, Sizer IW. A spectrophotometric method for measuring the breakdown of hydrogen peroxide by catalase. J Biol Chem 1952; 195: 133–140.
18. Giustarini D, Dalle-Donne I, Colombo R, Milzani A, Rossi R. An improved HPLC measurement for GSH and GSSG in human blood. Free Radic Biol Med 2003; 35: 1365–1372.
19. Larkin MA, Blackshields G, Brown NP, Chenna R, McGettigan PA, McWilliam H et al. Clustal W and Clustal X version 2.0. Bioinformatics 2007; 23: 2947–2948.
20. Simons PC, Vander Jagt DL. Purification of glutathione S-transferases by glutathione-affinity chromatography. Methods Enzymol 1981; 77: 235–237.
21. Lowry OH, Rosebrough NJ, Farr AL, Randall RJ. Protein measurements with the folin phenol reagent. J Biol Chem 1951; 193: 265–275.

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