Human arylamine N-acetyltransferase I is inhibited by the dithiocarbamate pesticide Thiram

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Abbreviations:
Thiram, tetramethylthiuram disulfide; DTC, dithiocarbamate; NAT1, arylamine N-acetyltransferase 1; XME, xenobiotic-metabolizing enzyme; 3,4-DCA, 3,4-dichloroaniline; 3,5-DCA, 3,5-dichloroaniline; AcCoA, acetyl-coenzyme A; IC_{50}, half maximal inhibitory concentration; k_{obs}, apparent first-order inhibition rate constant; k_{inact}, maximum rate inhibition constant; K_{i}, concentration of inhibitor to achieve half-maximal rate of inhibition; PNPA, p-nitrophenylacetate; PNP, p-nitrophenol; DMAB, 4-dimethylaminobenzaldehyde; 5-IAF; 5-iodoacetamide-fluorescein); DETC; S-methyl-N,N-diethylthiocarbamate; DETC-MeSO_{2}, S-methyl-N,N-diethylthiocarbamate sulfone; DTT, dithiothreitol; HPCL, high pressure liquid chromatography; SD, standard deviation.
Abstract

Thiram (tetramethylthiuram disulfide) is a representative dithiocarbamate (DTC) pesticide used in both the field and as a seed protectant. The widespread use of Thiram and other DTC pesticides has raised concerns for health as these compounds can exert neuropathic, endocrine disruptive and carcinogenic effects. These toxic effects are thought to rely, at least in part, on the reaction of Thiram (and certain of its metabolites) with cellular protein thiols with subsequent loss of protein function. So far, a limited number of molecular targets of Thiram have been reported including few enzymes such as dopamine β-hydroxylase, 11β-hydroxysteroid dehydrogenase and brain glycogen phosphorylase. We provide evidences that Thiram is an inhibitor ($K_I = 23 \ \mu\text{M}; k_{\text{inact}} = 0.085 \ \text{s}^{-1}; k_{\text{inact}}/K_I = 3691 \ \text{M}^{-1}\text{s}^{-1}$) of human arylamine $N$-acetyltransferase 1 (NAT1), a phase II xenobiotic-metabolizing enzyme (XME) which plays a key role in the biotransformation of aromatic amine xenobiotics. Thiram was found to act as an irreversible inhibitor through the modification of NAT1 catalytic cysteine residue as also reported for other enzymes targeted by this pesticide. We also showed using purified NAT1 and human keratinocytes that Thiram impaired the $N$-acetylation of 3,4-dichloroaniline (3,4-DCA), a major toxic metabolite of aromatic amine pesticides (such as Diuron or Propanil). As co-exposure to different classes of pesticides is common, our data suggest that pharmacokinetic drug-drug interactions between DTC pesticides such as Thiram and aromatic amine pesticides may occur through alteration of NAT1 enzymes functions.

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Introduction

Thiocarbamates (including dithiocarbamates) are an important class of pesticides. Among this group of chemicals, Thiram (tetramethylthiuram disulfide) is a representative dithiocarbamate (DTC) pesticide widely used as a fungicide for treatment of crops, vegetables, seeds and ornamental plants as well as a vulcanizing agent in rubber industry (Cereser et al., 2001; Rath et al., 2011; Mathieu et al., 2015). The widespread use of DTC pesticides has raised concerns for human health, in particular for agricultural workers (van Boxtel et al., 2010; Rath et al., 2011). In addition, the possibility of exposure to DTCs through contamination of foodstuff has been put forward as residues of DTC are found often in fruits, vegetables and cereals (van Boxtel et al., 2010). Exposure to DTCs such as Thiram is associated with various toxic effects including renal failure as well as neuronal and developmental toxicity. Reproductive, endocrine disruptive, carcinogenic and teratogenic effects were also reported (Rath et al., 2011; Rasaputra et al., 2013). Exposure to DTCs occurs mainly from their absorption through skin and/or by ingestion and inhalation. In addition, the lipophilic properties of DTCs allow them to pass through cell membranes and physiological barriers including blood brain and fetal-placental barriers (Frank et al., 1995; Rath et al., 2011). The toxic effects of DTCs are thought to rely, at least in part, on the formation of protein covalent adducts that perturb protein functions. In addition, DTC can also alter protein functions through chelation of metal ions (Rath et al., 2011; Viquez et al., 2012; Mathieu et al., 2015). Certain metabolites of DTCs (such as sulfoxide or sulfone forms) are also reactive. The toxic effects of DTC are thus thought to be due to the parent compounds and their metabolites (Jin et al., 1994; Staub et al., 1998; Mathieu et al., 2015; Mathieu et al., 2017). So far, Thiram pesticide has been clearly shown to impair the activity of few enzymes \textit{i.e.} dopamine \(\beta\)-hydroxylase, 11\(\beta\)-hydroxysteroid dehydrogenase, lysyl oxidase and brain glycogen phosphorylase (Lippmann and Lloyd 1969; Caroldi and De Paris 1995; Atanasov et al., 2003; Garbrecht et al., 2006; van Boxtel et al., 2010; Mathieu et al., 2017).

Arylamine N-acetyltransferases (NAT) are phase II XME which catalyze the transfer of an acetyl group from acetyl-coenzyme A (AcCoa) to the nitrogen or oxygen group of aromatic amine chemicals (Hein et al., 2000). As a consequence, these enzymes are key players in the detoxification and/or bioactivation of several aromatic amine drugs and carcinogens (Hein et al., 2000; Sim et al., 2012). In human, there are two NAT isozymes (NAT1 and NAT2) that differ in substrate specificity and tissue distribution. NAT2 is found mainly in liver and gut while NAT1 is ubiquitously expressed (Hein et al., 2000; Butcher and Minchin, 2012; Sim et al., 2012). In addition to xenobiotic metabolism, recent studies indicate that the NAT1 enzyme could...
be involved in other functions. For instance, NAT1 has been shown to act as a folate-dependent acetyl-coenzyme A hydrolase (Laurieri et al., 2014; Stepp et al., 2015). This NAT isozyme has also been reported to play a role in 5-methyltetrahydrofolate and S-adenosylmethionine metabolism (Whitam et al., 2013). More recent data indicate that human NAT1 is involved in methionine salvage pathway (Whitam et al., 2017). Among the very large panel of different aromatic amine chemicals (drugs, dyes, combustion-derived carcinogens, etc.) that are metabolized by NAT enzymes, it is known that these XME contribute significantly to the metabolism of several aromatic amine pesticides (also known as aniline/anilide pesticides) such as diuron, linuron, chlorotoluron or vinclozolin and their toxic by-products such as 3,4-dichloroaniline (3,4-DCA) or 3,5-dichloroaniline (3,5-DCA) (Tweedy et al., 1970; Tixier et al., 2002; Kim and Guengerich 2005; Rodrigues-Lima et al., 2006; Martins et al., 2009). In addition to the genetic mechanisms that govern the expression and activity of NAT enzymes (mainly polymorphisms), non-genetic factors such as certain chemicals (reactive oxygen species, drugs, heavy metals) can also act as inhibitors of these enzymes (Dairou et al., 2005; Rodrigues-Lima et al., 2008; Liu et al., 2008; Butcher and Minchin 2012; Dierolf et al., 2012; Duval et al., 2016). Interestingly, it has been shown that certain pesticides including organophosphates or DTCs, are able to alter xenobiotic metabolism pathways through inhibition of XME in particular CYP P450 (Brady et al., 1991; Hernández et al., 2013).

Using molecular, cellular and enzyme kinetics approaches, we report here that Thiram, a widely used DTC pesticide, is an irreversible inhibitor of human NAT1 enzyme (K_M=23 µM; k_{inact}=0.085 s^{-1}; k_{inact}/K_M=3691 M^{-1}.s^{-1}). We show that the N-acetylation of 3,4-DCA (a major toxic metabolite of aromatic amine pesticides) by purified human NAT1 was readily inhibited by Thiram. Accordingly, we found that exposure of HaCat cells to Thiram led to the impairment of NAT1-dependent acetylation of 3,4-DCA. Taken together, our data indicate that Thiram may impair the NAT1-dependent acetylation of aromatic amines such as 3,4-DCA. Moreover, as concomitant exposure to different type of pesticides is common (in particular in agriculture), our result suggests that pharmacokinetic drug-drug interactions may exist between DTC and aromatic amine pesticides through inhibition of NAT1 enzyme (Hernández et al., 2013).
Materials and Methods

Materials

Thiocarbamate pesticides (triallate, thiobencarb, cycloate, vernolate, S-ethyl-dipropylthiocarbamate (EPTC), molinate, ethiolate, pebulate, butylate), dithiocarbamate pesticides (ammonium pyrrolidinedithiocarbamate (AP) and Thiram), 3,4-dichloroaniline (3,4-DCA), p-aminobenzoic acid (PABA), p-nitrophenylacetate (PNPA) and acetyl-coenzyme A (AcCoA) were from Sigma-Aldrich (Saint-Quentin-Fallavier, France). The sulfone form of S-methyl-N,N-diethylthiocarbamate (DETC-MeSO₂) was from Toronto Research Chemicals Inc (Toronto, Canada). N-acetyl-3,4-DCA was purchased from Interchim (Montluçon, France). PD Minitrap G25 columns for buffer exchange were from GE Healthcare (Aulnay sous Bois, France). All other reagents were purchased from Sigma-Aldrich otherwise noted. The pesticides were dissolved in DMSO at 100 mM concentration.

Expression and purification of recombinant human NAT1

Human NAT1 enzyme was expressed and purified using BL21(DE3) E. coli strain transformed with a pET28a plasmid containing the cDNA of human NAT1 as previously described (Dairou et al., 2004). The purified enzyme was reduced by 10 mM DTT for 10 min in ice prior to dialysis against 25 mM Tris-HCl, 150 mM NaCl, pH 7.5 (reaction buffer). The protein concentration was determined by the Bradford reagent (Bio-rad, Marne la Coquette, France) and purity was assessed by SDS-PAGE and Coomassie staining. Purified recombinant human NAT1 was stored at -80 °C until use.

Assay of the activity of purified recombinant NAT1 using the PNPA method

Human NAT1 activity was assayed using the PNPA method. This approach allows to assay readily the activity of purified NAT enzyme in 96-well plates. This continuous NAT assay is cheap, rapid and has been used in several studies (Mushtaq et al., 2002; Wang et al., 2005; Malka et al., 2009; Laurieri et al., 2014). To this end, 80 µl of mixture containing the NAT1 enzyme was mixed with 10 µl of PABA (500 µM for final concentration) in reaction buffer. The reaction was initiated by the addition of 10 µl PNPA (2 mM final concentration). The formation of the product p-nitrophenol (PNP) was detected by measuring the absorbance at 405 nm. Incubations were conducted at 37°C in a 96-well plate and the reaction rate was determined by monitoring continuously the absorbance at 405 nm using a plate reader (BioTek, Colmar, France). Although
the non-specific hydrolysis of PNPA in absence of enzyme or PABA was found to be less than 1% of the hydrolysis of PNPA in presence of enzyme or PABA, the data were nonetheless corrected by subtracting the non-specific hydrolysis of PNPA in the absence of enzyme.

**Effect of Thiram on the N-acetylation of 3,4-DCA by purified recombinant NAT1**

The 4-Dimethylaminobenzaldehyde (DMAB) method was used to assess the effect of Thiram on the acetylation of 3,4-dichloroaniline (3,4-DCA) by purified NAT1. This NAT assay allows to measure the rates of N-acetylation of arylamine substrates by NAT enzymes. This approach is rapid, cheap and can be used in 96-well plates with either purified NAT enzymes or with cell extracts. This assay has been used in several studies where it replaced more tedious HPLC approaches (Sinclair et al., 1998; Kawamura et al., 2008; Russel et al., 2009; Laurieri et al., 2014). To this end, NAT1 (2.5 µM final concentration) was first incubated with Thiram (40 µM final concentration) for 30 min at 37 °C in a total volume of 50 µl. The mixture was then diluted (40 times) with reaction buffer (25 mM Tris-HCl, pH 7.5). Aliquots (3 µl) were removed and assayed in a total volume of 50 µl containing 3,4-DCA (500 µM final concentration) and AcCoA (500 µM final concentration) for 30 min at 37°C. The reaction was quenched by adding 40 µl of cold trichloroacetic acid (40% vol/vol in water). Finally, 100 µl of DMAB (5% w/v, in 9:1 acetonitrile and water) were added to the mixture and absorbance at 450 nm was measured in a 96-well plate reader. The amount of residual (non-acetylated) 3,4-DCA was determined from a standard curve of 3,4-DCA.

**Effects of pesticides on human NAT1**

Pesticides (0 – 100 µM final concentration) were independently mixed with human NAT1 (2.5 µM final concentration) in reaction buffer (25 mM Tris-HCl, 150 mM NaCl, pH 7.5). Total volume of the reaction was 50 µl. Upon incubation for 30 min at 37°C, aliquots (5 µl) were diluted (40 times) with reaction buffer and 80 µl were taken for measurement of the residual human NAT1 activity using the PNPA assay as described above. Enzyme activity was expressed as the percentage of the control (no pesticide). In all assays, the final concentration of enzyme was 50 nM. Controls were carried out in presence of DMSO.

To determine whether reducing agents could reactivate Thiram-inhibited human NAT1, the enzyme (2.5 µM final) was first preincubated with Thiram (40 µM final concentration, 30 min at 37 °C) and further
incubated with dithiothreitol (DTT, 1 – 10 mM final concentration) for 10 min at 37 °C in a total volume of 50 µl. Aliquots (5 µl) were diluted (40 times) with reaction buffer and 80 µl were taken for measurement of the residual human NAT1 activity using the PNPA assay as described above. Controls were carried out without Thiram, without DTT and with DTT only.

To investigate whether reducing agents could protect human NAT1 from the inhibitory effect of Thiram, the enzyme was incubated with Thiram (40 µM final concentration) in presence or absence of different concentration of DTT (1 – 10 mM final concentration) for 30 min at 37 °C in a total volume of 50 µl. Aliquots (5 µl) were diluted (40 times) with reaction buffer and 80 µl were taken for measurement of the residual human NAT1 activity using the PNPA assay as described above. Controls were carried out without Thiram, without DTT and with DTT only.

To test the ability of the NAT1 enzyme cofactor AcCoA to protect the enzyme from the inhibitory effect of Thiram, NAT1 was incubated with Thiram (40 µM final concentration) in presence or absence of AcCoA (1 mM final concentration) for 30 min at 37 °C in a total volume of 50 µl. Aliquots (5 µl) were diluted (40 times) with reaction buffer and 80 µl were taken for measurement of the residual human NAT1 activity using the PNPA assay as described above. Controls were carried out without Thiram, with AcCoA only and without AcCoA.

To determine whether the inhibition of human NAT1 by Thiram is irreversible, NAT1 (2.5 µM final concentration) was incubated with or without Thiram (40 µM final concentration) for 30 min at 37°C in a total volume of 100 µl. Aliquots (25 µl) were diluted (40 times) with reaction buffer to a total volume of 1 ml. Eighty µl were taken for measurement of the residual human NAT1 activity using the PNPA assay as described above. In parallel, 500 µl were buffer exchanged using a PD Minitrap G25 column (equilibrated with reaction buffer). Eighty µl of buffer-exchanged samples were assayed for residual human NAT1 activity using the PNPA assay as described above.

**Labeling of cysteine residues of human NAT1 with fluorescein-conjugated iodoacetamide**

The modification of NAT1 cysteine residues by Thiram was monitored using 5-IAF (5-iodoacetamide-fluorescein) labeling, as previously described (Duval et al., 2016). Briefly, human NAT1 (2.5 µM final concentration) was incubated with or without Thiram (0 – 25 µM final concentration) for 30 min at 37°C, in
25 mM Tris-HCl, 150 mM NaCl, pH 7.5. Then 5-IAF (100 µM, final concentration) was added and the mixture incubated for 10 min at 37 °C in the dark. Samples were then subjected to SDS-PAGE and transferred on nitrocellulose membrane. 5-IAF labeling was detected by fluorescence measurements ($\lambda_{\text{exc}}$: 492 nm; $\lambda_{\text{em}}$: 520 nm). The membrane was also probed for recombinant NAT1 protein by immunodetection using a monoclonal anti-6X Histidine-tag antibody (Sigma-Aldrich).

**Kinetic analysis of inhibition of NAT1 by Thiram**

For the kinetic analysis of Thiram-dependent NAT1 inactivation, NAT1 (1 µM final concentration) was incubated with Thiram (final concentration of 2.5–10 mM) at 37 °C in reaction buffer (total volume 50 µl). At various time intervals (30 s), aliquots (5 µl) were removed and diluted 40 times with reaction buffer. Eighty ul of each sample were assayed for residual human NAT1 activity using the PNPA assay as described above. The kinetic data were analyzed as reported in Cornish-Bowden (2001) and Copeland (2005) for irreversible inhibitors using OriginPro 8.1 program (RITME Informatique, Paris, France). The apparent-first order inhibition rate constant ($k_{\text{obs}}$) for each Thiram concentration, was calculated from the linear regression of the natural logarithm of the percentage residual activity versus time ($\ln (\% \text{ residual activity}) = k_{\text{obs}} \times t$, where $t$ is time) (Equation 1). The $k_{\text{obs}}$ values were plotted as a function of Thiram concentration and the data fitted to the following equation: $k_{\text{obs}} = (k_{\text{inact}} \times I)/(K_I + I)$ (Equation 2) where $k_{\text{obs}}$ is the apparent-first order inhibition rate constant, $k_{\text{inact}}$ is the first order maximum rate inhibition constant, $I$ is the concentration of inhibitor (Thiram) and $K_I$ the concentration of inhibitor that achieves half-maximal rate of inhibition. The effectiveness of the inhibition can thus be quantified using the second-order rate constant $k_{\text{inact}}/K_I$ (Copeland, 2005).

**Effects of Thiram on the acetylation of 3,4-DCA by cultured HaCat cells**

Acetylation of 3,4-DCA in cultured HaCat cells was monitored by High Pressure Liquid Chromatography (HPLC). This approach allows to monitor qualitatively and quantitatively both 3,4-DCA and N-acetylated 3,4-DCA in complex media such as the culture medium of cells. This method has been used widely used to assess the acetylation activity of NAT enzymes, including NAT1, in cellular contexts (Butcher et al., 2000; Butcher et al., 2004; Martins et al., 2009; Doll et al., 2010).

HaCat keratinocyte cells (a gift of Pr. B. Friguet, CNRS Université Paris 6 UMR 8256) were grown in 6-well plates at 37 °C under 5 % CO₂, in DMEM supplemented with 10% (v/v) fetal bovine serum. Cells were
seeded at a density of 1000 cells/well in triplicate and grown for 24 h. Prior to treatment, HaCat cells were washed with PBS buffer and then exposed to Thiram or DETC-sulfone (0 - 50 µM) for 1 h at 37 °C. 3,4-DCA (500 µM final) was then added to the culture medium. Aliquots of culture medium were taken every hour (for up to 3 h) for HPLC analysis of 3,4-DCA and N-acetylated 3,4-DCA. To this end, 20 µl of medium was mixed with 20 µl perchloric acid (15% in water). After centrifugation (10 min at 10,000 g), 20 µl of supernatant were injected into a Kromasil Eternity C18 reverse-phase HPLC column (Sigma Aldrich, St-Quentin-Fallavier, France) (Martins et al., 2009). 3,4-DCA and N-acetylated 3,4-DCA were used as standards and results were normalized using the protein concentration of whole cell lysates.

**Statistical analysis**

Analysis of Variance (ANOVA) or Student’s t-test was calculated by Prism 5.03 (GraphPad Software, La Jolla, USA). ANOVA analysis was followed by Dunnett’s multiple comparison test.
Results

The dithiocarbamate pesticide Thiram inhibits human NAT1

As stated above, thiocarbamate pesticides have been shown to react with thiol groups in proteins (Rath et al., 2011; Viquez et al., 2012; Mathieu et al., 2015). In addition, the structural similarity of the thioester group of the NAT cofactor AcCoA with thiocarbamates suggested that these chemicals could react with the catalytic cysteine of human NAT1 in a manner similar to that of AcCoA (Supplemental Figure 1). In addition, Disulfiram, a clinically used DTC has been shown to inhibit the activity of NAT1 (Malka et al., 2009). In order to determine whether thiocarbamate pesticides could impair the acetylation of aromatic amine xenobiotics by NAT1, we first investigated the impact of 11 different thiocarbamate pesticides (including two dithiocarbamates AP and Thiram) on human NAT1 activity (Supplemental Figure 1). Among the different compounds tested (data not shown), Thiram was the most effective at yielding almost full inhibition of NAT1 at 20 µM concentration (60 ±6 % SD and 90 ±9 % SD inhibition obtained with 10 and 20 µM Thiram, respectively). As shown in Figure 1, further analyses with Thiram confirmed that NAT1 was inhibited in a dose-dependent manner by this DTC pesticide with an IC$_{50}$ value of 9 ± 0.5 SD µM.

Human NAT1 is irreversibly inhibited by Thiram through the modification of its catalytic cysteine

To determine whether NAT1 cysteine residues are modified upon exposure to Thiram, we performed chemical-labeling experiments using 5-iodoacetamide-fluorescein (5-IAF), an alkylating agent that covalently and specifically labels free thiol groups. As shown in Figure 2A, the incubation of human NAT1 with increasing concentrations of Thiram led to a dose-dependent modification of NAT1 cysteine residues by Thiram which is evidenced by the dose-dependent loss of the fluorescence signal. In addition, the modification of NAT1 cysteine residues by Thiram correlated well with the dose-dependent inhibition of the enzyme (Figure 1). To investigate whether the loss of NAT1 activity was due to the modification of the catalytic cysteine residue present in the active site of human NAT1, we performed substrate protection experiments using AcCoA. This natural cofactor of NAT enzyme is known to acetylate specifically the catalytic cysteine of NAT1 thus protecting it from reaction with chemicals as described by Liu et al., (2008). As shown in Figure 2B, AcCoA fully protected the enzyme from inhibition by Thiram thus indicating that inhibition of NAT1 activity by Thiram occurred through the modification of the catalytic cysteine.
Thiram has been shown to inhibit irreversibly the activity of 11-β hydroxysteroid dehydrogenase 2 through a modification of an active site cysteine that could not be reversed by the reducing agent DTT (Atanasov et al., 2003; Garbrecht et al., 2006). Therefore, to further investigate the molecular mechanism of the inhibition of human NAT1 by Thiram, we determined the ability of the reducing agent DTT to reactivate the Thiram-inhibited enzyme. As shown in Figure 2C, DTT was found to poorly reactivate the inhibited enzyme with only 30% of the activity restored at high concentration of DTT (10 mM). As observed with 11-β hydroxysteroid dehydrogenase 2 (Atanasov et al., 2003), these results suggested that the irreversible inhibition of NAT1 was due to the formation of a stable adduct on the catalytic cysteine.

We also investigated the ability of DTT to prevent NAT1 from the inhibition by Thiram by co-incubating the enzyme with Thiram and DTT. As shown in Figure 2D, DTT was able to afford full protection against Thiram-dependent inhibition only at high concentration (10 mM). At 1 and 5 mM concentrations of DTT, the protection was partial (with ≈50% inhibition observed despite the presence of 5 mM DTT). This data suggest that Thiram reacts more rapidly with the catalytic cysteine of NAT1 than with the thiol group of DTT which is in agreement with the reactive nature of the catalytic cysteine residue of NAT1 (Rodrigues-Lima et al., 2001; Liu et al., 2008; Malka et al., 2009; Sim et al., 2012).

Finally, to confirm that Thiram acts as an irreversible inhibitor of human NAT1, Thiram-inhibited NAT1 was buffer-exchanged and assayed for NAT1 activity. As shown in Figure 3, buffer-exchange did not yield recovery of the activity of NAT1 enzyme previously exposed to Thiram. This experiment supports that NAT1 inhibition by Thiram occurs through an irreversible mechanism, which is agreement with the data obtained above (Figure 2).

**Kinetic analysis and determination of the second order rate constant of NAT1 inhibition by Thiram**

We then performed kinetic analysis to further characterize the inhibition of human NAT1 by Thiram. To this end, we carried out time course inhibition of the NAT1 enzyme at different Thiram concentrations (Figure 4A). The enzyme was found to be inhibited in a time and dose-dependent manner by Thiram as shown in the plot of $k_{obs}$ values (apparent-first order inhibition rate constants for each Thiram concentration) as function of Thiram concentration (Figure 4B). The data fitted well to the two-steps mechanism of irreversible inhibition equation ($k_{obs}=(k_{inact} x I)/(K_I+I)$) indicating that the inhibition by Thiram occurred through a two-step mechanism of inhibition that involves reversible binding of the inhibitor to enzyme followed by an irreversible
step (Copeland, 2005). The $K_I$ was found to be equal to 23 $\mu$M and the $k_{inact}$ to be equal to 0.085 s$^{-1}$, which give a second-order rate constant of $k_{inact}/K_I$=3691 M$^{-1}$s$^{-1}$. This rate of inhibition is higher than the rate of inhibition of NAT1 by acrolein, an aldehyde chemical, found to irreversibly inhibit the enzyme through a similar two-step mechanism ($k_{inact}/K_I$=57 M$^{-1}$s$^{-1}$) (Bui et al., 2013).

Thiram impairs 3,4-DCA acetylation by purified human NAT1 and in cultured HaCat cells

3,4-DCA is a major toxic metabolite of aromatic amine pesticides known to be biotransformed by NAT enzymes (Tixier et al., 2002; Martins et al., 2009; Yuan et al., 2017). We analyzed the impact of Thiram on the biotransformation of this pesticide by human NAT1. As shown in Figure 5A, incubation of purified NAT1 with Thiram led to the inhibition of the N-acetylation of 3,4-DCA. As skin is a major route of exposure to Thiram and aromatic amine pesticides, cellular experiments using the well-known human keratinocyte cell line HaCat were carried out. HaCat cells are known to express NAT1 enzyme and to display a functional NAT1-dependent aromatic amine acetylation pathway (Bonifas et al., 2010; Scheitza et al., 2012). HaCat cells were exposed to Thiram and 3,4-DCA, and the N-acetylation of 3,4-DCA by endogenous NAT1 was followed using HPLC methods (Rodrigues-Lima et al., 2006; Martins et al., 2009). As shown in Figure 5B, exposure to Thiram impaired the N-acetylation of 3,4-DCA mediated by NAT1 in a dose-dependent manner.

Thiocarbamates including Thiram, can be biotransformed into different reactive metabolites which can also react with cysteine residues. In particular, thiocarbamates can undergo oxidation, forming successively a sulfoxide and a final sulfone metabolite which can react with thiol groups in proteins (Lipsky et al., 2001; Viquez et al., 2012; Mathieu et al., 2015). As the sulfone metabolite of Thiram, DMTC-sulfone, is not commercially available, we investigated the impact of DETC-sulfone on recombinant NAT1. This compound differs only from DMTC-sulfone by a methyl group. In addition, Disulfiram, the dithiocarbamate that is the parent compound of DETC-sulfone, is also an irreversible inhibitor of NAT1 (Malka et al., 2009). As observed with Thiram, we found that exposure to DETC-sulfone led to a dose-dependent inhibition of recombinant human NAT1 (with an $IC_{50}$ of 1.7 ± 0.1 $\mu$M) (Supplemental Figure 2A). This inhibition was concomitant with a loss of 5-IAF fluorescence signal, suggesting that the inhibition was also associated with the modification of cysteine residues of the enzyme as shown for Thiram (Supplemental Figure 2B). Thus, altogether, these data supported that NAT1 is a target of Thiram pesticide but also of its sulfone metabolite.
Discussion

Thiocarbamates (including dithiocarbamates) are organosulfur compounds extensively used as pesticides. Exposure to these compounds is associated to different toxic effects (Rath et al., 2011; Mathieu et al., 2015). However, very little is known about the possible metabolic interactions between thio/dithiocarbamates and other class of xenobiotics including pesticides (Boobis et al., 2008; Hernández et al., 2013). Here we report that the xenobiotic-metabolizing enzyme NAT1 is inhibited by Thiram, a prototypic thiocarbamate widely used as pesticide. We provide mechanistic and kinetic evidence that the inhibition of NAT1 by Thiram occurs through the irreversible modification of the reactive catalytic cysteine of the enzyme based on a classical two-steps mechanism (Copeland, 2005). The toxic effects of Thiram are thought to rely, at least in part, on its reaction with cysteine residues in target proteins which drives their loss of function (Rath et al., 2011). Our results are in agreement with studies carried out on 11β-hydroxysteroid dehydrogenase showing that Thiram reacts with cysteine residue in the active site leading to its irreversible inhibition (Atanasov et al., 2003). Similarly, we found that Thiram-inhibited NAT1 was only slightly reactivated by high concentrations of the reducing agent such as DTT indicating that the disulfide adduct formed upon reaction of the catalytic cysteine with Thiram is stable and not readily reducible by DTT as also observed with 11β-hydroxysteroid dehydrogenase (Atanasov et al., 2003). We also found that oxidative metabolites of Thiram such as its sulfone metabolite may also impair the activity of NAT1. Interestingly, sulfone metabolites of thiocarbamate chemicals (such as DETC-sulfone the metabolite of Disulfiram) have been shown to react with cysteine residues and to also impair enzyme activity through covalent reaction with cysteines (Hu et al., 1997; Zimmerman et al., 2004).

So far, a limited number of enzymes have been reported to be inhibited by Thiram (Lippmann and Lloyd 1969; Caroldi and De Paris 1995; Atanasov et al., 2003; Garbrecht et al., 2006; van Boxtel et al., 2010; Mathieu et al., 2017). In addition to these enzymes, our data show that Thiram is also able to inhibit a phase II xenobiotic-metabolizing enzyme known to play a major role in the metabolism of arylamine xenobiotics. While other DTCs had been shown to alter the activity of certain cytochromes P450 or GST, so far no study had addressed the effect of Thiram on a XME (Rath et al., 2011; Mathieu et al., 2015). NAT enzymes are known to acetylate a large variety of aromatic amines xenobiotics including drugs, carcinogens, pesticides and their toxic metabolites such as 3,4-DCA and thus to impact their pharmacological and/or toxicological outcome (Kim and Guengerich 2005; Martins et al., 2009; Sim et al., 2012). We found that Thiram was able to impair
the NAT-dependent acetylation of 3,4-DCA, a well-known toxic metabolite of several aromatic amine pesticides (Tixier et al., 2002; Martins et al., 2009; Yuan et al., 2017). In addition, we also showed that exposure of cultured human keratinocyte cells to Thiram led to the impairment of the N-acetylation of 3,4-DCA by NAT1. Altogether these data point to potential pharmacokinetic drug-drug interactions between Thiram and aromatic amine xenobiotics metabolized by NAT1. In particular, as pesticides occurs frequently as mixtures, our results suggest possible metabolic interactions between DTCs such as Thiram and aniline pesticides (such as linuron or propanil) through alteration of NAT enzyme functions. Pharmacokinetic drug-drug interactions between different classes of pesticides have been reported (Hernández et al., 2013). These are often the result of one pesticide altering the absorption, distribution, metabolism or elimination of the others. For instance, it has been shown that certain organophosphate pesticides decrease the organism’s ability to detoxify pyrethroid pesticides due to esterase inhibition (Hernández et al., 2013).

To conclude, using a combination of molecular, kinetic and cellular studies, we provide evidence that Thiram, a widely used dithiocarbamate pesticide, impairs the NAT1-dependent acetylation of xenobiotics, including toxic aromatic pesticide metabolites. More broadly, our data suggest that pharmacokinetic drug-drug interactions between certain DTC pesticides and aromatic amine pesticides may occur through the inhibition of NAT1 enzyme.
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Authorship contributions

Participated in research design: XX, CM, FRL

Conducted experiments: XX, CM, JB, RD, LCB, FB

Performed data analysis: XX, CM, JB, RD, LCB, FB, JMD, FRL

Wrote or contributed to the writing of the manuscript: XX, CM, JB, FB, FRL
References


Footnotes

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Legend Figures

Figure 1. Dose-dependent inhibition of human NAT1 by Thiram. Human NAT1 was incubated with increasing concentrations of Thiram (0 - 40 µM final concentration) for 30 min at 37 °C, prior to activity measurement using the PNPA assay. Data represent mean values ± SD of triplicate experiments.

Figure 2. Effects of the reducing agent DTT and the cofactor AcCoA on the Thiram-dependent inhibition of NAT1.

(A) 5-IAF Labeling of free cysteines in human NAT1 treated with Thiram. Thiram-treated NAT1 (40 µM final) was further incubated with 5-IAF (100 µM final concentration) for 10 min at 37 °C. Samples were run on SDS-PAGE, transferred to nitrocellulose membranes and 5-IAF detected by measuring the fluorescence (λexc : 492 nm ; λem : 520 nm). NAT1 protein was revealed by immunodetection using an anti-6xHis-tag antibody.

(B) Protective effects of the NAT1 cofactor AcCoA against Thiram-dependent inhibition of human NAT1. Human NAT1 was incubated with Thiram (40 µM final) in presence of AcCoA (1mM) for 30 min at 37 °C and assayed for NAT1 activity using the PNPA assay. *, p < 0.05 compared to the control; #, p < 0.05 compared to the Thiram-treated enzyme. Data represent mean values ± SD of triplicate experiments.

(C). Reactivation of human NAT1 by the reducing agent DTT. Thiram-inhibited NAT1 (40 µM) was further incubated with DTT for 15 min prior to activity assay using the PNPA assay. *, p < 0.05 compared to the controls (no Thiram); #, p < 0.05 compared to the Thiram-treated enzyme (no DTT). Data represent mean values ± SD of triplicate experiments.

(D). Protective effects of the reducing agent DTT against Thiram-dependent inhibition of human NAT1. Human NAT1 was incubated with Thiram (40 µM final) in the presence of different concentrations of DTT (1, 5, 10 mM) for 30 min at 37 °C, and then assayed for NAT1 activity using the PNPA assay. *, p < 0.05 compared to the controls (no Thiram); #, p < 0.05 compared to the Thiram-treated enzyme (no DTT). Data represent mean values ± SD of triplicate experiments.

Figure 3. Effect of buffer-exchange on the inhibition of NAT1 enzyme by Thiram. NAT1 enzyme (2.5 µM) was first incubated for 30 min at 37°C with or without Thiram (40 µM). The samples were assayed for NAT1 activity using the PNPA assay before (left panel) and after buffer exchange with PD Minitrap G25
columns (right panel). *, p < 0.05 compared control (no Thiram). Data represent mean values ± SD of triplicate experiments. Statistics analysis was done by Student’s t-test.

Figure 4. **Kinetics analysis of human NAT1 inhibition by Thiram.**

(A) **Time- and concentration-dependent inhibition of NAT1.** NAT1 (1 µM) was incubated with Thiram different concentrations of Thiram (0, 2.5, 5 and 10 µM) at 37°C. Aliquots were taken every 30 s, diluted 40 times with reaction buffer. Then, the residual NAT1 activity was measured using the PNPA assay. The data were analyzed as reported in Cornish-Bowden (2001) and Copeland (2005) for irreversible inhibitors using OriginPro 8.1 program. Briefly, the apparent-first order inhibition rate constant (k_{obs}) for each Thiram concentration was obtained by linear regression from plots of the natural logarithm of the percentage residual activity versus time (equation 1, see Materials and methods section).

(B) **Determination of inhibition constants K_I and k_{inact}.** In order to obtain the first order maximum rate inhibition constant (k_{inact}) and K_I (the concentration of Thiram that achieves half-maximal rate of inhibition), the data were fitted to k_{obs} = (k_{inact} x I)/(K_I + I) (equation 2, see Materials and methods section). The second-order rate inhibition constant is equal to k_{inact}/ K_I. The R-squared value of the fit is shown.

Figure 5. **Thiram inhibits 3,4-DCA N-acetylation by recombinant human NAT1 and in HaCat cells.**

(A) **Inhibition of 3,4-DCA acetylation by purified human NAT1.** The enzyme was incubated with Thiram (40 µM final) for 30 min at 37°C prior to incubation with 3,4-DCA and AcCoA for 30 min at 37°C. NAT1 activity towards 3,4-DCA was then measured using the DMAB method as described in the Materials and Methods section. *, p < 0.05 when compared to control. Data represent mean values ± SD of triplicate experiments. Statistics analysis was done by Student’s t-test.

(B) **Thiram inhibits the endogenous N-acetylation of 3,4-DCA in cultured HaCat cells.** HaCat cells were exposed or not to different concentrations of Thiram (0-50 µM) for 1 h at 37°C in PBS Ca^{2+}/Mg^{2+}. Cells were then cultured in DMEM medium containing 500 µM 3,4-DCA for 3 h. Every hour, aliquots of the cell culture medium were taken and the amount of N-acetylated 3,4-DCA measured by HPLC. *, p < 0.05 when compared to the control. Data represent mean values ± SD of triplicate experiments.
Figure 2

(A) Western blot showing the effect of Thiram on 5-IAF and Anti-6xHis-tag proteins.

(B) Graph showing the activity (% of Control) of Control, AcCoA, Thiram, and Thiram + AcCoA. Thiram (40 μM) and DTT (mM) conditions are indicated.

(C) Graph showing the activity (% of Control) of Thiram (40 μM) and DTT (mM) conditions. Thiram and DTT concentrations are indicated.

(D) Graph showing the activity (% of Control) of Thiram (40 μM) and DTT (mM) conditions. Thiram and DTT concentrations are indicated.

* and # symbols indicate significant differences compared to control and Thiram treatments, respectively.
Figure 3
Figure 4
Figure 5

A

B

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