Characterization and Comparative Analysis of 2,4-Toluene Diisocyanate and 1,6-Hexamethylene Diisocyanate Haptenated Human Serum Albumin and Hemoglobin

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Characterization and comparative analysis of 2,4-toluene diisocyanate and 1,6-hexamethylene diisocyanate haptenated human serum albumin and hemoglobin

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Research paper

Diisocyanates (dNCOs) are low molecular weight chemical sensitizers that react with autologous proteins to produce neoantigens. dNCO-haptenated proteins have been used as immunogens for generation of dNCO-specific antibodies and as antigens to screen for dNCO-specific antibodies in exposed individuals. Detection of dNCO-specific antibodies in exposed individuals for diagnosis of dNCO asthma has been hampered by poor sensitivities of the assay methods in that specific IgE can only be detected in approximately 25% of the dNCO asthmatics. Apart from characterization of the conjugates used for these immunoassays, the choice of the carrier protein and the dNCO used are important parameters that can influence the detection of dNCO-specific antibodies. Human serum albumin (HSA) is the most common carrier protein used for detection of dNCO-specific-IgE and -IgG but the immunogenicity and/or antigenicity of other proteins that may be modified by dNCO in vivo is not well documented. In the current study, 2,4-toluene diisocyanate (TDI) and 1,6-hexamethylene diisocyanate (HDI) were reacted with HSA and human hemoglobin (Hb) and the resultant adducts were characterized by (i) HPLC quantification of the diamine produced from acid hydrolysis of the adducts, (ii) 2,4,6-trinitrobenzene sulfonic acid (TNBS) assay to assess extent of cross-linking, (iii) electrophoretic migration in polyacrylamide gels to analyze intra- and inter-molecular cross-linking, and (iv) evaluation of antigenicity using a monoclonal antibody developed previously to TDI conjugated to Keyhole limpet hemocyanin (KLH). Concentration-dependent increases in the amount of dNCO bound to HDI and TDI, cross-linking, migration in gels, and antibody-binding were observed. TDI reactivity with both HSA and Hb was significantly higher than HDI. Hb–TDI antigenicity was approximately 30% that of HSA–TDI. In conclusion, this data suggests that both, the extent of haptenation as well as the degree of cross-linking differs between the two diisocyanate species studied, which may influence their relative immuno- genicity and/or antigenicity.

1. Introduction

Diisocyanates (dNCOs) are highly reactive chemicals used as cross-linking agents in the manufacture of polyurethane products such as paints, elastomers, and adhesives (Vangronsveld et al., 2013; Petskon et al., 2000; Arnold et al., 2012). They are potent sensitizers and are a commonly reported cause of occupational chemical hypersensitivity reactions including asthma (Buyantsiwa et al., 2011; Ribeiro et al., 2014). 2,4-Toluene diisocyanate (2, 4-TDI) and 1, 6-hexamethylene diisocyanate (HDI) are among the most widely used isocyanates. Both have high vapor pressures (Sullivan and Krieger, 2001) and exposure often occurs through inhalation of vapors and aerosols during spraying operations at workplaces.

Immune-mediated hypersensitivity reactions to dNCOs include allergic rhinitis (Matheson et al., 2005), asthma (Mapp, 2001), hypersensitivity pneumonitis (Baur, 1995; Charles et al., 1976) and allergic contact dermatitis (Aalto Korte et al., 2012). Although most reported cases of isocyanate sensitization occur at workplaces (Rudzinski et al., 1998; Redlich and Karol, 2002; Hur et al., 2008), it has been suggested that non-occupational exposure to the general public may also occur through the use of “do-it-yourself” free diisocyanate containing commercial products such as polyurethane foams and sprays (Krone, 2004; Wilder et al., 2011). Once allergic sensitization to isocyanates occurs, asthmatic reactions may be triggered by exceedingly minute concentrations of isocyanates (Ribeiro et al., 2014; Ruwona et al., 2010; Wisnewski et al., 2012).

Diisocyanates are low-molecular-weight compounds that must first react with autologous proteins to produce a functional antigen (Wisnewski et al., 2004). The fate of the dNCO in the body and the
protein adducts responsible for immunological sensitization remain unknown (Mapp, 2001). Apart from reacting with proteins at the site of exposure, protein conjugation by dNCOs may also occur via glutathione (GSH) thio-carbamate intermediates. GSH is abundant in the airways and Wisnewski and colleagues demonstrated that albumin can be conjugated to TDI and HDI by GSH–TDI and GSH–HDI, respectively (Wisnewski et al., 2013). HSA is the most common carrier protein used for dNCO antibody immunoassays (Budnik et al., 2013) due to its prevalence in plasma (Wisnewski et al., 2000) to form dNCO adducts. Other molecules, such as keratin 18 (Wisnewski et al., 2000), tubulin (Lange et al., 1999), and the peptide glutathione (Lantz et al., 2001), have been found to be modified by dNCO exposure. Sabbioni and coworkers reported MDI bound to the N-terminal valine of Hb in MDI exposed rats and proposed Hb–MDI as a biological marker of MDI exposure (Sabbioni et al., 2000).

There is currently no simple diagnosis for dNCO-induced occupational asthma (OA) (Wisnewski, 2007). One approach that can potentially be used is testing for dNCO-specific IgE from a worker’s serum. For confirmation of the diagnosis of dNCO as the etiological agent of the occupational asthma, these assays have been reported to be specific (96–98%), but not sensitive (18–27%) (Ott et al., 2007). These low sensitivities have been attributed to both assay limitations and potential IgE-independent dNCO asthma mechanisms (Budnik et al., 2013). Immunoassay standardization is critical for improvement of immunoassay sensitivity and comparison of results across studies (Wisnewski et al., 2004). A number of factors that may confound results from these immunoassays include the choice of dNCO used, the carrier protein employed, dNCO–protein reaction conditions, and post-reaction processing and characterization of the haptenated protein.

Wisnewski et al., in separate studies, reported differences in reactivity between TDI and HDI toward glutathione (GSH). Albumin was conjugated to TDI and HDI by GSH–TDI (Wisnewski et al., 2011) and GSH–HDI (Wisnewski et al., 2013), respectively. From these two reports, the kinetics of GSH–HDI mediated albumin carbamoylation was substantially slower compared with those of GSH–TDI. The hydrolysis of aliphatic isocyanates is also much slower than aromatic isocyanates. However, the nature and extent of HDI and TDI conjugation, in vivo, to serum proteins has not yet been reported. Diisocyanate haptenated proteins have been used both to produce specific antibodies (Lemons et al., 2014; Ruwona et al., 2011) and to screen for dNCO specific antibodies in workers’ sera for diagnosis of OA (Tee et al., 1998). However, these conjugates are often poorly characterized and non-standardized.

Our previous work focused on characterization of methylene diphenyl diisocyanate (MDI)–HSA and MDI–Hb conjugates (Mihke et al., 2013). Although HSA is the most common carrier protein used for dNCO antibody immunoassays (Budnik et al., 2013; Wisnewski et al., 2000), other proteins, however, have also been found to be potentially modified by dNCO exposure (Wisnewski et al., 2000; Lange et al., 1999; Lantz et al., 2001; Sabbioni et al., 2000; Sabbioni and Beyerbach, 2000; Sabbioni et al., 2001) but the immunogenicity of added proteins other than albumins has not been reported.

One hypothesis is that the lack of a standard characterization protocol for conjugates used to screen for dNCO specific antibodies in workers’ sera is contributing to the reported low sensitivities and variability of these assay methods. Our previous work on MDI shed light on the need to use multiple methods to characterize these conjugates. In the present study we extend the characterization of the dNCO-protein conjugates from MDI to understand reactivity differences among dNCOs such as TDI and HDI that can impact assay sensitivities and standardization protocols. Quantification of the amount of TDI and HDI bound per mole protein was conducted by analyses of the corresponding hydrolysis products following enzyme hydrolysis of the conjugate, derivatization of the dianions and HPLC florescence detection (Mihke et al., 2013). Cross linking was evaluated using the 2, 4, 6-trinitrobenzene sulfonic acid (TNBS) assay, which is a primary amine-specific spectrophotometric probe. TNBS reacts with primary amines in proteins to produce a complex that absorbs at 420 nm. Loss of TNBS reactivity in dNCO-conjugated proteins occurs only when the dNCO cross-links two amine sites. This method, though not very sensitive, is very specific because only primary amines, the predominant sites found to be conjugated and cross-linked by dNCOs, react with TNBS. Gel electrophoresis was also used to qualitatively evaluate the extent of conjugation and cross-linking in dNCO conjugated proteins. Under denaturing conditions intermolecular cross-linked proteins and highly substituted proteins have a larger molecular size in comparison to un-conjugated protein and these tend to migrate slower. On the other hand, intramolecular cross-linking may prevent complete protein denaturation resulting in the migration similar to that of a smaller molecule that migrates faster. Proteomic mass spectrometry was employed to delineate TDI binding sites on Hb. Acrylonitrile added Hb and trimellitic anhydride (TMA)–adducted Hb were demonstrated to be antigenic (Wong et al., 2004; Pien et al., 1988), so it is crucial to understand the reactivity of Hb to different dNCOs as well as to dNCO specific monoclonal antibodies relative to a well-documented dNCO reactive protein HSA.

2. Materials and methods

2.1. Chemicals

Unless otherwise specified, all reagents were acquired from Sigma-Aldrich (St. Louis, MO, USA) and used without further purification. Dichloromethane (reagent grade) was purchased from J.T. Baker/Avantor Performance Materials (Center Valley, PA, USA). Sodium tetra borate, sodium hydroxide, hydrochloric acid, 98% sulfuric acid, and N-acetyl glycine were purchased from Fisher Scientific (Fair Lawn, NJ, USA).

2.2. Preparation of TDI–HSA/Hb adducts

TDI–protein adducts were prepared as described previously for MDI–HSA/Hb conjugates (Mihke et al., 2013). Briefly, 0.5 mg/ml protein solutions were prepared in 0.01 M PBS (pH 7.4). TDI (42.3 μl) was dissolved in 1 ml dry aceton and diluted ten times to make stock solution for 40:1 TDI:protein. Serial dilutions of TDI in acetone were performed to make stock solutions for 10:1, 5:1, and 1:1 TDI:protein. Fifty microliters of TDI stock solution was added to 5 ml of 0.5 mg/ml protein with mixing, resulting in TDI:protein molar conjugation ratios of 1:1, 5:1, 10:1, and 40:1. Samples were then incubated at room temperature (RT) for 1 h with mixing. Following incubation, samples were dialyzed for 18 h at 4 °C against 4 L of distilled deionized water using 12,000–14,000 MWCO dialysis tubing (Spectrum Laboratories, Inc., Rancho Dominguez, CA) and stored at 4 °C until analysis.

2.3. Preparation of HDI–HSA/Hb adducts

For preparation of HDI–protein adducts, 0.5 mg/ml protein solutions were prepared in 0.01 M PBS (pH 7.4). HDI (47.3 μl) was dissolved in 1 ml dry acetone and diluted ten times to make stock solution for the 40:1 HDI:protein conjugation ratio. Serial dilutions of HDI in acetone were prepared to make solutions for 10:1, 5:1 and 1:1 HDI:protein conjugation ratios. Conjugations, dialysis and sample storage were performed as described for TDI samples.

2.4. Analysis of number of moles of dNCO bound per mole protein

TDI/HDI-conjugated proteins (2 ml aliquots) were hydrolyzed by incubating with 1 ml of 3 M H2SO4 at 100 °C for 16 h. Toluene diamine (TDA) and hexamethylene diamine (HDA) standards (Sigma-Aldrich, St. Louis, MO, USA) were spiked into protein standards (1–16,000 ng/ml) and were run in parallel with conjugates. Following hydrolysis, samples and standards were cooled to RT and 5 ml of saturated sodium hydroxide was added. Samples were vortexed,
and put in an ice bath to cool for 10 min. The resulting TDA and HDA from samples and standards were extracted into 6 ml of dichloromethane and the solvent was subsequently evaporated at 40 °C under N2 to 1 ml. The dichloromethane extracts were then back-extracted into 500 μl of 0.5% H2SO4. Saturated borate buffer (250 μl, pH 8.5) and 450 μl of acetonitrile were added to 250 μl of H2SO4 extract and vortexed for 1 min. Fluorescamine (50 μl of 14.4 mg/ml in acetonitrile) was added. This was vortexed for 1 min, and 100 μl was injected onto a Supelco LC-SI C18 column (25 cm 4.6 mm, 5 μm, Supelco, Bellefonte, PA, USA). Samples and standards were analyzed on a Shimadzu Prominence high-performance liquid chromatography system (HPLC) (Shimadzu, Columbia, MD, USA) consisting of an online vacuum degasser (model DGU-20A5), a quaternary pump (model LC-20AT), an auto sampler (model SIL-10AD-VP), and a fluorescence detector (model RF-10AXL). The HPLC system was controlled by EZ Start software version 7.3 (Lab Alliance, State College, PA, USA). Samples and standards were eluted from the column at 1 ml/min over 20 min using a linear gradient of 10% to 50% acetonitrile/water over 13 min and held at 50% for 5 min. The resulting TDA/HDA–fluorescamine complex was excited at 410 nm, and emission was measured at 510 nm.

2.5. Assessment of cross-linking: TNBS assay

The trinitrobenzene sulfonic acid (TNBS) assay was used to evaluate the extent of cross-linking in TDI–HSA and HDI–HSA conjugates (Snyder and Sobocinski, 1975). TNBS (5%, w/v) was diluted 1:5.48 with 0.1 M borate buffer, pH 9.3. To 500 μl of sample, 12.5 μl of diluted TNBS was added, mixed and incubated for 30 min at RT. Absorbance at 420 nm was measured on a Beckman Coulter spectrophotometer (model DU 800, Beckman Coulter, Somerset, NJ, USA).

2.6. Assessment of cross-linking: Sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE)

For denaturing gels, HSA, Hb, and TDI/HD1/HSA-Hb conjugates were mixed with Laemmli sample buffer containing 2-mercaptoethanol. Samples were run on 8% and 12% polyacrylamide gels. Following electrophoretic separation of proteins, the gels were stained with Imperial™ protein stain (Pierce, Rockford, IL, USA) and destained in water. Unmodified/unconjugated HSA, HB and Bio-Rad pre-stained molecular weight markers (Life Science, Hercules, CA) were used for relative molecular weight determination.

2.7. Trypsin digestion of hemoglobin samples

For identification of TDI conjugation sites on Hb by ultra-performance liquid chromatography quadruple time-of-flight mass spectrometry (UPLC–qTOF MS), 200-μl aliquots of TDI–Hb samples were incubated with tributylphosphine for 30 min at RT to reduce the disulfide bonds, followed by alkylation with iodoacetamide for 1 h at RT. Alkylation was quenched by further addition of tributylphosphine for 15 min at RT. Porcine trypsin in 25 mM NH4HCO3 was then added at a 40:1 (protein/trypsin) ratio. Samples were incubated overnight at 37 °C. The next day, 12 μl of 10% trifluoroacetic acid (TFA) was added to stop trypsin digestion.

2.8. Ultra-performance liquid chromatography (UPLC)

Tryptic peptides of Hb and TDI–Hb were separated on a Waters nanaACQUITY UPLC system (Waters, Milford, MA, USA). Aliquots (1 μl) of the digest mixture were injected and trapped/desalted on a 5-μm Symmetry C18 trapping column (180 μm × 20 mm) with 99.5/0.5 A/B (A: 0.1% formic acid; B: 0.1% formic acid in acetonitrile) at a flow rate of 15 μl/min for 1 min. Separation was performed on a 1.7-μm BEH130 C18 analytical column (100 μm × 100 mm) using gradient elution at a flow rate of 400 nl/min and a gradient of 99:1 to 60:40 A/B over 90 min.

2.9. Tandem Mass Spectrometry (MS/MS) analysis of Hb peptides

The eluent from the UPLC system was directed to the nano-electrospray source of a Waters SYNAPT MS qTOF mass spectrometer. Positive ion nano-electrospray was performed using 10-μm Pico-Tip (Waters) emitters held at a potential of +3.5 kV. The cone voltage was held constant at +40 V for all experiments. Dry nitrogen desolvation gas was supplied to the instrument via a nitrogen generator (NitroFlowLab, Parker Hannifin, Haverhill, MA, USA). [Glu] 1-Fibrinopeptide B (100 fmol/μl in 75:25 A/B) was supplied to an orthogonal reference probe, and the [M + 2 H]2+ ion (m/z 785.84265 u) was measured as an external calibrant at 30-s intervals. Ultra-high purity (UHP) argon was used as collision gas. Spectra were acquired in an “MSe” fashion (Hettick et al., 2012; Hettick and Siegel, 2011). Alternating 1-s mass spectra were acquired. The collision energy was set to 6 eV (1-s low energy scan) and a 15- to 30-eV ramp (1-s high energy scan).

2.10. Data analysis for TDI binding sites on Hb

Data were analyzed with BioPharalynnax version 1.2 (Waters), a software program for analysis of peptide mass maps and identification of sites of modification on known protein sequences. Identification of an isocyanate binding site involved observing a potential peptide–dNCO conjugation product with less than 30 ppm m/Dm mass error in the analyte peptide mass map, comparing analyte and control peptide mass map from unmodified Hb showing that observed m/z and chromatographic retention time are unique to analyze, and observing MS/MS data containing bn- and yn-type ions consistent with the assigned sequence and modifier.

2.11. Immunoassay for conjugates: ELISA for TDI–HSA/Hb

Binding of IgG1 monoclonal antibody (mAb) 60G2 raised against TDI–KLH (Ruwona et al., 2011) to TDI-conjugated HSA and Hb was analyzed using an indirect enzyme-linked immunosorbent assay (ELISA). The development and characterization of the 60G2 mAb has been previously described by Ruwona et al. (2011). Ninety-six-well plates (Corning, NY, USA) were coated with TDI-protein conjugates overnight at 4 °C. After washing three times with PBST (PBS with 0.05% Tween 20), plates were blocked with 3% skim milk/PBST (SMPBST) for 1 h at 37 °C. Plates were then incubated on a shaker for 1 h with 2 μg/ml 60G2 mAb at RT, washed three times with PBST and incubated for 1 h at 37 °C with alkaline phosphatase conjugated AffiniPure goat, antihuman IgG(H + L) (Promega, Madison, WI) diluted 1:5000 (v/v) in SMPBST. Following incubation, plates were washed 3 times with PBST and binding of the 60G2 mAb to the conjugates was visualized using 0.5 mg/ml p-nitrophenyl phosphate (Sigma-Aldrich, CAS Number 4264-83-9) in alkaline phosphatase substrate. The optical density was measured at 405 nm after 30 min using a Molecular Devices SpectraMax M4 Multi-mode Microplate Reader (Sunnyvale, CA, USA).

2.12. Statistical analysis

Data are presented as mean and standard deviation (SD). Analysis of variance (ANOVA) was employed for comparing the effect of dNCO and protein on the extent of conjugation and crosslinking on proteins. Differences were considered significant at a p < 0.05. N = 3/group, where N is number of replicates.

3. Results

3.1. Mapping TDI binding sites on Hb

TDI-Hb conjugates were digested with trypsin, and resultant peptides were analyzed by UPLC–MS/MS to determine TDI binding sites.
Examination of the tandem mass spectra of the tryptic peptides allowed assignment of conjugation sites on Hb as previously described (Mhike et al., 2013). Hb has 2 alpha and 2 beta subunits and mass spectrometry allowed identification of the parent subunit from which each binding site originated. Table 1 shows the concentration-dependent specific binding sites identified for TDI on Hb.

A TDI concentration dependent increase in the number of binding sites was observed and a total of eight binding sites were identiﬁed at the highest concentration of TDI used, including the N-terminal valine on both the alpha and beta chains. TDI bound to three lysines on the alpha chain and three additional lysines on the beta chain. At the lowest TDI concentration used, only the two N-terminal valines of the alpha and beta chains were bound. Increasing TDI concentrations increased the number of sites bound to a maximum of eight at 40:1 TDI:Hb.

3.2. Quantitation of TDI and HDI in Hb and HSA

TDI and HDI-conjugated HSA and Hb were hydrolyzed under strong acidic conditions and the resultant TDA and HDA were derivatized with fluorescamine and quantitated using HPLC. Quantitation of the number of moles of TDI and HDI bound to Hb and HSA is reported in Table 2. On a per mole basis, TDI was more reactive to both Hb and HSA than HDI over the entire concentration ranges used in this study. This agrees with ﬁndings from Wisnewski et al. who found that the rate of HSA carbamoylation from TDI–GSH derived TDI was higher than carbamoylation from HDI–GSH derived HDI (Wisnewski et al., 2013). Table 2 also demonstrates that HSA was more reactive to TDI than Hb. A similar trend was noted for HDI.

3.3. Cross-linking in TDI– and HDI–HSA: TNBS assay

Table 3 shows a concentration-dependent loss of available primary amines with increasing TDI and HDI concentrations and, thus, an increase in the amount of dNCO cross-linking of protein residues (Mhike et al., 2013). At TDI and HDI concentrations ranging from 1:1 to 10:1, the degree of cross linking is not statistically different between the 2 diisocyanates. However at 40:1 dNCO:HSA, TDI has a signiﬁcantly higher degree of cross linking than HDI (P-value <0.01). The TNBS assay could not be used to evaluate cross-linking in Hb conjugates because of spectral interference at 420 nm, the wavelength at which the absorbance of the complex between TNBS and primary amine is measured.

3.4. Qualitative assessment of conjugation and crosslinking in TDI/HDA–HSA and TDI/HDA–Hb: Gel electrophoresis

Polyacrylamide gel electrophoresis was used to evaluate the extent of binding and cross-linking. Intermolecular cross-linked and highly substituted proteins will migrate at a slower rate than the unconjugated protein, whereas extensive intramolecular cross-linking may prevent complete protein denaturation, causing an apparent migration of a molecule smaller than the unconjugated protein. Fig. 1 shows an 8% SDS-PAGE gel of 0.5 mg/ml HSA reacted to TDI. Significant spreading of the HSA band was observed at the 40:1 TDI:HSA conjugation ratio. Fig. 2 is a 12% denaturing gel of 0.5 mg/ml Hb reacted to TDI. Denaturation of Hb resulted in the incomplete dissociation of the alpha and beta subunits that migrated at molecular weights of approximately 14 kDa 28 kDa. Shift in migration due to conjugation to TDI was not observed. In contrast, HDI–HSA and HDI–Hb conjugates produced band spreading and shifts in migration/band spreading for both HSA and Hb conjugates. (Figs. 3 and 4).

3.5. ELISA assessment of TDI–HSA and TDI–Hb

HDI–Hb conjugates reactivity to antibodies was not evaluated due to the lack of an HDI speciﬁc antibody. Although there is cross reactivity between HDI and monoclonal antibodies raised against TDI, the reactivity was too low and close to detection limit to make any quantitative analysis in agreement with HPLC results. Binding of IgG mAb 60G2 to TDI-conjugated HSA and Hb was analyzed using an indirect enzyme-linked immunosorbent assay (ELISA). Fig. 5 shows the immunoassay results of conjugated proteins following titration into the ELISA plate at protein concentrations from 97.66 ng/ml to 25 μg/ml. Immuno reactivity of 60G2 to the conjugated Hb was higher at 40:1 TDI:protein than 10:1 in both HSA and Hb conjugates. 60G2 was more reactive to TDI–HSA than TDI–Hb at both 40:1 and 10:1 TDI:protein.

4. Discussion

Our previous study employed several techniques to evaluate MDI–HSA and MDI–Hb conjugates (Mhike et al., 2013). In the current study, we extended use of this methodology to compare TDI–HSA, TDI–Hb, HDI–HSA, and HDI–Hb conjugates. The objective was to compare the extent of conjugation of TDI and HDI to proteins (HSA and Hb), evaluate differences in the extent of cross-linking using the TNBS assay between TDI and HDI on HSA, and assess reactivity of TDI conjugated HSA and Hb with a monoclonal antibody (IgG 60G2 mAb) that recognizes TDI conjugated proteins. The methodology used herein is relevant for the charac-terization and standardization of dNCO haptenated protein for specific

![Table 1](https://example.com/table1.png)

| Amino acid specific binding sites observed for TDI on hemoglobin. |
|-----------------|-----------------|
| 1:1  | 5:1  | 10:1  | 40:1  |
| Alpha subunit  |       |       |       |       |
| Val  | x     | x     | x     | x     |
| Lys 11 | x    | x     | x     | x     |
| Lys 16 |       | x     |       |       |
| Lys 40 |       |       |       |       |
| Beta subunit  |       |       |       |       |
| Val  | x     | x     | x     | x     |
| Lys 17 |       | x     |       |       |
| Lys 144 |       | x     |       |       |
| Lys 61 |       |       |       | x     |

![Table 2](https://example.com/table2.png)

| Mole of TDI and HDI bound to Hb or HSA quantified using HPLC. |
|-----------------|-----------------|
| dNCO:protein | Average moles of dNCO | Average moles of dNCO |
| dNCO:protein | bound | bound |
| TDI:HSA        |       |       |
| 1:1             | 0.51 ± 0.56 | 0.04 ± 0.41** |
| 5:1             | 2.65 ± 0.23  | 0.44 ± 0.45** |
| 10:1            | 5.06 ± 0.43  | 0.82 ± 0.36** |
| 40:1            | 12.86 ± 0.56 | 2.79 ± 0.40** |
| HDI:HSA        |       |       |
| 1:1             | 0.02 ± 0.21  | 0.02 ± 0.21** |
| 5:1             | 0.57 ± 0.02  | 1.33 ± 0.23** |
| 10:1            | 1.48 ± 0.09  | 0.32 ± 0.60** |
| 40:1            | 4.31 ± 0.06  | 1.03 ± 0.59** |

** Extent of HDI binding to HSA is statistically different than TDI binding.

Comparison of dNCO binding to HSA vs. to Hb.

* P < 0.05.
** P < 0.01.
*** P < 0.001.

![Table 3](https://example.com/table3.png)

<table>
<thead>
<tr>
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</thead>
<tbody>
<tr>
<td>dNCO: protein molar ratio</td>
<td>Loss of TNBS amine reactivity (%)</td>
</tr>
<tr>
<td>TDI:HSA</td>
<td>2.4-TDI:HSA</td>
</tr>
<tr>
<td>Negative control</td>
<td>0.00 ± 0.00</td>
</tr>
<tr>
<td>1:1</td>
<td>25.46 ± 2.34</td>
</tr>
<tr>
<td>5:1</td>
<td>30.24 ± 2.89</td>
</tr>
<tr>
<td>10:1</td>
<td>32.88 ± 3.14</td>
</tr>
<tr>
<td>40:1</td>
<td>48.35 ± 3.76</td>
</tr>
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** Extent of TDI cross-linking statistically different than HDI (P < 0.01).
antibody detection. Although knowledge of specific sights bound at lower conjugation ratios may have value in the development of bio-monitoring of dNCO conjugates in biological fluids, the measurement and characterization of in vivo formed species is beyond the scope of the present work. Differences in reactivity between TDI and HDI conjugated to HSA and Hb were observed using the HPLC quantification of moles dNCO bound per mole protein. HSA was more reactive to both TDI and HDI than Hb. This may be indicative of the structural differences between the two proteins. Hb, with four polypeptide subunits (two alpha and two beta) and an iron-containing porphyrin ring, may mask potential binding sites, thus affecting its reactivity with dNCOs. This contrasts sharply with HSA, a single polypeptide with 17 pairs of disulfide bridges and 1 free cysteine. TDI was more reactive to both HSA and Hb than HDI at pH 7.4. These results agree with earlier findings where HSA was found to be the most modified protein in the blood of dNCO exposed subjects (Budnik et al., 2013).

MS/MS was used to delineate specific TDI binding sites on Hb. A concentration-dependent increase in the number of binding sites was observed across the entire TDI concentration range employed. Only the N terminal valines on both the alpha and beta subunits were observed at 1:1 TDI:Hb and these were conserved at all concentrations studied, suggesting that these sites are the kinetically favored reactive sites. Non-terminal amino acids of the beta subunit were bound by TDI only from 10:1 TDI:Hb concentrations and higher, while non-terminal amino acid binding sites on the alpha subunit were observed at 5:1 TDI:Hb. The non-terminal TDI binding sites observed on Hb were all lysine residues, specifically lysines 11, 16, and 40 of the alpha subunit and lysines 17, 144, and 61 of the beta subunit. Some of the TDI binding sites observed in this study were comparable to the MDI binding sites reported in our previous study (Mhike et al., 2013). In addition to the N-terminal valines of the alpha and beta subunits, lysine 66 was also observed at 1:1 MDI:Hb. Only lysine 40 of the alpha subunit and lysine 61 of the beta subunit were bound by both MDI and TDI. Lysines 11 and 16 of the alpha subunit and lysines 17 and 144 of the beta subunit were only observed in TDI. In contrast, lysine 7 of the alpha subunit and lysines 8, 65 and 66 of the beta subunit were only observed with MDI. The differences in the binding sites between MDI and TDI can give an insight into the possibility for conformational and structural differences in the resultant conjugates, which may potentially affect their antigenicity and immunogenicity.

The TNBS assay, which has traditionally been used to assess chemical adduction with amino groups (Lemus et al., 2001), was employed in this study to evaluate cross-linking in TDI-HSA and HDI-HSA conjugates. A concentration dependent loss of available primary amines with increasing TDI and HDI concentrations was observed, indicating an increase in the amount of dNCO cross-linking of protein residues. At lower TDI and HDI concentrations (1:1–10:1 TDI/HDI:HSA), the degree of cross-linking was similar for both dNCOs. At 40:1 dNCO:HSA, TDI had a higher degree of cross-linking than HDI. A 62% loss of primary amine reactivity was observed at 40:1 TDI:HSA compared with a 48% loss of amine reactivity at 40:1 HDI:HSA (P < 0.01). A similar comparison could not be
made for hemoglobin conjugates due to spectral interference at 420 nm, the wavelength at which the absorbance of the TNBS-amine complex was measured.

The ability of the TDI conjugates to be bound by TDI-specific antibody was evaluated using an indirect ELISA format. The ELISA format employed in the current study used an anti-TDI–protein IgG that was produced in our lab against TDI-KLH (Ruwhona et al., 2010) as the primary antibody and an alkaline phosphatase-labeled anti-IgG as the detection antibody. Both TDI-HSA and TDI-Hb reacted to the anti-TDI–protein monoclonal antibody, indicating that TDI conjugated Hb can be antigenic. TDI-Hb antigenicity with the 60G2 mAb was however 30% lower than that observed for TDI-HSA. This is in agreement with HPLC results where TDI binding to HSA was significantly higher than to Hb suggesting that the absolute number of moles of dNCO bound rather than the specific protein bound is a greater determinant for recognition by the 60G2 mAb. The immunogenicity or antigenicity of in vivo dNCO-added hemoglobin has not yet been reported in the literature, although our data suggest that dNCO haptenated HSA is superior to conjugated Hb for the detection of dNCO specific antibody.

Cross-linking and extent of conjugation was visualized using denaturing gel electrophoresis. Alteration of band migration and band spreading was evident for both HDI and TDI conjugated HSAs at the highest conjugation ratio, however, a clear migration shift/band spreading was only evident in HDI:HSA at the lower binding ratios (Figs. 1 and 3). Hemoglobin subunits did not completely dissociate under denaturing conditions as evidenced by the protein band at approximately 28 kDa. Shift in migration of TDI bound Hb subunits was not observed, and only clearly observed at the highest HDI conjugation ratio of 40:1 (Figs. 2 and 4). These findings are in contrast to what was previously observed for MDI, where clear conjugation-dependent shifts in migration were observed down to a 1:1 conjugation ratio (Mhike et al., 2013). One possible explanation for the differences seen between SDS–Page between that observed for HDI, MDI and TDI bound proteins may be that location of the 2 TDI NCO groups located on the benzene ring are spatially closer to each other than in MDI or HDI which may produce differences in comparative migration of the conjugates in the SDS-Page gels.

Increasing molar ratio for conjugation increased extent of conjugation, degree of cross-linking, gel migration and reactivity with dNCO specific monoclonal antibody binding. It is therefore difficult to dissect the specific influences of intra- and intermolecular cross-linking from total dNCO bound on the overall antigenicity of the resultant conjugates.

The bifunctional electrophilic nature of the isocyanates makes it very difficult to dissect out the components critical to dNCO specific antibody recognition.

Increases in total amount of dNCO bound, intra- and intermolecular crosslinking, and
dNCO self-polymerization on proteins, and as well as recognition by the 60G2 mAb all increase were demonstrated with increasing conjugation ratios of dNCO:protein. The mAb 60G2 is extremely well characterized with respect to binding specificity (30). It recognizes both 2,4-TDI and 2,6-TDI bound HSA, bound mouse serum albumin, and bound keratin. It has slight reactivity toward MDI–HSA, HDI–HSA and HSA conjugated to 2,5- and 3,4-dimethyl phenylisocyanate. It has no reactivity toward phenyl isocyanate, 2-toluene isocyanate, 4-toluene isocyanate or toluene diisothiocyanates. Although, dNCO specific IgE and IgG from dNCO exposed individuals was not tested against the various dNCO conjugated proteins, others have reported that recognition by patient sera antibodies is dependent on immunosassay procedure, conjugation method and predominant exposure dNCO form to that individual (Campo et al., 2007; Ye et al., 2006). Comparison of specific antibody prevalence in dNCO workers is difficult in the absence of detailed dNCO–HSA characterization. Until the relative contribution of the multiple dNCO conjugation products to dNCO immunogenicity and antigenicity can be determined, we believe that dNCO antigen preparations used for standardized screening of workers’ sera or research applications should undergo as complete quantitative chemical characterization as possible similar to that outlined in the present manuscript.

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