Characterization of human thioredoxin-like 2 (Txl-2): a novel microtubule-binding thioredoxin predominantly expressed in the cilia of lung airway epithelium and spermatid manchette and axoneme

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SUMMARY

We describe here the cloning and characterization of a novel member of the thioredoxin family, named thioredoxin-like protein 2. The Txl-2 open reading frame codes for a protein of 330 amino acids consisting of two distinct domains: an N-terminal domain typical of thioredoxins and a C-terminal domain belonging to the NDP-kinase family, separated by a small interface domain. The Txl-2 gene spans approximately 28 kb, is organized into 11 exons and maps at locus 3q22.3-q23. A splicing variant lacking exon 5 (Δ5Txl-2) has also been isolated. By quantitative real-time PCR we demonstrate that Txl-2 mRNA is ubiquitously expressed, with testis and lung having the highest levels of expression. Unexpectedly, light and electron microscopy analyses show that the protein is associated with microtubular structures such as lung airway epithelium cilia and the manchette and axoneme of spermatids. Using in vitro translated proteins, we demonstrate that full-length Txl-2 weakly associates with microtubules. In contrast, Δ5Txl-2 splicing variant specifically binds brain microtubules preparations containing microtubule-binding proteins with very high affinity. Importantly, Δ5Txl-2 also binds to pure microtubules, proving that it possesses intrinsic microtubule-binding capability. Taken together, Δ5Txl-2 is the first thioredoxin reported to bind microtubules and might therefore be a novel regulator of microtubule physiology.
INTRODUCTION

Thioredoxin (Trx) is a small ubiquitous protein (12 kDa) that is conserved in all organisms from lower prokaryotes to human and functions as a general protein disulfide reductase. The redox activity of thioredoxin resides in the sequence of its conserved active site Cys-Gly-Pro-Cys (CGPC) which undergoes reversible oxidation of the two cysteine residues from a dithiol to a disulfide form (1). Thioredoxin is maintained in its active reduced form by the flavoenzyme thioredoxin reductase, a selenocysteine containing protein that uses the reducing power of NADPH (1). Several functions have been assigned to thioredoxin, mostly dependent on its redox activity, including regulation of transcription factor DNA binding activity, antioxidant defense, modulation of apoptosis and immune response (2). Moreover, abnormal thioredoxin expression has been correlated with a number of pathological situations such as cancer, Alzheimer’s and Parkinson’s diseases (3). The three-dimensional structure of thioredoxin is conserved through evolution and consists of five central stranded $\beta$-sheets externally surrounded by four $\alpha$-helices (4). The active site is located in a protrusion of the protein between the $\beta$-2 strand and the $\alpha$-2 helix (4). The conserved active site sequence and the three-dimensional structure of thioredoxin is the hallmark for the thioredoxin family.

During recent years, the number of thioredoxin family members has increased substantially. Based on protein sequence organization, two distinct groups of thioredoxins can be distinguished: Group I includes those proteins that exclusively encode a thioredoxin domain, while Group II is composed of fusion proteins of thioredoxin domains plus additional domains. Among those belonging to Group I are *E. coli* Trx-1 (5), the three yeast thioredoxins (6,7) and mammalian Trx-1 and Trx-2 (8,9). Examples of Group II thioredoxins are *E. coli* Trx-2 (10), *Chlamydomonas* DLC14 and DLC15 proteins (11), mammalian Txl-1 and the spermatid-specific thioredoxins Sptrx-1.
and Sptrx-2 (12-14). Until our discovery of Txl-2, Sptrx-2 was the only mammalian member of the family where two different known protein domains are present in the same polypeptide, as Sptrx-2 is a fusion protein of an N-terminal thioredoxin domain followed by three nucleoside diphosphate (NDP) kinase domains. Similar domain structure is also found in sea urchin axonemal protein IC1 (15).

NDP-kinases (also known as nm23) constitute another well-known family of structurally and functionally conserved proteins identified across a wide range of species from bacteria to human. NDP-kinases catalyse the transfer of \(\gamma\)-phosphates between nucleosides and deoxynucleoside di- and tri-phosphates, playing a pivotal role in maintaining a balanced pool of nucleotides (16,17). In addition to the kinase function, nm23 proteins have been implicated in cell growth, cancer progression and development (17,18). Similar to thioredoxins, humans have several NDP-kinases (termed nm23-H1 to H8, of which nm23-H8 is also known as Sptrx-2). NDP kinases can also be classified into two groups based on sequence alignment and phylogenetic analysis (19). Group I is composed of nm23-H1 to H4 which all share a similar genomic organization consisting of a hexameric three-dimensional structure and the classical enzymatic activity of NDP kinases. Group II encompasses nm23-H5 to H8 genes, which are defined by a more divergent sequence, including a sequence of the active site that is not strictly conserved. Furthermore, nm23-H5, H7 and H8 have been shown to have a tissue-specific distribution mainly in testis/spermatozoa (14,20,21).

We report here the characterization of a novel human fusion protein composed of an N-terminal thioredoxin domain followed by an NDP kinase domain, named Txl-2. Based on the considerations above, Txl-2 should be classified as a member of the Group II of both thioredoxin and nm23 family of proteins. Txl-2 shares clear homology with Sptrx-2 with the difference that the latter has three NDP kinase domains following the
thioredoxin domain. In accordance with nm23 nomenclature, Txl-2 should also be denoted as nm23-H9.
MATERIALS AND METHODS

cDNA cloning of human Txl-2 gene: The Basic Local Alignment Search Tool (BLAST) (22) was used to perform a survey of different databases at the National Center for Biotechnology Information (http://www.ncbi.nlm.nih.gov/) to identify new entries encoding potential novel members of the thioredoxin family. Using the thioredoxin domain of human Sptrx-2 as bait (14), we found the EST entry AI341589 to encode a putative thioredoxin-like sequence. Based on this sequence, the nested forward primers F1 (5´-GGGCAGCAGGAAGAAGGATATTGC-3´) and F2 (5´-CAGGTCAACATCAGCACACAACAGC-3´) were used for 3´-RACE on a human testis cDNA library (Clontech). Based on the sequence obtained, the nested forward primers R1 (5´-CACCAGCCTAGATAGACATCAACA-3´) and R2 (5´-CTCGATCCTCATCTTCTGGAAGAG-3´) were used for 5´-RACE in the same library. The resulting sequences were used to amplify by PCR the full-length and Δ5Txl-2 cDNA of human Txl-2 from the same library. The amplification products were cloned in the pGEM-Teasy vector (Promega) and sequenced in both directions.

Quantitative real-time PCR. Human cDNA MTC panels were purchased from Clontech (Palo Alto, California, USA), oligonucleotide primers were purchased from Cybergene AB (Stockholm, Sweden) and TaqMan probes and GAPDH kits were purchased from PE Applied Biosystems (Warrington, UK). To detect total Txl-2 mRNA, the forward primer was 5´-GGGCAGCAGGAAGAAGGAA-3´ (nucleotides 30-48), the reverse primer was 5´-TTAGTCCTTTGGAACTGAGCATTTC-3´ (nucleotides 115-91) and the Txl-2 TaqMan probe was 5´-CCTGCAGGTCAACATCAGCACCCA-3´ (nucleotides 54-77). To selectively detect only full-length Txl-2, the forward primer was 5´-TCGTCTTGATGTCCTCGAAAGTAC-3´ (nucleotides 234-258), the reverse primer was
5'-CCTCTAACCACAGCCACCAGTT-3' (nucleotides 323-302) and the Txl-2 long TaqMan probe was 5'-GAGCCAACCTTTCTGTGGATTATGCAGGAGGAG-3' (nucleotides 189-212). All TaqMan probes were fluorescein labeled with the reporter dye FAM and quencher dye TAMRA. All primer/probe sets were designed to cross an exon-intron boundary to prevent detection of genomic DNA. 90 µl mastermix containing 400 nM primers, 400 nM TaqMan probe, and 1xTaqMan Universal Mastermix (PE Applied Biosystems) was added to 1.8 µl cDNA before aliquoting in triplicate to a 96-well microtiter plate. The cDNA was amplified using an ABI PRISM 7700 thermocycler (PE Applied Biosystems) under the following conditions: 1 cycle at 50°C for 2 min and 95°C for 10 min, followed by 40 cycles at 95°C for 15 s and 60°C for 1 min. The amount of GAPDH and Txl-2 mRNA was calculated using the Standard Curve Method (separate tubes, following the instructions in User Bulletin 2; PE Applied Biosystems). Fluorescence intensity was measured during the PCR run. A graph was drawn with the threshold cycle CT value versus the logarithm of the amount of serially diluted control cDNA. Using this graph and the CT value of GAPDH and TACC samples, the relative amount of TACC mRNA adjusted for GAPDH was calculated. All real-time PCR experiments were carried out in triplicate and performed a minimum of three independent times with similar results.

To determine the relative amounts of full length and alternatively spliced Δ5Txl-2 in samples, plasmids containing either form were quantified and serially diluted to prepare standard curves for analysis at the same time as sample tissue cDNA. As a secondary control, samples were also prepared containing various mixtures of Txl-2 full length and Δ5Txl-2 plasmids. Each sample was amplified using the Txl-2 primer and probe sets to determine the quantity of either total or full-length Txl-2.
Expression and purification of human Txl-2. The ORF encoding human Txl-2 was cloned into the BamHI-EcoRI sites of the pGEX-4T-1 expression vector (Pharmacia) and used to transform *E. coli* BL21(DE3). A single positive colony was inoculated in 1 liter of LB medium plus ampicillin and grown at 37°C until A<sub>600</sub> = 0.5. The production of the fusion protein was induced by addition of 0.5 mM IPTG and growth was continued for another 3.5 h. Overexpressing cells were harvested by centrifugation and frozen until use. The cell pellet was resuspended in 40 ml 20 mM Tris-HCl, 1 mM EDTA and 150 mM NaCl plus protease inhibitor cocktail at the concentration recommended by manufacturer (Sigma-Aldrich). Lysozyme was added to a final concentration of 0.5 mg/ml with stirring for 30 min on ice. 1% sarkosyl was added and cells disrupted by 10 min sonication and the supernatant was cleared by centrifugation at 15,000 x g for 30 min and loaded onto a glutathione sepharose 4B column (Pharmacia Biotech). Binding to the matrix was allowed to occur for 2 h at room temperature. Thrombin (5U per mg fusion protein) was used to remove GST by incubation overnight at 4°C. The resulting protein preparation was then subjected to ion exchange chromatography using a HiTrap Q column (Pharmacia Biotech) and human Txl-2 was eluted using a gradient of NaCl. For gel filtration chromatography, the Txl-2 preparation from ion exchange chromatography was applied to a Superdex G-75 prep. grade column (Amersham Pharmacia Biotech) under non-denaturing conditions, pre-equilibrated with the same buffer as the protein preparation. Protein concentration was determined from the absorbance at 280 nm using a molar extinction coefficient of 24,310 M<sup>-1</sup> cm<sup>-1</sup>, calculated using the Protean Program included in the DNASTAR Software Package (DNASTAR Inc., Madison, USA). An identical protocol was used to purify recombinant Δ5Trx-2 variant and the protein concentration was determined using the same molar extinction coefficient as that of the full-length form.
**Mass spectrometry analysis.** Reduced Txl-2 was prepared by incubation in the presence of 250 mM DTT on ice for 30 min, and unreduced controls were treated under identical conditions but in the absence of DTT. The samples were diluted ten times with 0.1% (v/v) trifluoroacetic acid and mixed with an equal volume of a saturated solution of sinapinic acid in 33% (v/v) acetonitrile and 0.1% (v/v) trifluoroacetic acid. An aliquot of 0.5 µl of this mixture was crystallized on a microcrystalline layer that had been prepared with a saturated solution of sinapinic acid (Fluka) in ethanol (23). The spectra were acquired using a Reflex III mass spectrometer from Bruker (Germany) and calibrated using the high mass protein standard from Aglient Technologies that contained bovine serum albumin (66430.2 Da), equine cardiac myoglobin (16951.5 Da), and equine cardiac cytochrome c (12359.2 Da). Data processing and evaluation was carried out with the XMASS software from Bruker.

**Antibody production.** Purified GST-hTxl-2 was used to immunize rabbits (Zeneca Research Biochemicals). After six immunizations, serum from rabbits was purified by ammonium sulfate precipitation. Affinity purified antibodies were prepared using a cyanogen bromide-activated Sepharose 4B column, onto which 0.5 mg recombinant Txl-2 fragment had been coupled using the procedure recommended by the manufacturer (Pharmacia). The specificity of the antibodies was tested by western blotting using recombinant and *in vitro* translated human Txl–2 and Δ5Txl-2. Immunodetection was performed with horseradish peroxidase-conjugated donkey anti-rabbit IgG diluted 1:5000 following the ECL protocol (Amersham Corp.).

**Mouse/rat testis and epididymis sample preparation, immunocytochemistry and electron microscopy.** Adult male Sprague-Dawley rats and CD mice were anesthetized and testes and epididymes were fixed by perfusion through the abdominal aorta and
heart, respectively, either with 0.5% glutaraldehyde and 4% paraformaldehyde in 0.1M phosphate buffer containing 50 mM lysine, pH 7.4 or with 4% paraformaldehyde (mice only) or in Bouin’s fixative (for light microscopy). Tissues destined for Lowicryl embedding (for EM) were immersed in the respective fixatives for 2h at 4°C, washed three times in phosphate buffer and incubated with phosphate buffer containing 50 mM NH₄Cl for 1 h at 4°C. Tissues were subsequently washed in buffer, dehydrated in graded methanol up to 90% and infiltrated and embedded in Lowicryl K4M. Thin sections were mounted on Formvar nickel coated grids for immunogold labeling. Bouin’s fixed rat and human tissue were washed extensively in 75% alcohol before being completely dehydrated in ethanol and embedded in paraffin. Paraformaldehyde fixed tissue were washed in phosphate buffer, dehydrated and embedded in paraffin.

For light microscopic immunocytochemistry 5 µm paraffin sections were deparaffinized and hydrated through a graded series of ethanol concentrations before immunoperoxidase localization with anti-Txl-2 antibody by standard procedures (24). For electron microscopic immunocytochemistry ultrathin Lowicryl sections on Formvar coated nickel grids were immunogold labeled according to the procedure of Oko et al. (1996) (25). Staging of the cycle of the seminiferous epithelium and determining the steps of spermiogenesis were done according to the classifications of Leblond and Clermont (26).

**Mouse lung sample preparation, immunocytochemistry and electron microscopy.**

Adult C57Bl6 mice were were killed through cervical dislocation, the trachea was cannulated and the cannula was tied firmly in place. The anterior chest wall was removed and the lungs dissected out. The lungs were infused via the tracheal cannula with 4% paraformaldehyde, pH 7.4 at 20 cm H₂O pressure and maintained at this pressure for five minutes. The lungs were subsequently kept in fixative overnight at 4°C. After fixation the
lungs were dehydrated through a graded series of ethanol. Finally the lobes were separated and placed into individual cassettes and embedded in paraffin. The central portions of the blocks were sectioned at 5 µm intervals and the sections were mounted on glass slides, deparaffinized and hydrated before immunoperoxidase localization with anti-Txl-2 antibody. To block unspecific binding of secondary antibodies, sections were incubated in blocking solution (5% normal goat serum). Primary antibody was added at a 1:20 dilution in blocking solution and in control experiments 0.03 mg/ml blocking peptide was included. After several washes, the Vectastain-ABC kit (Vector Laboratories, OH) was used for visualization. Sections were slightly counterstained with Mayer’s hematoxylin, dehydrated and mounted.

Indirect immunogold labeling was used at the electron microscope level to specifically localize Txl-2. Small pieces (1 mm³) of adult mouse lung were fixed in 4% paraformaldehyde in 0.1 M Sorensens buffer (pH 7.4) for 4 h at 4°C. After washing in several changes of Sorensens buffer at 4°C, the tissues were dehydrated in a graded series of methanol at progressively lower temperatures to -20°C. The tissue pieces were then infiltrated and embedded in Lowicryl K4M (SPI Supplies, West Chester, PA) at -35°C. Lowicryl blocks were polymerized by ultraviolet illumination for 24 h at -35°C and an additional 48 h at -10°C.

Lowicryl thin sections were placed on formvar-coated nickel grids and incubated face-down on drops of blocking solution consisting of 1% BSA in 50 mM Tris (pH 7.6) with 100 mM sodium chloride. After 30 min, the grids were transferred to drops of primary antibody diluted 1:10 in blocking solution and left overnight in a humidity chamber at 4°C. The following day, the grids were washed face-down on wells containing 50 mM Tris (pH 7.6) with 100 mM sodium chloride for 3 x 10 min. The grids were then incubated on blocking solution for an additional 30 min prior to being transferred to drops of secondary antibody (goat F(ab’2) anti-rabbit IgG conjugated to 10 nm colloidal
gold; Ted Pella) diluted 1:30 in blocking solution. After 1 h, the grids were washed as previously described and then rinsed in distilled water. Immunolabeled sections were counterstained with methanolic uranyl acetate followed by lead citrate.

**Microtubule binding assay**

MTs were purified from rat and mouse brain essentially as described previously (27), resulting in MT preparations that contain microtubules and MT-binding proteins including MAP and kinesins. In short: brain samples were extracted in ice-cold BRB80 buffer containing protease inhibitors pepstatin, leupeptin and PMSF and centrifuged at 55,000 rpm for 15 min at 4 °C. Supernatants containing depolymerized MT were removed and stored on ice. MT extracts were next supplemented with 0.5 mM GTP, 15 U/ml hexokinase and 20 mM D-glucose to deplete ATP. Next the extracts were warmed to 30 °C and 5 µM taxol was added. After 5 min incubation at 30 °C, 15 µM taxol was added (20 µM total). AMP-PNP (2 mM) was added to stabilize kinesin heavy chains on MTs. The extracts were layered onto sucrose cushions and spun at 40,000 rpm for 20 min at 22 °C. Cushion was washed 2x with BRB80 before being removed to avoid contamination of the MT pellet. MT pellets were resuspended in 100 µl BRB80 and depolymerized on ice for 20 min. GTP (0.5 mM) and AMP-PNP (2 mM) were added to the MT preparations. Extracts were warmed to 30 °C and polymerized and centrifuged as above. MT pellets were subjected to another round of depolymerization and polymerization as above. Final pellets of polymerized MT were resuspended in 10% glycerol/BRB80 and flash frozen in liquid nitrogen.

Pure polymerised microtubules were made from purified tubulin (CytoSkeleton Inc.) as follows: 10 µl pure tubulin (10 mg/ml) were mixed in BRB80 buffer with 5 µl glycerol and 3.3 µl Taxol (0.15 mM) and water was added to 50 µl. This reaction was incubated at 33 °C for 15 min. The mix was loaded on a glycerol cushion of equal volume,
and spun at 22,000 rpm for 30 min at room temperature. The MT pellet was recovered and dissolved in 50 µl BRB80 buffer/10 % glycerol.

To analyze binding of Txl-2 and Δ5Txl-2 to MT, both forms were transcribed and translated in vitro in the presence of ³⁵S-cysteine using the TNT reticulocyte transcription and translation system (Promega). Radiolabeled proteins were incubated with polymerized brain MT or with pure MT at 30 °C for 15 min in the presence of AMP-PNP (2 mM), GTP (0.5 mM) and Taxol (20 µM). MT were pelleted at 30,000 rpm for 15 min at 22 °C. Supernatants were saved for SDS-PAGE analysis. Two subsequent pelleting reactions were performed. Aliquots of both supernatants and final pellets were boiled in SDS sample buffer and analyzed by electrophoresis on 10 % SDS-PAGE gels, the gels dried and exposed to KODAK BIOMAX film. Intensity of bands was quantitated by image analysis. In indicated experiments MTs were preincubated with anti-Txl-2 antibodies or with human recombinant Txl-2.
cDNA cloning, sequence analysis, genomic organization and chromosomal localization of human Txl-2 gene. By sequence comparison with human Trx-1 (8), we found that Genbank EST entry AI341589 (from pooled germ cell tumors) encoded a putative novel human thioredoxin-like sequence. Therefore, we designed specific primers based on AI341589 sequence and performed 5’- and 3’-RACE PCR analysis using a human testis cDNA library to clone the full-length cDNA sequence of this novel protein. The complete sequence of the cDNA obtained consists of an ORF of 990 bp, a 5’-UTR of 27 bp and a 3’-UTR of 432 bp upstream of the poly(A)’ tail (Fig. 1A). In addition, the PCR also rendered a smaller band, which corresponded to a putative splicing variant (see below). Human Txl-2 ORF encodes a protein of 330 amino acids with a calculated molecular mass of 36.9 kDa and a pl of 4.73. Analysis of the human Txl-2 sequence identified two distinct domains: an N-terminal domain (comprising the first 105 residues) similar to thioredoxins and a C-terminal domain composed of one NDP kinase domain (Fig. 1B). Both domains are separated by a small interface domain. The Txl-2 protein domain organization resembles that of Sptrx-2 (14), with the exception that Sptrx-2 has three NDP kinase domains. Regarding the N-terminal thioredoxin domain, some of the structural amino acids that are conserved in previously characterized mammalian thioredoxins (including Sptrx-2) such as Asp-26, Trp-31, Pro-75 or Gly-91 (numbers refer to those of human Trx-1) are also conserved in Txl-2. However, other residues shown to be essential for catalysis, maintenance of three-dimensional structure or protein-protein interactions are substituted, for instance Phe-11, Ala-29, Pro-40, Asp-58 or Lys-81 (4) (see figure 2B of (14)).

Additionally, the Txl-2 C-terminal domain consists of one NDP kinase domain. Following the nomenclature for NDP kinase protein family, Txl-2 should be considered
as the ninth member of this family and therefore termed nm23-H9 (19). A protein alignment and phylogenetic analysis of all human NDP kinase domains shows that Txl-2 belongs to NDP kinase group II (Figures 2A and 2B) (19). The protein alignment of Txl-2 thioredoxin domain with all the previously identified thioredoxin proteins has been reported elsewhere (14). The 3′-UTR of human Txl-2 mRNA contains a short interspersed repetitive element (SINE) of the Alu family spanning from bases 1156 to 1439 (Fig. 1A). This is not the first report of the presence of such a repetition in a member of the NDP kinase family as nm23-H6 mRNA also harbors an ALU sequence located in the 3′-UTR of its mRNA (21). Interestingly, both proteins are testis-specific. In X-ray crystallography and site-directed mutagenesis studies, nine residues essential for catalysis and stability of nm23 proteins have been identified (17,19). Human Txl-2 has five of these nine conserved residues regarded as crucial for enzymatic activity (Figure 2A). Furthermore, the sequence of the active site (NAVH) matches the consensus for this family of proteins (NXXH) where X can be any residue (Figure 1A and 1B).

A comparison of the protein sequence with PROSITE database (28) identified, along with the above-mentioned thioredoxin and NDP kinase domains, several potential phosphorylation sites and an RCC1 (regulator of chromosome condensation-1) signature present between residues 222-232 (Figure 1A).

A homology search in the Human Genome Sequence Database (http://www.ncbi.nlm.nih.gov/genome/guide/human/) identified the Txl-2 genomic region to be localized to chromosome 3q22.3-q23 (entry NT_025664.5|Hs3_25820) (Figure 3). Using the Genomatix Software (http://www.genomatix.de/) we have determined that Txl-2 gene spans approximately 28 kb and is organized into 11 exons and 10 introns, all conforming to the GT/AG rule (Supplemental data 1). Analysis of the splicing variant that appeared during the cloning PCR indicated that it lacked exon 5.
Differential expression pattern of Txl-2 mRNA in various human tissues. Initially, we used multiple-tissue Northern blots to determine the size and tissue distribution of human Txl-2 mRNAs using either the ORF or the thioredoxin domain as probes. However, we were unable to detect any signal despite the use of low stringency conditions and extended time of exposure (data not shown). To improve the sensitivity of our studies and to provide a means of quantification, we therefore used real-time PCR to determine Txl-2 mRNA levels in a variety of human adult tissues (Fig. 4A). Our results demonstrate that total Txl-2 mRNA in adult tissues is very low. However, highest levels are found in testis and lung while lower levels are found in brain, thymus, spleen, prostate, kidney and ovary. Txl-2 mRNA is virtually absent in colon, liver and heart. We also determined the relative amounts of full-length Txl-2 versus alternatively spliced Δ5Txl-2 in the three tissues with highest mRNA content: testis, lung and brain. As shown in Figure 4B, Δ5Txl-2 is the predominant mRNA form in all three cases. However, while in testis and lung the ratio Δ5Txl-2/Txl-2 is approximately 60:40, in brain this ratio is clearly shifted to a much higher relative amount of the spliced variant, 90:10.

Expression and enzymatic activity of human Txl-2 protein. Recombinant human Txl-2 migrated at 36 kDa in good agreement with its theoretical size (Figure 5A, inset). Members of the NDP kinase family have been described to have an oligomeric structure in their native conformations (19). To evaluate whether this was also the case for Txl-2, we performed gel filtration chromatography and found two peaks corresponding to the monomeric and the dimeric conformation indicating that, in its native form, Txl-2 might be found in equilibrium between both forms (Figure 5A). Analysis of the fractions eluting from the column by SDS-PAGE ruled out the possibility of protein contamination or degradation to explain the appearance of two peaks (data not shown). Next, we used MALDI-TOF mass-spectrometry analysis to determine whether the dimeric conformation
is maintained by disulfide bonding. Figure 5B shows Txl-2 spectra in which two peaks can be identified, corresponding to the monomeric and dimeric form, respectively. Incubation of the Txl-2 protein with DTT buffer resulted in a decrease of the dimeric peak, further demonstrating that disulfide bonds are responsible, at least in part, for the dimeric conformation (Fig. 5A).

Enzymatic activity of thioredoxin is usually assayed by the capacity to reduce the disulfide bonds of insulin using either DTT as artificial reductant or NADPH and thioredoxin reductase as a more physiological reducing system (1). We were unable to detect any enzymatic activity in either enzymatic assay, using full length Txl-2 or the Δ5Txl-2 variant (data not shown). We also determined whether Txl-2 behaves as dominant negative, competing with Trx-1 for binding to thioredoxin reductase. As shown in Figure 5C, increasing amounts of full length Txl-2 weakly, but consistently, compete with human Trx-1 in the thioredoxin reductase enzymatic assay, thus suggesting that the absence of enzymatic activity of Txl-2 is not due to a lack of interaction with the reductase.

On the other hand, NDP kinases catalyze the transfer of a terminal phosphate residue from NTPs to NDPs according to a “ping-pong” mechanism. The first step of this reaction consists of the autophosphorylation of the enzyme at a conserved histidine of the active site (29,30). Txl-2 was unable to undergo autophosphorylation under the same experimental conditions that allowed positive control yeast NDP kinase autophosphorylation (Anna Karlsson, personal communication) (31).

**Cellular and subcellular localization of Txl-2 protein.** As determined by real-time PCR, testis and lung were the tissues with the highest amount of Txl-2 mRNA. We therefore selected these two tissues to address the cellular and subcellular localization of Txl-2 protein. Thioredoxins and NDP kinases display a high degree of amino acid
identity between humans and rodents (LocusLink, http://www.ncbi.nlm.gov/LocusLink/). Due to sample availability and considering the identity between humans and rodents, we decided to perform immunohistochemical analysis on mouse and rat samples. As shown in Figure 6, Txl-2 labeling is readily detected in the cilia of the mouse lung airway epithelium. Preincubation of the antibodies with human recombinant Txl-2 abolished the signal. Similarly, Figure 7 demonstrates the presence of Txl-2 in rat testis seminiferous tubules, where the protein is associated with the spermatid tail and manchette, a microtubular structure assumed to participate in the elongation of the spermatid head and storage of proteins for later delivery to the developing tail (32). Again, antibody preadsorbed with the human recombinant protein gave no labeling thus confirming the specificity of the antibodies and their cross-reactivity in rodent samples. Surprisingly, a strong and specific labeling in the testis blood vessels was also obtained. Identical results were obtained with mouse testis sections (data not shown).

The finding of Txl-2 protein in such specific cellular structures, prompted us to investigate in more detail its subcellular localization, because the possibility existed that Txl-2 is somehow associated with microtubules. Surprisingly, by immunogold electron microscopy, we found Txl-2 in association with microtubules both in lung and testis: Txl-2 signal was over MT that make up lung cilia as well as over the MT-containing spermatid manchette and axoneme (Figures 8 and 9). In addition, in testis blood vessels Txl-2 colocalized with fibrillar components of smooth muscle tissue (Figure 10).

**Microtubule binding activity of human Txl-2.** The immunolocalization data strongly suggested that Txl-2 might be able to bind microtubules. This would constitute the first example of such a characteristic for thioredoxins. To prove this possibility, we performed in vitro microtubule binding assays using $^{35}$S-labeled in vitro translated
proteins. First, translated Txl-2 and Δ5Txl-2 proteins were analyzed for binding to brain MT containing MBPs. As shown in Figure 11A, upper panel, full-length Txl-2 binds only weakly to microtubules in this assay as demonstrated by the presence of recombinant protein in both the microtubule pellet (p; lane a, upper panel) as well as in supernatant 2 (s2; lane c, upper panel), a consequence of release of Txl-2 from microtubules during the course of the experiment. Preincubation of the in vitro translated Txl-2 either with specific antibodies or unlabeled recombinant protein did not result in a significant decrease of the binding (lanes d and g; upper panel, respectively). The appearance of two bands when the Txl-2 construct is translated in vitro is probably a consequence of the use of an internal methionine as translational start in the in vitro system (Figure 1A). In agreement with the immuno electron microscopy data which suggested that Δ5Txl-2 protein can colocalize with microtubules, Δ5Txl-2 protein is clearly present in the pellet fraction and not in the supernatant 2 demonstrating that Δ5Txl-2 is stably bound to microtubules in this assay (Figure 11A lower panel). Moreover, the binding is specific as preincubation of Δ5Txl-2 with unlabeled recombinant protein or anti-Txl-2 antibodies decreases binding to 49% and 28%, respectively, of the levels achieved with the Δ5Txl-2 protein alone (lanes d and g, lower panel, respectively). Since brain MT preparations contain besides microtubules, also microtubule-binding proteins such as MAP, kinesins, etc, these experiments demonstrate that Txl-2 associates with MT, but cannot distinguish direct from indirect MT binding. To distinguish these two mechanisms we compared MT-binding for both Txl-2 and Δ5Txl-2 using pure microtubules and brain microtubules. The results are shown in Figure 11B. The data show that Δ5Txl-2 binds with high affinity to pure microtubules (lane i) with very little material in the second supernatant (lane h). There is no significant difference in Δ5Txl-2 MT-binding using pure or brain MT. This proves that Δ5Txl-2 MT-binding is direct and establishes Δ5Txl-2 as the first genuine
microtubule-binding thioredoxin. As with brain MT, full length Txl-2 associates significantly more weakly with pure or brain MTs (lanes c and f, respectively).
DISCUSSION

Microtubules are fibrous cytoskeletal components of the eukaryotic cell cytoplasm that serve to perform a wide variety of functions, such as cell motility and division, organelle transport and morphogenesis. In addition, microtubules are the main components of the complex and highly organized axonemal structures found in cilia and flagella (33). There is evidence that some members of the NDP-kinase family associate with microtubular structures (34-38). In contrast, only four proteins containing a thioredoxin domain have been reported to be associated to microtubular structures, and in all cases in lower eukaryotes: LC14 and LC15 proteins from Chlamydomonas flagellum (11) and the dynein intermediate filaments IC1 from sea urchin sperm axoneme and IC3 of the ascidian, Ciona intestinalis (15,39). The last two proteins are composed of an N-terminal thioredoxin domain followed by three NDP kinase domains. Similar domain structure has been found in Sptrx-2 (14), a human protein exclusively expressed in the spermatozoon fibrous sheath, a structure that surrounds the axoneme and outer dense fibers of sperm tail principal piece (unpublished results). However, Sptrx-2 is not likely to interact with sperm tail microtubules despite the clear domain homology with the IC proteins as they are kept apart from each other by the outer dense fibers.

We report here a novel protein (Txl-2 and Δ5Txl-2, a splicing variant lacking exon 5) consisting of an N-terminal thioredoxin domain followed by one NDP-kinase domain in close similarity to Sptrx-2 (14). Indeed, a phylogenetic analysis of the separate domains places both proteins in the same tree branch. However, a comparison of the expression pattern of both proteins shows important differences. While Sptrx-2 is a testis-specific protein (14), Txl-2 is more ubiquitous and has lower overall expression levels, although it is more abundant in testis and lung. Unexpectedly, an immunohistochemical analysis of Txl-2/Δ5Txl-2 expression in these two tissues shows the protein to be in close association
with microtubular structures: cilia of the lung airway epithelium and the manchette and axoneme of the spermatids. Furthermore, Txl-2/ ΔTxl-2 is found within the fibrillar components of the testis blood vessel smooth muscle.

The electron microscopic study suggested that Txl-2 might represent the first MT-binding thioredoxin family member, which would have a considerable impact on its potential functions. This possibility was approached by in vitro MT-binding assays. These experiments, using pure microtubules, established definitively that ΔTxl-2 could bind directly to microtubules, without the aid of microtubule-associated proteins such as kinesins or MAPs. In contrast the full length Txl-2 protein has a much reduced affinity for MT.

At this stage we do not know if Txl-2, ΔTxl-2 or both associate with MT in lung and spermatids, because the polyclonal antibodies raised against full-length Txl-2 recognize both forms. In light of the quantitative RT PCR data and the in vitro MT-binding experiments, it appears most likely however that ΔTxl-2 rather than Txl-2 is the major MT-binding form. The difference in affinity may be explained in two ways. First, the MT-interacting area in ΔTxl-2 may be disrupted by the presence of exon 5 in the Txl-2 form, or exon 5 sequences cause a change in Txl-2 conformation affecting MT-binding. Further in vitro analysis of ΔTxl-2 deletion mutants can address this point. Second, cell or tissue-specific factors might be required to modulate the MT binding activity of ΔTxl-2, which are not present in the in vitro MT binding assay. Analysis of brain may help resolve this possibility, since in that tissue ΔTxl-2 is by far the major form. Be that as it may, our results demonstrate for the first time that genuine microtubule-binding members of the thioredoxin family exist.

At present we can only speculate on the mechanism of binding of ΔTxl-2 to microtubules and its role in microtubule physiology. Cysteine residues in tubulin are actively involved in regulating ligand interactions and microtubule formation both in
vivo and in vitro (40). As thioredoxins are considered as general protein disulfide reductases, it is conceivable that the cysteine residues at the active site of Δ5Txl-2 interact with tubulin sulfhydryl groups therefore playing a critical role as a regulator of microtubule stability and maintenance. Moreover, as Δ5Txl-2 has four additional structural cysteines it is plausible that disulfide bonding involving any of these residues could play a role in the binding mechanism. Studies are in progress to analyze the kinetics of Δ5Txl-2 binding to tubulin.

Txl-2/ Δ5Txl-2 has an RCC1 signature in its C-terminal NDP-kinase domain, a motif that is not present in any other member of the nm23 or thioredoxin family. RCC1 is a chromatin-bound guanine-nucleotide exchange factor for the small nuclear GTPase Ran, a Ras-related protein (41). RCC1 has been implicated in nuclear-cytoplasmic transport, mitotic spindle nucleation and nuclear membrane formation (42). Interestingly, both, a somatic and a testis-specific form of Ran have been described in mammals (43,44). The testis-specific variant of Ran is found in the spermatid manchette, a location where RCC1 has also been found (42,44). The spermatid manchette is a transient microtubular structure that develops during spermiogenesis and caudally surrounds the spermatid nucleus (45). The manchette has been recently proposed to be a transient storage location for both signalling and structural proteins involved in nucleocytoplasmatic trafficking or eventually sorted to the centrosome and the developing spermatid tail (32,46). This sorting mechanism has been named intramanchette transport (IMT), in close reference to the so-called intraflagellar transport (IFT) (46,47). Both IMT and IFT involve molecular motors (primarily kinesis and dyneins) mobilizing a multicomplex protein raft to which cargo proteins or precursors for the assembly of the axoneme of a flagellum or a cilium are bound and transported to the tip of the axoneme (46). Thus, the potential association of Txl-2/ Δ5Txl-2 with Ran by its RCC1 motif may be an indirect mechanism of binding to microtubules.
As Txl-2 is also expressed in lung cilia, the potential interacting proteins or substrates are not likely to be testis-specific but rather common to the microtubular infrastructure. The fact that Txl-2 contains thioredoxin and NDP-kinase domains, both thought to be catalytic domains, in addition to its low abundance suggests that Txl-2 is more likely to act as a signalling/enzymatic protein rather than to have a structural role. The acquisition of a thioredoxin domain in flagellar proteins occurs early in evolution and is most likely a consequence of molecular or enzymatic requirements for a specific function in flagellum movement (11). In this context, it has recently been reported that the two thioredoxin domain-containing proteins from *Chlamydomonas* flagella, DLC14 and DLC15 (11) might regulate, by a redox mechanism, the ATPase activity of the flagella outer dynein arm (48). In addition, *Chlamydomonas* flagella p72 and sea urchin sperm axoneme IC1 proteins have been reported to have NDP-kinase activity and suggested to be suppliers of the GTP for microtubule assembly (49). Interestingly, an exhaustive proteomic analysis of human ciliary proteins has found nm23-H5, nm23-H7 and Txl-2 as the only members of the NDP-kinase family expressed in human axonemal extracts, adding further support to our data here reported on axonemal localization of Txl-2 (50). Based on this, a model could be proposed in which Txl-2 is responsible for GTP generation in cilia and flagella axoneme in these organisms. Furthermore, the presence of an RCC1 signature in Txl-2 suggests that it could interact with Ran and supply the GTP required for its enzymatic activity. Thus, it may be anticipated that disruption of a Ran-RCC1-Txl-2 pathway might lead to alteration in the normal development of spermatozoa and other phenotypes associated with defective cilia physiology. Txl-2 expressed in bacteria is inactive in both the thioredoxin assay and the autophosphorylation assay. A similar situation was found for Txl-2’s closest homologue Sptrx-2 as well as other thioredoxins and NDP-kinases (12,14,20). Taken together, the lack of enzymatic activity of bacterially produced Txl-2, despite its interaction with thioredoxin reductase, indicates
that posttranslational modifications that might expose the active site upon conformational changes as well as interaction with other proteins or additional cofactors (cell or tissue-specific) might be required for Txl-2 to function. Alternatively, since Txl-2 binds to microtubules, the weak interaction detected \textit{in vitro} with thioredoxin reductase may become physiologically significant \textit{in vivo}.

The prominent expression of Txl-2 in microtubular structures in lung and testis may make it as a candidate gene for diseases such as primary ciliary dyskinesia (PCD), an an autosomal recessive disorder (OMIM 242650) characterized by a failure of proper ciliary and flagellar movement whose clinical manifestations are chronic respiratory infections, male infertility and situs inversus (51,52). The motility of cilia and flagella is generated in the axoneme, which has been estimated to be composed of more than 250 polypeptides (53). The axoneme consists of a core of 9 peripheral + 2 central microtubule doublets connected by outer and inner dynein arms (composed of heavy, intermediate and light chains) plus other accessory proteins. Electron microscopy studies of sperm of patients affected by PCD reveal multiple phenotypes with anomalies in both microtubule and dynein arm organization, indicating that PCD is a genetically highly heterogeneous disease (51). To determine whether any breakpoint in the region where the human Txl-2 gene maps has been reported to be associated with any of the PCD phenotypes, we screened the Mendelian Cytogenetics Network Database (http://www.wjc.ku.dk/databases) and found one translocation [46,XY, t(3,8) (q23;p23)] with an associated trait of azoospermia/oligospermia. Interestingly, two more breakpoints in the Txl-2 region are associated with hydrocephalus, a phenotype that is assumed to be a consequence of malfunctioning of the cilia of ependymal cells that facilitate circulation of the cerebral spinal fluid (54).

In summary, we have identified and characterized a new protein composed of thioredoxin and NDP-kinase domains that directly binds to microtubules. Its axonemal
localization in sperm flagella and lung cilia indicates that it is a component of the axonemal machinery taking part in regulation of ciliary and flagellar movement and that Txl-2 is a potential susceptibility candidate gene involved in the PCD phenotype.
ACKNOWLEDGEMENTS

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REFERENCES

FIGURE LEGENDS

Figure 1. A) Nucleotide and amino acid sequence of human Txl-2 gene and protein. Numbers on the left indicate amino acids and on the right nucleotides. Thioredoxin and NDP-kinase active sites are boxed. The RCC1 signature is double underlined and the potential polyadenylation signal in the 3’-UTR is underlined. Exon 5 (absent in Δ5Txl-2 variant) within the open reading frame and the SINE repeat within the 3’-UTR are shadowed. B) Domain organization of Txl-2 and Δ5Txl-2 proteins. Numbers indicate the amino acids flanking the thioredoxin, interface and NDP-kinase domains.

Figure 2. A) Primary sequence comparison between the NDP kinase domains of the human Nm23/NDP kinase proteins. The numbering at the top refers to the Nm23-H1 sequence while the numbers on the left indicate the amino acid residue for each respective full-length protein. Identical (black boxes) and conserved (grey boxes, according to (55)) residues within all sequences are indicated. The asterisk denotes residues involved in catalysis and stability (19). Note the amino acid insertion in Txl-2 where the RCC1 signature (MCSGPSHLLIL) is located. B) Phylogenetic tree of human NDP-kinases. The scale indicates the number of substitution events per hundred bases. The percentage bootstrap values (based on 1000 replications) are given on the nodes of the tree.

Figure 3. Chromosomal localization and genomic organization of human Txl-2 gene. The human Txl-2 gene is located between the markers D3S1576 and D3S3586 at 144.46-145.33 cM from the top of the linkage group of human chromosome 3 (based on deCODE high-resolution recombination map of human genome (56)). By comparing this location with other genes in the region we have mapped Txl-2 gene to 3q22.3-q23 flanked
by the genes \textit{DKFZP434A043} and \textit{MRAS}. Human Txl-2 gene is organized in 11 exons, and exon 5 (black) is absent in the \( \Delta 5 \text{Txl-2} \) variant.

**Figure 4. Relative expression of Txl-2 mRNA in human tissues.** Real-time PCR was used to quantify Txl-2 mRNA using the Standard Curve Method. Values were normalized to GAPDH. (A) Bars represent the mean amount of total Txl-2 mRNA for each tissue as determined in triplicate samples. (B) Real-time PCR was used to determine the relative amounts of full-length and alternatively spliced \( \Delta 5 \text{Txl-2} \) in human testis, lung and brain. All experiments were performed in triplicate and similar results were obtained in at least 3 separate experiments. The value for testis was set to 100.

**Figure 5. A) Gel filtration chromatography of human Txl-2.** Elution profile of human Txl-2 applied to a Superdex 75 column in a volume of 5 ml, at a final concentration of 2 mg/ml and a flow rate of 5 ml/cm\(^2\)/h. The inset shows the SDS-PAGE on the fractions eluted from the Superdex 75 column demonstrating the integrity of Txl-2. **B) MALDI-TOF analysis of human Txl-2.** The spectrum shows the single and double charged ions as well as the dimeric form of Txl-2. A decrease in the dimeric form is obtained upon incubation with DTT. **C) Dominant negative activity of human Txl-2.** The dominant negative effect of full length Txl-2 on Trx-1 enzymatic activity was carried out in the presence of NADPH and thioredoxin reductase. The assay conditions were identical to those previously described (13) except that increasing amounts of Txl-2 protein were added to the mix prior to initiating the reaction. Lysozyme was used as a control at the highest molar ratio. The reaction was initiated by adding 5 \( \mu \)l calf thymus thioredoxin reductase (50 \( A_{412} \) units) and stopped after 20 min by the addition of 6 M HCl and 1 mM DTNB. The experiment was repeated three times.
Figure 6. Immunohistochemical localization of Txl-2 protein in lung. Paraffin sections of mouse lung, immunoperoxidase stained with affinity purified anti-Txl-2 antibodies diluted 1/20. A) Large intrapulmonary airway. B) Small intrapulmonary airway. Both locations exhibit clear immunostaining specifically localized to the cilia of ciliated epithelial cells (arrows). No staining is observed in non-ciliated epithelial cells (arrow-heads). C) Immunoreactivity is substantially reduced when Txl-2 blocking peptide is included with the primary antibody. Bar, 10 µm.

Figure 7. Immunohistochemical localization of Txl-2 protein in testis. Paraffin sections of rat seminiferous tubules, immunoperoxidase stained with affinity purified anti-Txl-2 antibodies diluted 1/100. A) Survey section showing obvious immunostaining of the arterioles (Ar). B) No immunoreactivity is detectable when anti-Txl-2 is preincubated with human recombinant Txl-2. C) At higher magnification, during the spermatid elongation phase (steps 8-14), immunostaining was apparent at the periphery of the spermatid nucleus and beyond (arrows), suggesting labelling of the microtubular manchette. D) At this magnification it is clear that the smooth muscle (Sm) wall of the arteriole is immunoreactive. E) Fainter immunostaining of the sperm tails (arrows) in the lumen is also apparent. Bar, 10 µm.

Figure 8. Immunogold electron microscopy localization of Txl-2 protein in lung. A) Electron micrographs (EM) of sections through mouse lung ciliated epithelium immunogold labelled with anti-Txl-2. Labelling is found in close association with the microtubules of the axoneme. B) Mouse cilia labelled with anti-Txl-2 preincubated with human recombinant Txl-2. No immunogold labelling of the microtubules of the axoneme is evident. Bar 0.2 µm.
Figure 9. Immunogold electron microscopy localization of Txl-2 protein in testis. Electron micrographs (EM) of sections through spermatid heads and tails immunogold labelled with anti-Txl-2. A) Elongating spermatid in step 10 showing labelling of the microtubular manchette (M). A, acrosome; Nr, nuclear ring of the manchette; N, nucleus. B) Spermatid in step 12 showing labelling of the manchette (M) and of the axoneme (Ax) C) Spermatid tails in step 15 showing labelling of the axoneme. R and L, ribs and longitudinal columns of the fibrous sheath, An, anlagen of the fibrous sheath, ODF, outer dense fibers. D) Spermatid in step 12 labeled with anti-Txl-2 preincubated with human recombinant Txl-2. No immunogold labelling of manchette nor axoneme (Ax) is evident. CL, cytoplasmic lobe, NR, nuclear ring. Bar, 0.2 µm.

Figure 10. Immunogold electron microscopy localization of Txl-2 protein in testicular arterioles. EM of testicular arteriole wall showing immunoreactivity in smooth muscle (Sm) cell. In contrast, the endothelium (En) does not appear to be immunogold labelled. CT, connective tissue, M, mitochondria; Lu, lumen. Bar, 0.2 µm.

Figure 11. Txl-2 and Δ5Txl-2 bind microtubules in vitro with differential affinity. A) 35S-labeled Txl-2 (FL) and Δ5Txl-2 (AS) were synthesized by in vitro translation and incubated with rat brain microtubules (MT). Binding reactions were pelleted through a cushion, washed and this procedure was repeated. Resulting supernatant 1 (s1), supernatant 2 (s2) and MT pellets (p) were analyzed for Txl-2 and Δ5Txl-2 protein by SDS-PAGE and autoradiography. Upper panel, Txl-2 binding experiments (FL), and lower panel, Δ5Txl-2 binding experiments (AS). Lanes a–c: proteins were incubated with brain MT. Lanes d–f: FL and AS proteins were analyzed for MT binding after preincubation of MT with recombinant Txl-2 protein. Lanes g–i: FL and AS proteins were analyzed for MT binding after preincubation of MT with anti-Txl2 antibodies. Binding of
Txl-2 (FL %) and Δ5Txl-2 (AS %) proteins relative to untreated samples (arbitrily set at 100 %) is indicated. B) To distinguish direct from indirect MT-binding, 35S-labeled Txl-2 (FL; lanes a - f) and Δ5Txl-2 (lanes g - l) were synthesized by in vitro translation and incubated with purified rat brain microtubules (brain MT) or with pure microtubules prepared by polymerisation of pure tubulin (pure MT). Pelleted MTs were washed and supernatants and pellets analyzed as described for Figure 12. S1 = supernatant 1, s2 = supernatant 2, p = pellet. Note strong binding of Δ5Txl-2 to pure microtubules.
**A**

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**B**

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Sadek et al. (2002) Figure 1
Figure 2

Sadek et al. (2002)
B

Nm23-H1
Nm23-H2
Nm23-H3
Nm23-H4

Txl-2
Nm23-H7B
Sptrx-2C
Nm23-H5
Nm23-H7A
Sptrx-2B
Nm23-H6

Sadek et al. (2002) Figure 2 continuation
Sadek et al. (2002) Figure 4

**A**

![Bar graph showing gene expression levels across different tissues.](chartA)

**B**

![Bar graph comparing gene expression in Testis, Lung, and Brain.](chartB)
Figure 5

A. Absorbance 280 nm vs. Fraction number (Elution time in hours). Peaks indicate the elution of Albumin, Ovoalbumin, Chymotrypsinogen A, and Ribonuclease A.

B. Mass spectrometry analysis with m/z values for -DTT and +DTT samples, showing absolute intensity for peaks at 18432, 36824, 36845, 18465, 73584, and 73525.
**Figure 5 Continuation**

Sadek et al. (2002)
Sadek et al. (2002) Figure 8
Sadek et al. (2002) Figure 9
A

\[
\begin{array}{cccccc}
& p & s_1 & s_2 & p & s_1 & s_2 & p & s_1 & s_2 \\
\text{Txl-2} & 100 & 100 & 90 & 100 \\
\Delta 5\text{Txl-2} & 100 & 49 & 28 & 28 \\
\end{array}
\]

B

\[
\begin{array}{cccccc}
& s_1 & s_2 & p & s_1 & s_2 & p \\
\text{pure MT} & 100 & 49 & 28 & 28 \\
\text{brain MT} & 100 & 49 & 28 & 28 \\
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Sadek et al. (2002) Figure 11
Table 1. Intron-exon organization of human Txl-2 gene.

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Sadek et al. (2002) Supplemental data 1