Quantitative Proteomic Analysis Using Isobaric Protein Tags Enables Rapid Comparison of Changes in Transcript and Protein Levels in Transformed Cells

Richard D. Unwin‡§, Andrew Pierce‡, Rod B. Watson¶, David W. Sternberg∥, and Anthony D. Whetton‡§‡‡

Isobaric tags for relative and absolute quantitation, an approach to concurrent, relative quantification of proteins present in four cell preparations, have recently been described. To validate this approach using complex mammalian cell samples that show subtle differences in protein levels, a model stem cell-like cell line (FDCP-mix) in the presence or absence of the leukemogenic oncogene TEL/PDGFRβ has been studied. Cell lysates were proteolytically digested, and peptides within each sample were labeled with one of four isobaric, isotope-coded tags via their N-terminal and/or lysine side chains. The four labeled samples are mixed and peptides separated by two-dimensional liquid chromatography online to a mass spectrometer (LC-MS). Upon peptide fragmentation, each tag releases a distinct mass reporter ion; the ratio of the four reporters therefore gives relative abundances of the given peptide. Relative quantification of proteins is derived using summed data from a number of peptides. TEL/PDGFRβ leukemic oncogene-mediated changes in protein levels were compared with those seen in microarray analysis of control and transfected FDCP-mix cells. Changes at the protein level in most cases reflected those seen at the transcriptome level. Nonetheless, novel differences in protein expression were found that indicate potential mechanisms for effects of this oncogene. *Molecular & Cellular Proteomics 4:924–935, 2005.

The development of approaches for measurement of relative expression of proteins between two (or more) samples is an essential aspect of systems biology. A common technique for this type of proteomic study has been two-dimensional gel electrophoresis, where proteins are separated by isoelectric point and molecular weight, and spot patterns are compared by sophisticated computer algorithms. Proteins of interest are then identified by MS. However, gel-based approaches have drawbacks, including amount of material required, reproducibility, and limiting of sensitivity by protein loss. These are balanced against recent advances in protein staining that allow intragel comparison of protein quantity from two or three samples (1).

An alternative approach to proteomic analysis is LC-MS (2). This provides an increased sensitivity compared with gel-based approaches and can catalogue protein present in a sample. However, relative quantitation of proteins using LC-MS is challenging. Quantification by analyzing two samples in parallel and comparing their mass spectrometric profiles is not feasible. Isotopic labeling of peptides, however, does allow two samples to be analyzed in a single experiment. Isotope-coded affinity tagging using ICAT reagent technology indicates peptide source, with peak height giving relative quantity (3). Comparing the ICAT reagent approach with two-dimensional gel electrophoresis, however, demonstrates that neither offers comprehensive coverage of a proteome (4). This is true in part because many proteins (and therefore peptides) do not contain cysteine, the amino acid used for covalent attachment of the isotopomer in ICAT reagents. Thus, much information can be discarded in the form of non-labeled peptides, whereas two-dimensional gel electrophoresis excludes many large, hydrophobic, and basic proteins.

Stable isotope labeling with amino acids in cell culture uses isotopes of essential amino acids (for example deuterated leucine) to label cells in culture (5). The samples are mixed, proteolytically digested, and run in LC-MS experiments. All leucine-containing peptides appear as “heavy” and “light” peaks, giving relative protein abundance. This elegant method can only be used on cultured cells; it is unsuitable for study of primary material. For ICAT reagent and stable isotope labeling with amino acids in cell culture technologies, the labeled peptides have different masses in an MS scan; this increases the number of measurable peptides and therefore the sensitivity of the method.
the complexity of the MS spectra and necessitates that MS/MS is therefore performed on the same peptide (the heavy and light labeled versions) twice, wasting analysis time.

Novel labeling reagents can overcome some of the limitations described above. Isobaric tags for relative and absolute quantitation (iTRAQ)\(^1\) use reagents that enable up to four samples to be analyzed within the same experiment. The labels consist of a protein-reactive group that labels all free amines (i.e. will label at the N terminus of all peptides and also the side chain of internal lysine residues), a balance group and a reporter group (6). The labels are isobaric, with a different distribution of isotopes between the reporter and balance groups. Hence, each labeled peptide appears at the same mass in an MS scan, but upon fragmentation in the mass spectrometer, the label dissociates and releases the reporter group as a singly charged ion of masses 114.1, 115.1, 116.1, or 117.1, respectively. Relative peak area indicates the contribution of each sample to total peptide present, providing a measure of relative abundance. The balance group is also lost and the remaining peptide fragments, which all have addition of the same mass (i.e. the protein reactive) group provide data from which to infer the peptide sequence.

The t(5,12) translocation found in chronic myelomonocytic leukemia results in the expression of the leukemogenic tyrosine kinase TEL/PDGFR\(^\beta\) and activation of the PDGFR\(^\beta\) tyrosine kinase domain (7). This stem cell disease has been modeled by expressing the TEL/PDGFR\(^\beta\) in a multipotent hematopoietic stem cell line, FDCP-Mix. The effects of oncogenic expression can be subtle yet lead to profound changes in cellular development. Perhaps unlike signaling for proliferation or apoptotic suppression, the appropriate tools for immediate analysis of potential effectors of altered development are not freely available. Herein, we report the validation and use of iTRAQ reagents on the FDCP-Mix TEL/PDGFR\(^\beta\) system as a paradigm for rapid, systematic definition of oncogenic processes using proteomics and the value of iTRAQ in permitting direct comparison of transcriptome data.

**EXPERIMENTAL PROCEDURES**

**Cell Line Preparation and Culture—**FDCP-Mix cells were transduced with TEL/PDGFR\(^\beta\) using a murine stem cell retroviral vector as described previously (8). Cells were routinely cultured in Fishers medium with 20% (v/v) horse serum supplemented with 10 ng/ml IL-3 (R&D Systems, Minneapolis, MN). Conditions for inducing myeloid differentiation FDCP-Mix cells were induced to differentiate by two methods as described previously (9). Briefly cells were cultured in Iscove’s modified Dulbecco’s medium supplemented with pre-selected fetal calf serum (20% (v/v)) and a combination of recombinant murine granulocyte macrophage-colony-stimulating factor (50 units/ml; Biogen IDEC, Zug, Switzerland), recombinant human macrophage-colony-stimulating factor (5 ng/ml; Amgen Biologics, Thousand Oaks, CA) plus recombinant murine IL-3 (0.1 ng/ml; Calbiochem, Nottingham, UK). Cells were prepared with a Cytospin centrifuge and stained with May Grunwald-Giemsa, and differential morphology was scored for greater than 100 cells per slide.

**Western Blotting—**Western blotting was carried out with standard protocols using a monoclonal antibody to the kinase domain of PDGFR\(^\beta\) or anti-phosphotyrosine antibodies (BD PharMingen, Oxford, UK). The phosphoprotein content (serine, threonine, and tyrosine) was measured by separation of total cell lysates on 10%T SDS-polyacrylamide gels and staining with Pro-Q diamond stain (Molecular Probes, Leiden, The Netherlands) per the manufacturer’s instructions.

**iTRAQ Reagent Labeling—**An overview of the workflow is shown in Fig. 1. 1–2 \(\times\) 10\(^6\) cells were lysed in 250 \(\mu\)l of 0.5 \(M\) triethylammonium bicarbonate (Sigma-Aldrich, St. Louis, MO) + 0.05% (w/v) SDS on ice for 20 min with regular vortexing. Protein was centrifuged at >10,000 \(\times\) g for 20 min at 4 °C, supernatant was removed, and protein quantified using the modified Bradford protein assay (Bio-Rad Laboratories). Protein (50 or 100 \(\mu\)g) in 20 \(\mu\)l of 0.5 \(M\) triethylammonium bicarbonate/0.05% SDS was reduced by addition of 2 \(\mu\)l of 50 mM tris-(2-carboxyethyl)phosphine and incubation at 60 °C for 1 h. Reduced cysteine residues were then blocked by addition of 1 \(\mu\)l of

![](image)

**Fig. 1. Workflow for the identification of changes in the proteome induced by TEL/PDGFR\(^\beta\) in the FDCP-Mix cell line, using iTRAQ reagents.** Three LC-MS/MS runs were performed to this protocol. To determine reproducibility of both the labeling reaction and the MS analysis internal controls were included. Confirmation of relative quantitation was thereby derived. The expression between FDCP-Mix and FDCP-Mix-Tel/PDGFR\(^\beta\) cells was compared twice in each MS analysis (114 versus 117 and 116 versus 117). The strong cation exchange step shown was employed to remove free iTRAQ reagent as well as to fractionate peptides for separate analyses by reversed-phase LC/MS/MS. PROQUANT is a program designed to integrate data from these isobaric tag experiments for relative protein quantification.

\(^1\) The abbreviations used are: iTRAQ, isobaric tags for relative and absolute quantitation; PDGFR\(^\beta\), platelet-derived growth factor receptor subunit \(\beta\); C/EBP, CCAAT/enhancer-binding protein.
200 μm methylmethanethiosulfate in isopropanol and incubation at room temperature for a further 10 min. Protein was then digested by addition of 10 μl of trypsin at 0.5 μg/μl and incubated at 37 ℃ overnight.

To label the peptides with iTRAQ reagent (Applied Biosystems, Warrington, UK), one unit of label (defined as the amount required to label 100 μg of protein) was thawed and reconstituted in 70 μl of ethanol, with vortexing for 1 min. The reagent solution was added to the digest and incubated at room temperature for 1 h. Labeling reactions were then pooled before analysis.

Peptide Fractionation and Mass Spectrometry—To remove excess, unbound iTRAQ reagent and to simplify the peptide mixture before reversed-phase LC-MS/MS, peptides were washed and fractionated off line using a strong cation exchange column (Applied Biosystems).

In brief, the peptide mixture was diluted 10-fold in loading buffer (10 mM potassium phosphate in 25% (v/v) acetonitrile), pH 3.0), and the pH was checked to ensure it remained between 2.5 and 3.3. Sample mixture was slowly injected onto the strong cation exchange cartridge and was washed with a further 1 ml of loading buffer to remove salts, tris-(2-carboxyethyl)phosphine, and unincorporated iTRAQ reagent. Samples were eluted from the column using 500-μl volumes of elution buffer (10 mM potassium phosphate in 25% (v/v) acetonitrile) containing increasing concentration of KCl. Salt concentrations used were 50, 100, 150, 200, 250, 300, 350, and 500 mM. Each salt fraction was then concentrated and dried in a SpeedVac (Thermo Electron, Waltham, MA).

Dried peptide fractions were resuspended in 250 μl of 2% (v/v) acetonitrile/0.1% (v/v) formic acid. For each analysis, 60 μl of the peptide sample was loaded onto a 15-cm reversed phase C18 column (75 μm i.d.) using an UltiMate pump (LC Packings, Amsterdam, The Netherlands) and separated over a 120-min solvent gradient from 5.9% (v/v) acetonitrile/0.1% (v/v) formic acid to 41% (v/v) acetonitrile/0.1% (v/v) formic acid on-line to a QSTAR XL mass spectrometer (Applied Biosystems). Data was acquired using an independent data acquisition protocol in which, for each cycle, the two most abundant multiply charged peptides (2+ to 4+) above a 10 count threshold in the MS MS spectrum were selected for MS/MS. Each peptide was selected twice and then dynamically excluded (±50 milli-charge units) for 40 s.

Data Analysis—Data were searched against a mouse KBMS3.0 protein database from the Celera Discovery System (Applied Biosystems). The database allowed for iTRAQ reagent labels at N-terminal residues, internal K and Y residues, and the methylmethanethiosulfate-labeled cysteine as fixed modification, plus one missed cleavage. Search parameters within ProQUANT were set with an MS tolerance of 0.15 Da, an MS/MS tolerance of 0.1 Da, and a minimum confidence score of 20. ProQUANT pooled data from all LC-MS runs. Assessment of these parameters for peptide and protein identification is described in Supplemental Table I.

Transcriptome Analysis—RNA from FDCP-Mix and TEL/PDGFRβ FDCP-Mix samples was prepared using TRIzol (Invitrogen) in triplicate samples and then cleaned using MiniElute RNAeasy Clean up kit (Qiagen, Valencia, CA) per the manufacturer’s instructions. Transcriptome analysis was undertaken using murine MOE430A Affymetrix chips by the CR-UK Affymetrix microarray facility, Paterson Institute (Manchester, UK). -Fold changes were calculated from the scaled data, where appropriate t-test analysis was applied.

RESULTS

Expression of TEL/PDGFRβ in the Multipotent Hematopoietic Cell Line FDCP-Mix Inhibits Differentiation but Has No Effect on Growth Factor Dependence—FDCP-Mix cells were transduced to express TEL/PDGFRβ (Fig. 2a). Previous experience with Ba/F3 cells transfected with BCR/ABL and TEL/PDGFRβ have shown that TEL/PDGFRβ has comparatively small effects on tyrosine phosphorylation (data not shown). However, TEL/PDGFRβ has small but significant effects on protein tyrosine phosphorylation and also significantly affects total phosphoprotein level (Fig. 2, b and c).

Differentiation-blocked cell lines can become growth factor-independent when expressing TEL/PDGFRβ (10). This did not occur in TEL/PDGFRβ-FDCP-Mix cells. FDCP-Mix cells differentiate to form mature cells when cultured in the appropriate cytokines (11). Expression of TEL/PDGFRβ inhibited this development (Fig. 2d). Culture conditions that induce myeloid development to 100% postmitotic cells (macrophages, neutrophils) in control FDCP-mix cells gave no such effect in TEL-PDGFRβ-transfected cells. After 20 days in culture, colony-forming clonogenic cells were still present that had primitive myeloid cell or blast cell morphology. This effect on differentiation led us to systematically analyze potential differences in transfected and non-transfected cells using transcriptomic and proteomic methods. To do this, the proteomic method required validation.

Analysis of a Standard Mix of Proteins Using iTRAQ Reagent Labels—A defined six-protein mix that had been enzymatically digested with trypsin was used to confirm the accuracy of ratiometric quantitation of the iTRAQ reagents. The tryptic digest was halved, and each half was labeled with either reagent 116 or 117. These differentially labeled digests were mixed at various ratios (1:1, 2:1, and 1:3) and analyzed by LC-MS/MS. A representative spectrum is shown in Fig. 3a. Relative quantitation of proteins by iTRAQ reagent technology was both accurate and reproducible for five proteins (Fig. 3b). The sixth protein in the standard mixture was not detected with sufficient peptides to allow quantitation. Overall 117:116 ratios of 0.9699, 0.5885, and 3.1748 were obtained. We confirmed that no peptides remained “unlabeled” by analysis of parent ion masses derived for the MS analysis and comparison with theoretical tryptic digests of the six proteins. All isobaric forms of the iTRAQ reagent tag labeled equally efficiently.

Identification of TEL/PDGFRβ Induced Alterations in the Proteome—The potential value of the system was further examined on lysates from mammalian FDCP-Mix cells and FDCP-Mix cells transduced with the TEL/PDGFRβ leukemic oncogene. In these experiments, microarray data was available for comparison from the FDCP-Mix cells described above. The experimental design is outlined in Fig. 1. All four labels were used, allowing the use of internal controls and replicates within the same LC-MS experiment for this paradigm study. This experimental design allowed the accuracy of quantitation to be verified by including a 1:1 ratio of control sample (114:116) and a 2:1 ratio of the TEL/PDGFRβ FDCP-Mix cell (117:115), as shown in Fig. 1. Furthermore, an internal replicate for the comparison of TEL/PDGFRβ-expressing and non-expressing cells is thus provided. The experiment was
run in triplicate. Labeling efficiency with iTRAQ reagents was high (>99%), and both intra- and interexperiment quantitation was highly reproducible, with overall ratios of 1.064, 0.967, and 1.093 when samples were mixed 1:1 and 2.127, 2.128, and 2.217 when samples were mixed 2:1. It is noteworthy that ratios detected were normally distributed; therefore, statistical tests such as Student's t test can be applied. In addition, replication within the experiment achieved an acceptable standard, with an average 116:114 ratio of 1.072 ± 0.07 (S.D.; n = 3) (Fig. 4a), and the replicate FDCP-Mix versus FDCP-Mix-TEL/PDGFRβ analyses (117:114 versus 117:116) had a correlation coefficient of 0.93, confirming the reproducibility of the technique. Examples of the spectra produced by iTRAQ reagent-labeled peptides are shown in Fig. 5, a–c, where peptides showing no change between samples, down-regulation or up-regulation by TEL/PDGFRβ expression are illustrated.

Comparison of iTRAQ™ Reagent Data to cDNA Microarray—Relatively few changes are seen in cDNA microarrays from TEL/PDGFRβ-transfected and control FDCP-Mix cells, reflecting the relatively small effect of this oncogene on protein tyrosine phosphorylation. This paucity of change in the transcriptome led us to confirm changes using proteomics. Analysis of microarray data revealed that, with a 1.5-fold change cut-off, a total of 105 transcripts show a significant increase with TEL/PDGFRβ expression, and 159 transcripts show decrease (see Supplemental Table I).

The data derived permitted comparison of individual transcript/protein levels. All of the differences identified by iTRAQ reagents (Table I), along with 87 unchanged proteins, were analyzed, giving a set of 100 proteins/transcripts in total as examples of the data recorded. The average ratio of the

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**Fig. 2. Consequences of expression of TEL/PDGFRβ in FDCP-Mix cells.** Expression and consequence of TEL/PDGFRβ in transfected cells as analyzed by Western blot analysis of total cell lysates using anti-PDGFRβ (a) and anti-phosphotyrosine (b) antibodies. c, level of total protein phosphorylation as assessed by staining whole-cell lysate with Pro-Q Diamond. In each of the three gel images, lane 1 indicates control cells and lane 2 indicates TEL/PDGFRβ-transfected cells. d, effect of TEL/PDGFRβ on differentiation as assessed by culturing control and TEL/PDGFRβ-transfected FDCP-Mix cells in 0.1 ng/ml IL-3 and 50 units/ml granulocyte macrophage-colony-stimulating factor plus 5 ng/ml macrophage-colony-stimulating factor. Cells were seeded at 1 × 10⁶/ml, and morphology was determined on days 0, 4, and 7. Results shown are the mean of three experiments; S.E. was <10% in all cases. Late progenitor cells indicate metamyelocytes, myelocytes, and promonocytes.
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a

LDAINENJ

m/z, amu

116.1

117.1

b

| Sample ratio | Beta-Gal | Albumin | Transferrin | Lactoglobulin | Lysozyme |
|--------------|----------|---------|-------------|---------------|----------|
| 1:1          |          |         |             |               |          |
| 2:1          |          |         |             |               |          |
| 1:3          |          |         |             |               |          |
mRNA expression between FDCP-Mix and FDCP-Mix-TEL/PDGFRβ cell lines using iTRAQ reagents. For each specific transcript, the ratio determination in the internal control experiment comparing 100 μg of control protein labeled with either the 114 or 116 iTRAQ reagent labels. The x-axis indicates the number of peptides identified (using MS/MS) from each protein. Each spot represents one protein identified. Dotted line shows the average 116:114 ratio across the complete data set.

**FIG. 4.** Ratiometric analysis of proteins from FDCP-mix and FDCP-mix-TEL/PDGFRβ cell lines using iTRAQ reagents. a, plot to show the reproducibility of ratio determination in the internal control experiment comparing 100 μg of control protein labeled with either the 114 or 116 iTRAQ reagent labels. The x-axis indicates the number of peptides identified (using MS/MS) from each protein. Each spot represents one protein identified. Dotted line shows the average 116:114 ratio across the complete data set. b, reproducibility of the iTRAQ reagent-relative quantitation: the ratios for five representative proteins over three experiments using the FDCP-Mix and FDCP-Mix-TEL/PDGFRβ cell lines are shown. Error bars are defined as in Fig. 2b. c, plot showing the distribution of the oncogene/control (117:114) ratios of proteins identified in the iTRAQ reagent experiment, plotted against the number of peptides identified and therefore quantified from that protein. Unchanged proteins are shown by ■, and differentially expressed proteins are shown by ●. Dotted line represents the mean FDCP-Mix TEL/PDGFRβ:FDCP-Mix ratio from the complete data set.

**FIG. 3.** Ratiometric analysis of five standard proteins labeled with iTRAQ reagents 116 or 117 and mixed at 1:1, 1:2, and 3:1 ratios. a, a representative spectrum of peptide LDAINENJ (where J represents an internal lysine labeled with iTRAQ reagent via the ε-amino acid in its side chain) from β-lactoglobulin derived for a 2:1 mixture of 116- and 117-labeled samples. The peptide sequence is derived from the fragment ion pattern; inset, the relative abundances were determined by comparison of the areas under the 116.1 and 117.1 peaks. b, summary of the ratios for the five standard proteins analyzed. Error bars are S.D. of n observations, where n is equal to the number of peptides identified in that protein.
TABLE I

The relative quantification of protein levels and mRNA levels in FDCP-Mix cells and TEL/PDGFRβ-FDCP-Mix cells

For comparison, 100 proteins (randomly selected proteins plus those in which changes between the two cell lines were seen using the isobaric tag quantification approach) were selected. Random selection was made via search databases (generating the accession codes shown: emb, EMBL; gb, GenBank™; pir, Protein Information Resource; rf, RefSeq (NCBI); spt, Swiss-Prot; trm, TrEMBL). The total peptides columns refers to the number of peptides from that protein fragmented within three replicate experiments. The relative ratios for each protein were calculated as a weighted value combining averages from three replicate experiments. Ratios shown are: average control/control, an average (1) oncogene/control, containing data from comparison of 117:116 ratios. Also shown is the fold change in the ratio of protein in the TEL/PDGFR

| Accession | Protein | Total peptides | Average control/control | Average (1) oncogene/control | Average (2) oncogene/control | -Fold cDNA |
|-----------|---------|----------------|------------------------|-----------------------------|-----------------------------|-----------|
| pir | S31975 | 14-3-3 protein ε | 7 | 1.115 | 1.061 | 0.981 | 1.034 |
| rf | NP_853613.1 | 14-3-3 protein γ | 8 | 1.093 | 1.072 | 1.006 | 1.042 |
| spt | P63101 | 14-3-3 protein ζ | 8 | 1.111 | 1.043 | 0.968 | 1.106 |
| spt | P47911 | 605 ribosomal protein L6 | 10 | 1.059 | 0.785 | 0.745 | 0.995 |
| rf | NP_031501.1 | Acidic ribosomal phosphoprotein P0 | 7 | 1.065 | 0.789 | 0.759 | 1.028 |
| spt | P51881 | ADP, ATP carrier protein, fibroblast isomform | 18 | 1.108 | 0.999 | 0.919 | 0.940 |
| spt | Q60604 | Adseverin | 36 | 1.066 | 1.000 | 0.974 | 1.040 |
| spt | P47738 | Aldehyde dehydrogenase, mitochondrial | 5 | 0.509 | 1.074 | 1.034 | 0.926 |
| spt | P56480 | ATP synthase β chain, mitochondrial | 10 | 1.123 | 1.238 | 1.127 | 1.058 |
| Emb | CAA27396.1 | β-Actin | 64 | 1.123 | 1.061 | 0.956 | 0.988 |
| spt | Q9CWJ9 | Bifunctional purine biosynthesis protein | 7 | 1.079 | 1.691 | 1.661 |
| Emb | CA05361.1 | Bip | 13 | 1.138 | 0.976 | 0.885 | 0.835 |
| gb | AA62450.1 | Calnexin | 15 | 1.012 | 1.010 | 1.021 | 0.984 |
| Spt | P14211 | Calreticulin (CRP55) | 41 | 1.109 | 0.939 | 0.867 | 0.963 |
| Spt | P97742 | Carnitine O-palmitoyltransferase | 4 | 1.167 | 1.024 | 0.915 | 1.199 |
| pir | HHMS60 | Chaperonin groEL | 13 | 0.892 | 1.538 | 1.757 | 0.913 |
| rf | NP_031662.1 | Chaperonin subunit 2 (β) | 9 | 1.032 | 0.746 | 0.749 | 1.078 |
| spt | P18760 | Cofilin, non-muscle isoform | 5 | 1.081 | 1.077 | 1.015 | 1.008 |
| prf | 1513495A | Cu/Zn superoxide dismutase | 10 | 1.110 | 1.056 | 0.973 | 0.996 |
| rf | NP_031834.1 | Cytochrome c, somatic | 9 | 1.088 | 0.887 | 0.829 | 0.885 |
| spt | P49717 | DNA replication licensing factor MCM4 | 5 | 1.075 | 0.878 | 0.827 | 1.234 |
| Trm | Q8BM3 | Dolichyl-diphosphooligosaccharide | 6 | 1.126 | 0.917 | 0.821 | 1.085 |
| sp | O70251 | Elongation factor 1-α | 7 | 1.038 | 0.952 | 0.935 | 0.957 |
| spt | P10126 | Elongation factor 1-α | 12 | 0.709 | 0.797 | 0.754 | 1.080 |
| spt | P08113 | Endoplasmic precursor (GRP94) | 17 | 0.999 | 1.106 | 1.176 | 0.876 |
| rf | NP_075608.1 | Enolase 1, α neuron | 28 | 1.093 | 1.159 | 1.073 | 1.036 |
| rf | NP_031933.1 | Eugaryotic translation elongation factor 2 | 27 | 1.117 | 0.771 | 0.705 | 0.977 |
| spt | P26040 | Ezrin | 12 | 1.081 | 0.969 | 0.917 | 0.976 |
| Trm | Q9EOQ | Fatty acid synthase (Fasn protein) | 6 | 1.022 | 0.821 | 0.824 | 1.003 |
| spt | P35550 | Fibulilim (Nucleolar protein 1) | 8 | 1.050 | 0.933 | 0.900 | 1.030 |
| rf | XP_127565.4 | Filamin B, α | 22 | 1.052 | 0.862 | 0.836 |
| rf | XP_207130.3 | Filamin, α | 10 | 1.002 | 0.908 | 0.953 |
| spt | P05064 | Fructose-bisphosphate aldolase A | 28 | 1.054 | 1.260 | 1.224 | 1.041 |
| rf | NP_032181.1 | Glucose phosphate isomerase 1 | 13 | 1.052 | 1.037 | 1.018 | 1.033 |
| spt | P26443 | Glutamyl dehydrogenase, mitochondrial | 7 | 1.114 | 1.055 | 0.988 | 0.986 |
| rf | NP_032110.1 | Glyceraldehyde-3-phosphate dehydrogenase | 29 | 1.060 | 1.126 | 1.087 | 0.987 |
| spt | P04187 | Granzyme B(G,H) | 12 | 1.111 | 1.194 | 1.118 | 0.987 |
| rf | NP_034566.1 | H2A histone family, member X | 27 | 1.089 | 0.817 | 0.767 | 1.189 |
| pir | Q19893 | H3.3 like histone MH921 | 8 | 1.066 | 0.799 | 0.758 | 0.997 |
| rf | NP_034610.1 | Heat-shock protein 1α | 16 | 1.051 | 0.835 | 0.810 | 0.948 |
| Emb | CA56631.1 | Histone 1A, H1a | 16 | 1.054 | 1.146 | 1.107 | 1.040 |
| rf | NP_056501.1 | Histone 1, H1c | 19 | 1.122 | 0.949 | 0.875 | 0.827 |
| rf | NP_835503.1 | Histone 1, h2bg | 32 | 1.062 | 1.261 | 1.217 | 1.009 |
| pir | S5110 | Histone H2A | 45 | 1.142 | 0.846 | 0.762 | 1.138 |
| gb | AAH58529.1 | Histone H4 | 30 | 1.111 | 0.928 | 0.856 | 1.095 |
| spt | Q99020 | hnRNP A/B | 8 | 1.049 | 1.123 | 1.097 | 0.985 |
| gb | AAH06694.1 | hnRNP K protein | 11 | 1.179 | 1.077 | 0.927 | 1.129 |
Regarding the iTRAQ experiments, all five transcriptomic differences were mirrored in the protein changes seen. It is noteworthy that, with respect to developing studies on mechanisms of transformation, a set of proteins showed

tained five differences. These were cathepsin G, L-plastin, Mast cell protease 8, myeloperoxidase, and protein disulfide isomerase. Regarding the iTRAQ experiments, all five transcriptomic differences were mirrored in the protein changes seen. It is noteworthy that, with respect to developing studies on mechanisms of transformation, a set of proteins showed

| Accession | Protein | Total peptides | Average (1) oncogene/control | Average (2) oncogene/control |
|-----------|---------|----------------|-------------------------------|-------------------------------|
| NP_034829.1 | Lactate dehydrogenase 1, A chain | 36 | 1.124 | 1.040 |
| JN0066 | Leukotriene-A4 hydrolase | 8 | 1.073 | 0.868 |
| CA65761.1 | M2-type pyruvate kinase | 51 | 1.104 | 1.137 |
| AAA9509.1 | Malate dehydrogenase | 9 | 1.036 | 1.000 |
| NP_620084.1 | Methylene-tetrahydrololate dehydrogenase 1 | 5 | 1.045 | 0.954 |
| NR_034963.1 | Mesoestranol | 10 | 1.080 | 0.975 |
| Q9EQK1 | NADP+-specific isocitrate dehydrogenase | 11 | 1.136 | 0.930 |
| Q8VDD5 | Nonmuscle heavy chain myosin II-A | 24 | 1.063 | 1.004 |
| P09405 | Nucleolin (protein C23) | 17 | 1.093 | 0.892 |
| Q8VDW0 | Nuclear RNA helicase, DECD variant of DEAD box family | 13 | 1.088 | 1.085 |
| NP_077155.1 | Nucleolar protein 5A | 4 | 1.053 | 1.248 |
| Q6937 | Nucleophosmin | 13 | 1.099 | 0.799 |
| AA6557 | Nucleoside diphosphate kinase | 9 | 1.103 | 0.962 |
| Q01176 | Nucleoside diphosphate kinase B | 7 | 1.129 | 0.906 |
| P24369 | Peptidyl-prolyl cis-trans isomerase B | 6 | 1.110 | 1.330 |
| NP_032933.1 | Peptidyl-prolyl isomerase A | 10 | 1.119 | 1.114 |
| SPT | Phosphoserine aminotransferase | 5 | 1.051 | 1.117 |
| NP_035202.1 | Profilin 1 | 9 | 1.055 | 1.046 |
| P27773 | Protein disulfide isomerase A3 | 11 | 0.984 | 0.954 |
| NP_033417.1 | RAN, member RAS oncogene family | 14 | 1.067 | 1.216 |
| Q61599 | Rho GDP-dissociation inhibitor 2 | 4 | 1.110 | 1.026 |
| NP_038749.1 | Ribosomal protein L7a | 8 | 1.005 | 0.937 |
| NP_079862.1 | Ribosomal protein L15 | 8 | 0.951 | 0.862 |
| NP_075029.1 | Ribosomal protein L23 | 18 | 1.119 | 0.928 |
| NP_035430.1 | Ribosomal protein S7 | 6 | 1.105 | 0.929 |
| NP_084043.1 | Ribosomal protein S9-like | 7 | 1.123 | 0.929 |
| NP_080809.1 | Ribosomal protein S13 | 9 | 1.120 | 0.903 |
| AAA16796.1 | Ribosomal protein S18 | 7 | 1.070 | 0.801 |
| Q69843 | Structure-specific recognition protein 1 | 6 | 1.071 | 0.709 |
| NP_063932.1 | Succinate-CoA ligase, GDP-forming, α | 7 | 1.101 | 1.117 |
| AAA4038.1 | T complex polypeptide 1 | 7 | 1.070 | 0.905 |
| P26639 | Talin 1 | 12 | 0.972 | 1.076 |
| Q9WVA4 | Transgelin 2 | 10 | 1.047 | 1.006 |
| P40142 | Transketolase | 16 | 1.074 | 0.835 |
| JL0149 | Transplantation antigen P198 | 5 | 1.043 | 0.839 |
| ISMST | Triosephosphate isomerase | 11 | 1.055 | 1.310 |
| NP_666228.1 | Tubulin, β 2 | 54 | 1.109 | 0.642 |
| AAA43053.1 | Valosin containing protein | 15 | 1.068 | 1.019 |

Differentially expressed proteins as determined by iTRAQ

| Accession | Protein | Total peptides | Average (1) oncogene/control | Average (2) oncogene/control |
|-----------|---------|----------------|-------------------------------|-------------------------------|
| Q64333 | 10-kDa heat shock protein | 5 | 1.079 | 1.513*** |
| P46634 | 60S ribosomal protein L10 | 12 | 1.072 | 0.624* |
| AAA21937.1 | Aldo-keto reductase family 1, C13 | 22 | 1.105 | 0.970* |
| P28293 | Cathepsin G | 9 | 1.059 | 0.617*** |
| A54935 | DnaK-type molecular chaperone hsc70 | 37 | 1.057 | 0.813** |
| NP_853613.1 | Eukaryotic translation initiation factor 5A | 7 | 1.137 | 0.498*** |
| AAA37666.1 | 84-kDa heat-shock protein | 34 | 1.087 | 0.712*** |
| NP_031542.1 | Heterogeneous nuclear ribonucleoprotein D | 7 | 1.068 | 1.382** |
| Q61233 | L-plastin | 40 | 1.081 | 0.748* |
| P43430 | Mast cell protease 8 | 3 | 1.062 | 0.318*** |
| A45935 | 60S ribosomal protein L10 | 12 | 1.072 | 0.624* |
| Q7TMS4 | Myeloperoxidase protein | 10 | 1.035 | 0.307*** |
| P09103 | Protein disulfide isomerase | 27 | 1.122 | 0.696*** |
| Q60931 | Voltage-dependent anion-selective channel protein 3 | 5 | 1.058 | 0.638** |

* Statistical significance of the change as determined by Student t test is depicted (*, p < 0.05; **, p < 0.01; *** , p < 0.001).
changes (subtle, albeit statistically significant) in the proteome but not in the transcriptome. These included a 60S ribosomal protein subunit, aldo-keto reductase family member C13, and eukaryotic translation initiation factor 5A. This initial scan of the value of the iTRAQ approach therefore reveals that it will offer advantages over transcriptome analysis. Furthermore, specific proteins identified using iTRAQ LC-MS/MS were not assayed within the microarray. These included Filamins A and B, and bifunctional purine biosynthesis protein PURH. This assessment was confirmed after searching using alternative protein names, gene names, and accession numbers. iTRAQ therefore offers objective, non-selective sample analysis.

In the five changes showing consistency at transcript and protein level (cathepsin G, mast cell protease 8, myeloperoxidase precursor, protein disulfide isomerase, and l-plastin) the -fold changes in this set were remarkably similar to the -fold changes detected in their mRNA levels. The iTRAQ reagent and cDNA microarray data sets also agree in that most of the changes identified are subtle (2-fold or less).

**DISCUSSION**

iTRAQ reagent technology is a newly developed method for relative quantification of proteins from up to four samples. It has immense potential to improve the sensitivity and quality of mass spectrometric analysis of the proteome. Validation of the approach is reported here using a defined protein mixture and cell lysates from a disease model. iTRAQ reagent labels all peptides at their N terminus, along with free amines in lysine side chains, hence all of the peptide population is labeled, allowing more peptides to be quantified from each protein and increasing the quality of the data obtained by this approach. We have demonstrated that it is possible, with both standard proteins and whole-cell lysates, to label four peptide mixtures to completion and, using LC-MS/MS, to identify the relative quantities of the peptide emanating from four samples. The labeling protocol for iTRAQ reagents is simple, with relative quantification of proteins from up to four samples. It can be achieved using iTRAQ reagents and multiple subcellular fractions, multidimensional chromatography, plus ion exclusion lists in iTRAQ reagent-based experiments.

Several proteins are differentially expressed in the presence of TEL/PDGFRβ. Heterogeneous nuclear ribonucleoprotein D, shown to be increased by TEL/PDGFRβ expression, is an mRNA binding protein that has been implicated in tumorigenesis (13) and that is a target of the leukemogenic oncogene BCR/ABL (14). In our study TEL/PDGFRβ expression decreased Hsc70 levels, whereas previous studies have shown that increased expression of Hsc70 inhibits transformation (15). Myeloperoxidase protein, decreased by TEL/PDGFRβ, is expressed in early myeloid progenitors (16); therefore, its decreased levels in the differentiation-blocked TEL/PDGFRβ-expressing cells may provide clue as to the mechanism for this block. Myeloperoxidase expression is regulated by transcription factors such as Pu.1 and the C/EBP family (17). These proteins regulate myelopoiesis and loss of either Pu.1 or C/EBPα leads to compromised ability to produce mature

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2 A. D. Whetton and C. A. Evans, unpublished observations.

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**Fig. 5. Identification of differentially expressed proteins.** a, representative MS/MS spectrum showing a peptide, FTQAGSEVSALLGR, from ATP synthase β that does not show an expression change upon TEL/PDGFRβ expression and the iTRAQ reagent peaks used for relative quantitation (inset). b, representative MS/MS spectrum showing a peptide, VFFASWR, from myeloperoxidase, a protein that shows a decrease in expression with TEL/PDGFRβ and the iTRAQ reagent peaks used for relative quantitation showing the decreased levels of the 115 and 117 labels compared with control protein labeled with 114 and 116 (inset). c, representative MS/MS spectrum showing a peptide VVLDDJDYFLFR (where J represents an internal lysine labeled with iTRAQ reagent via the ε-amino acid in its side chain) from 10-kDa heat-shock protein that shows an increase in expression with TEL/PDGFRβ and the iTRAQ reagent peaks used for relative quantitation showing the increased levels of the 115 and 117 labels compared with control protein-labeled with 114 and 116 (inset).
cells (18, 19). C/EBPα transcript levels are decreased 1.5-fold (p = 0.017) in TEL/PDGFRβ-transfected cells, although Pu.1 and other C/EBP family members are unchanged at the transcript level. Thus, the data we have derived allow further experiments on the mechanism of differentiation blockade in these cells. These can include pre-enrichment and selective searching for ions from C/EBPα in our iTRAQ reagent experiments to allow relative quantitation of transcription factor levels. In addition, Cathepsin G, which is also reduced in expression, is a serine protease highly expressed in promyelocytes (20, 21); it has a role in hematopoietic stem cell mobilization and differentiation and so may also play a role in the TEL-PDGFRβ-mediated differentiation block. Mutations in such proteases have been implicated in neutropenia (22, 23). Perhaps even more relevant is that cathepsin L has been shown to locate to the nucleus and regulate transcription via a proteolytic mechanism (24). Cathepsin G may have a similar function.

Comparison of this data set with a cDNA microarray data set from the same cell line provides relatively high levels of agreement between transcripts and protein level. All five of the changes from the 100-transcript sample set were also detected by iTRAQ; of these, all showed a similar level of change. However, the iTRAQ reagent approach identified changes in proteins that are caused by post-transcriptional effects, in that no change is seen in the levels of mRNA. A comparison of this type was previously almost impossible, because stable isotope or gel-based approaches tend to focus on identifying proteins whose expression changes, rather than the relative abundances of all proteins in a sample.

In conclusion, we have shown that iTRAQ protein labeling reagents can be employed to successfully identify proteins in which expression is potentially modified. This has the advantage of using multiple samples in a single LC-MS/MS run. The iTRAQ reagent produced high quality, reproducible data regarding relative expression levels in up to four samples. Comparison of the iTRAQ reagent data with cDNA microarray data suggests a high degree of similarity; all changes in a subset of the cDNA microarray are replicated by iTRAQ reagent analysis. However, iTRAQ reagent experiments also defined several other changes not detected by the cDNA array. iTRAQ reagent technology has great value as a new method for relative quantification of proteins in enriched complexes, organelles, and whole cell lysates.

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