5D proteomic approach for the biomarker search in plasma: Acute myeloid leukaemia as a case study

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Acute myeloid leukaemia (AML) is a type of cancer affecting all ages but it is more common in adults, as compared to children. Recent advancements in proteomics and mass spectrometry tools, offer a comprehensive solution to study the molecular complexity of diseases, such as cancers. This study is focused on the proteomic profiling of AML in comparison to healthy control for which, a systematic 5D proteomic approach for the fractionation of pooled plasma samples was used. Methodology includes depletion of Top-7 abundant proteins, ZOOM-isoelectric focusing (ZOOM-IEF), two-dimensional gel electrophoresis (2-DGE), and matrix-assisted laser desorption/ionization mass spectrometry (MALDI-MS) analysis followed by the validation of identified biomarker proteins using enzyme linked immunosorbent assay (ELISA). Up-/down-fold changes in concentration of proteins were observed in 2-DGE of AML in comparison with the healthy control and a total of 34 proteins were identified in fractioned plasma. Among them, fifteen proteins were significantly differentiated and five proteins; SAA1, complement factor C7, ApoE, plasminogen, and ApoA1 were later verified by ELISA in individual samples, which showed that SAA1 and plasminogen could be used as potential biomarker for AML.

5D proteomic profiling using various approaches including mass spectrometry are being effectively used as a powerful tool for the understanding of diseases and identification of biomarkers particularly in cancers6. However, identification of reliable and sensitive protein biomarker depends on many factors specially the proteomic fractionation strategy being applied. A number of studies based on mass spectrometry, combining proteomic technologies such as two-dimensional gel electrophoresis (2-DGE), difference gel electrophoresis...
is cost effective, include greater statistical power, improved ability to compare results and validation of models. Blood collection tubes, containing K2-ethylenediaminetetraacetic acid (K2-EDTA). The plasma was separated by centrifugation at 2,200 × g for 10 min at 4 °C. Pool of individual samples was used in this study, because pooling is cost effective, increase greater statistical power, improved ability to compare results and validation of models. Pooled plasma has already been applied in many studies. To make healthy pool, equal volumes of each individual healthy plasma sample were mixed to obtain the healthy Pakistani pooled plasma. Similarly, AML samples were also pooled by combining equal amounts of every AML specimen. Pooled plasma samples were then subjected to aliquoting, and stored at −80 °C until further processing.

1D SDS-PAGE analysis of samples. One-dimensional sodium dodecyl sulphate polyacrylamide gel electrophoresis (1D SDS-PAGE) was performed for comparative analysis of healthy and diseased samples on X Cell SureLock system (Invitrogen). Chemicals and reagents for 1D SDS-PAGE were purchased from Invitrogen (USA). β-mercaptoethanol, sucrose, and Tris HCL were purchased from Sigma Aldrich (USA).

Depletion of abundant proteins through MARS column. Depletion of top seven most abundant proteins was carried out using Multiple Affinity Removal Column (MARS) Hu-7 (4.6 × 50 mm), purchased from Agilent (USA) on ÄKTA ™ FPLC system (GE Healthcare, Sweden). The column has an affinity for the albumin, IgG, IgM, transferrin, antipyrin, haptoglobin (HPT), and filtrypsin, haptoglobin. Protein inhibitors; ethylene diamine tetra-acetic acid (EDTA), leupeptin, pepstatin-A, and phenyl methylsulphonyl fluoride (PMSF) were purchased from Sigma Chemicals (USA). 500 mL Vacuum Filter/Storage Bottle System, 0.22 μm was purchased from Corning (USA). The plasma sample (300 µL) was depleted according to the kit protocol. The plasma amount used was according to the column capacity, and required sample size for multiple replicate depletions. 1D SDS-PAGE analysis of all fractions, including bound and unbound fractions was performed to check the sample recovery after depletion. Unbound fractions were pooled and concentrated using 5 kDa molecular weight cut off (MWCO) tubes for both pools. For concentrating the pooled bound fractions, the sample was centrifuged several times at 3,500 rpm for 15 min at 4 °C to obtain an amount of 200 µL. Enrichment efficiency was checked by loading the equivalent to 0.1 µL plasma from the fractions before concentrating and after concentrating.

Reduction and alkylation. Urea was purchased from Invitrogen (USA), tris(hydroxymethyl)aminomethane (Tris) from Boehringer Mannheim (Germany), dichlorodiphenyltrichloroethane (DDT) and iodoacetamide (IAM) from SERVA (Germany). Acetone, and trichloroacetic acid (TCA) were purchased from Fisher Scientific (UK) and Scharlau (Spain), respectively. For denaturation and pH adjustment, concentrated depleted sample was adjusted to 8 M urea and 20 mM Tris. To reduce the proteins, the sample was adjusted to 20 mM DTT, and subsequently for alkylation, to 50 mM IAM. To quench the alkylation, the sample was adjusted to 1% TCA solution, were mixed and placed at −20 °C for 1 hr. to precipitate the proteins. The tube was then centrifuged at 5,000 rpm for 30 min to have the proteins pelleted down. The pellet was washed two to three times with ice-cold acetone.

ZOOM-IEF. The ZOOM-IEF® Fractionator Combo Kit including ZOOM-IEF fractionator was purchased from Invitrogen (USA). The sample pellet was dissolved in ZOOM buffer by vortexing for 5 min, followed by
Fractionation was performed according to established protocol\(^2\). 1D SDS-PAGE analysis of all five ZOOM fractions was performed to check the efficiency of IEF, and to compare the samples from healthy persons with AML patients.

2-D gel electrophoresis. 2-DGE was performed on Bio-Rad PROTEAN IEF cell. The ReadyPrep 2-D Starter Kit, ReadyStrip IPG Strips, Ready Gel precast gel, mineral oil, 10X Tris/glycine/SDS Buffer, and paper wicks were purchased from Bio-Rad (USA). Buffers were prepared according to the kit protocol. 125 µL of freshly prepared rehydration/sample buffer for 7 cm IPG strip was added in a conical centrifuge tube containing the sample pellet, and vortexed to dissolve the pellet. The sample was centrifuged to settle down the fine particles. The whole procedure was performed according to the kit protocol. Gel images were taken through Gel DOC 800 system (Bio-Rad, USA).

Mass spectrometric analysis. Analysis was performed using MALDI-TOF-TOF MS (Ultraflex III, Broker Daltonics Germany). Mass spectrometric profile was obtained by flexAnalysis version 3.0. (Bruker Daltonics). The protein spots of interest were extracted from the stained gels using the manual cutting procedure and digested according to formerly mentioned protocol\(^3\)\(^,\)\(^4\).

The digested peptides were analysed using standard protocol\(^5\). Briefly, the samples were mixed with equal amounts of freshly prepared α-cyano-4-hydroxycinnamic acid in acetonitrile (ACN) in 1:1 ratio. Calibration of MALDI-TOF was carried out in the reflector positive mode using peptide calibrant standard I (Bruker Daltonics). A 337-nm nitrogen laser and a 2 GHz digitilizer were used. Mass spectra were obtained with 25 KV of ion acceleration, 6 KV lens potential, and high gating strength to deflect ions with a mass below 500 m/z values. Spectra were obtained in the mass range of 500–3000. Every spectrum was the sum of 2000 laser shots within the same spot (200 shots/position) and intensity of 20–40%.

ELISA analysis. ELISA was performed for five proteins, serum amyloid A (SAA1), plasminogen, apolipoprotein E (ApoE), complement factor C7 and apolipoprotein A1 (ApoA1) on Thermo Fischer Scientific ™ Multiskan™ FC Microplate Photometer (USA). ELISA kits were purchased from Crystalchem (USA) and Assaypro (USA). Plasma samples of 15 healthy individuals and 18 AML patients were diluted according to the kit protocol, using 1X diluent. Standard and diluted samples were dispensed into wells. After one-hour incubation, the samples were aspirated and the wells were washed 4 times using 1X wash buffer. The antibody-HRP conjugate was then applied for 20 min. On completion of this step, 4 times washing was carried out again using 1X wash buffer. The substrate solution was then added and left for 10 min for reaction completion. Stop solution was added and readings were taken at 450 and 630 nm. This procedure with some modifications according to the manufacturer’s protocol was applied to validate our results for selected proteins; complement factor C7, plasminogen, and ApoE, for which ELISA kits were purchased from Assaypro (USA).

Statistics and data analysis. The analysis of 2-DGE images, detection of spots, spot matching, and semi-quantitative statistical analysis were performed using the Bio-Rad PDQuest version 8.0.1. Bio-Rad (USA). Master gel was used to compare the gels of healthy and AML pool. The analysis involved matching the gels, differences and similarities in spots pattern, background subtraction, and removal of artefacts (horizontal and vertical streaks). T-test was used to study the differential protein expression among protein spots from healthy and AML pool gels, with \( p < 0.05 \) and four-fold change in spot intensity was selected as threshold. After automated matching, the detected spots were manually edited for greater accuracy.

After MALDI-MS the protein identification was carried out through MASCOT database search – Matrix Science, based on mass fingerprinting (PMF) using Swiss-Prot and NCBIinr databases. Peptide modification which we have chosen as a fixed modification during the search, was carboxymethylation of cystein. The oxidation of methionine was used as variable modification. The maximum number of missed cleavages was set to 1, peptide tolerance 100 ppm/1 Da, and \( p < 0.05 \) were used to identify proteins.

Identified proteins were further subjected to Gene Ontology (GO) based analysis to know the function. The connections between differently expressed proteins with each other and their connections with the other proteins were assessed by the STRING: EMBL (European Molecular Biology Laboratory) software\(^6\). Minimum required interaction score of medium confidence 0.400 was used.

Results

Fractionation of plasma samples. Using a systematic strategy (as shown in Fig. 1) pool plasma samples of AML and healthy subjects were first depleted for the top-7 abundant proteins through a MARS column. The resultant FPLC spectrum showed a clear separation of unbound and bound fractions, and was in the agreement with the figure provided by the manufacturer (Fig. S1). The resultant low-abundant proteins in flow through were then analysed by 1D SDS-PAGE for depletion efficiency which showed an effective protein depletion (Fig. S2). After depletion, the unbound portion was concentrated using 5 kDa MWCO tubes, followed by enrichment efficiency checking by 1D SDS-PAGE and found to be acceptable (Fig. S3).

The unbound portion, after enrichment by 5 kDa MWCO and protein precipitation, was further resolved by ZOOM-IEF over a pH range of 3.0 to 10 into five fractions of different pH ranges; pH: 3.0–4.6, pH: 4.6–5.4, pH: 5.4–6.2, pH: 6.2–7.0, and pH: 7.0–10.0 (Fig. S4). 1D SDS-PAGE analysis of all fractions showed that among all fractions two of pH: 5.4–6.2 and pH: 6.2–7.0 from AML and healthy samples pool had many protein bands with some differential pattern in the molecular weight range of 10–266 kDa. Therefore, these two fractions were subjected to further analysis. A comparative 1D SDS-PAGE picture of these two fractions from AML and healthy samples pool is shown in Fig. 2.
These two fractions were then mixed to make a single fraction of pH range 5.4 to 7.0. After protein precipitation, 2-DGE of this fraction was performed on IPG strip of pH range 4 to 7 followed by electrophoretic separation. Master gel image was created by combining the spots from both AML and healthy gels (Fig. 3a). That was used for comparison of AML gel with healthy gel image (Fig. 3a and b). The comparison between healthy and AML samples is shown in the scatter plot (Fig. S5). Out of 182 spots, 42 spots were having a 4X quantity difference, while 137 spots were with significance level ≥95%. The spots which were with 4X quantity difference and ≥95% significant both were 41. Altogether, 182 gel spots from the control and the AML samples, when analysed by MALDI-TOF MS, led to the identification of 34 distinct proteins and/or their respective isoforms and subunits (Fig. 4). The list of identified proteins with details such as theoretical and experimental pI and molecular weight,
MASCOT score, sequence coverage, etc. is shown in Table 1. A sample Mascot Score Histogram of hemopexin protein is given in supplementary section (Fig. S6).

Using PDQuest, we further analysed the 2-DGE spots to identify most significant and consistently dysregulated 15 proteins in AML cases (showing ≥4-fold increase/decrease in spot intensity and with significance level more than 95% in t-test), in comparison to the control subject. The results unequivocally showed variability in the levels of the SAA1, HPT, complement factor B, CD5 antigen-like, kininogen-1, fibrinogen gamma chain, C4b-binding protein alpha chain, complement factor 7, ApoA1, ApoE, plasminogen, apolipoprotein A-IV, prothrombin, fibronectin, and gelsolin. The former 6 proteins were found to be up-regulated, while the latter 9 were down-regulated in the AML, as shown in column graphs (Fig. S7).

**Gene ontology (GO) analysis.** Based on the known or postulated biological functions of the proteins as found in the GO consortium using homo sapiens taxon, the functions of the identified 34 proteins could be categorized on the basis of their functions as binding (28%), enzyme regulation activity (7%), homeostasis (7%), structural (2%), receptor mediated activity (15%), catalytic (14%), biological process regulation (25%) while for 2% proteins the functions are not yet known (Fig. S8).

**Validation through ELISA.** ApoA1 was found to be down-regulated in AML pool (10.67 mg/dL), in agreement with the spot intensity in the 2-DGE image of AML, as compared to the healthy pool (18.15 mg/dL). However, when the individual samples of AML were analysed, the mean plasma concentration of ApoA1 in the healthy group was 10.55 ± 4.6 mg/dL, whereas in the AML subjects it was found to be 14.59 ± 6.2 mg/dL i.e., higher, quite contrary to the expected results (Fig. 5). We applied t-test with Welch's correction (One-way ANOVA) to compare the variances. The mean concentrations were found to be significantly different with $p < 0.0396$, while the variances were not significantly different in individual samples.

SAA1 was found to be up-regulated in AML pool sample (1.15 mg/dL), in comparison to the pool sample of healthy individuals (0.89 mg/dL). Validation results of individual samples also showed significant differences when validated through ELISA. The mean plasma concentration of SAA1 in the individual samples of control was 0.69 ± 0.19 mg/dL whereas it was found to be 1.04 ± 0.38 mg/dL i.e., high in the AML subjects (Fig. 5). Mean concentrations were found to be significantly different with $p < 0.0029$, and the variances were also significantly different in individual samples with $p < 0.0162$ by applying statistical analysis.

![Figure 2](image-url)
Plasminogen was found to be down-regulated in AML patients in comparison to the healthy individuals through ELISA validation and found to agree with 2-DGE results. When pool samples of AML and healthy individuals were compared, the value of plasminogen was found to be low in AML (216.42 µg/mL), in comparison to healthy pool (244.87 µg/mL). Similarly, the mean plasma concentration of plasminogen in healthy and AML individuals was 269.8 ± 58.74 µg/mL, and 250.1 ± 69.31 µg/mL, respectively (Fig. 5). Statistical analysis showed significant difference of mean concentrations with p < 0.3844, and the variances with p < 0.5372.

Figure 3. Comparison of AML pool and healthy pool 2-DGE image. Highlighted spot numbers are those who are more than 95% significant and quantity changes more than 4-fold, which were further analysed. (a) Master gel, created by adding spots from AML pool and healthy pool gel images using PDQuest software. (b) 2-DGE map of healthy pool in the range of pH 5.4–7.0. (c) 2-DGE map of AML pool in the range of pH 5.4–7.0.

Figure 4. Identified variants and subunits of proteins in 2-DGE image through MALDI-MS and MASCOT database searching. [1, 39, 46: hemopexin, 2: alpha-1β-glycoprotein, 3, 20: kininogen-1, 4: Vit-D-binding protein, 5: transthyretin, 6, 9: apolipoprotein A-I, 7, 8, 52: ceruloplasmin, 10, 51, 55: complement factor H, 11: human serum amyloid-P component, 12, 13, 14, 32: apolipoprotein E, 15: apolipoprotein A-IV, 16, 35, 44: haptoglobin, 17, 53: alpha 2-macroglobulin, 18: fibronectin, 21: complement factor 7, 22: CD5 antigen-like, 23, 40: prothrombin, 24: complement C1r-subcomponent, 25: complement factor B, 26: complement C4-A, 27: alpha-1 microglobulin, 28, 29, 48: fibrinogen gamma chain, 30, 31, 47: fibrinogen beta chain, 33, 49, 50: plasminogen, 34, 41, 43: serum amyloid A-I, 36: retinol binding protein, 37, 39: C4b-binding protein alpha chain, 38: gelsolin, 39: human serum albumin, 42: serum amyloid A-IV, 43: haemoglobin β-component, 45: complement C4, 54: fibrinogen alpha chain].
In AML, complement factor 7 (C7) protein was found to be up-regulated in comparison to healthy individuals, contrary to 2-DGE results, where we found C7 was down-regulated in AML pool gel, in comparison to healthy pool gel. Although not a big difference, the complement C7 protein value in healthy pool was 94.68 ± 17.92 µg/mL versus 98.09 µg/mL in AML pool. Plasma mean value was 84.62 ± 17.92 µg/mL in healthy, while 104.6 ± 36.96 µg/mL in AML patients (Fig. 5).

T-test with Welch’s correction showed no significant difference between AML and healthy with p value 0.0537, while F-test to compare variances showed significant difference among study groups with p value 0.0090.

ApoE was found to be up-regulated in AML in contrast to 2-DGE results. ApoE value in healthy pool was 517.79 µg/mL versus AML pool was 564.82 µg/mL. Mean value in AML patients was found to be 504.4 ± 214.2 µg/mL, versus 720.2 ± 387.5 µg/mL in healthy group (Fig. 5). T-test with Welch’s correction gave no significant difference with p value 0.0532, and when variances were compared through F-test, a significant difference with p value 0.0300 was observed.

**Discussion**

The combination of 2-DGE and mass spectrometry offers a powerful tool to investigate the proteomic expression profiles to identify biomarkers which might serve as indicators of the disease. Recently many potential protein biomarkers have been reported in the literature for the diagnosis of AML e.g. ApoE, complement factor H, HPT, apolipoprotein A-N, SAA1, and gelsolin using proteomics techniques. Also various MS based studies have already been done on cell lines. In this study, we have investigated differential proteomic profile pattern among AML and healthy samples in plasma by employing multi-dimensional fractionation strategy to identify biomarker proteins.
The major bottleneck in analysing plasma proteins is to analyse low abundant proteins in the presence of high abundant proteins\(^3\). Therefore, to overcome this difficulty, we first depleted the samples for high abundant proteins to unmask the proteins which are present in very low amount. The depletion column with antibodies for the top 7 abundant proteins, including albumin, fibrinogen and HPT were used. While doing depletion, we have noticed that 100% depletion of plasma samples was not achieved, because in flow-through (unbound fraction), we have also identified peaks of albumin, all three chains of fibrinogen (\(\alpha\), \(\beta\) and \(\gamma\)), and HPT (Fig. 4). In terms of specificity of immunodepletion, contaminants in bound and eluted portion have been already reported by

Figure 5. Scatter plot with standard deviation achieved through ELISA results of individual samples of healthy and AML subjects. (a) ApoA1 protein, (b) SAA1 protein, (c) plasminogen, (d) complement factor C7, (e) apolipoprotein E.
manufacturers and researchers. Albumin may be present due to non-specific interactions with other proteins in the plasma, which is called the "sponge-effect". HPT and three chains of fibrinogen were visible in unbound fraction possibly due to the narrow-range techniques applied, or the fragments may have low binding efficiency to the column. The presence of target proteins (to be depleted through column) as contaminants have already been reported.

After depletion, many low abundant proteins were visualized by IEF. In-solution IEF eases the detection of low abundance proteins and increases the detection range as large amounts of proteins of specific pH can be loaded on the gel. In 2-DGE maps, most of the identified proteins were represented by multiple spots in AML and control group (Fig. 4). Slight to moderate shifts in pI or mass were between theoretical and experimentally-calculated values has already been reported in most of the cases. Both observations i.e., representation of single protein by multiple spots and variations in theoretical and experimentally calculated pI/MW, seem to be the result of post-translational modifications, especially glycosylation, affecting the electrophoretic mobility of the proteins, as reported earlier.

To gain an insight of the biological functions and interactive links that are known to be associated with the differentially expressed proteins in the dataset, STRING software program was used. Interactive links between 15 proteins of the dataset could be traced as presented in Fig. 6. Blue lines show phylogenetic co-occurrence between proteins, whereas light blue lines represent database evidence. The line thickness is a rough indicator of the power of the association. The visualizations among protein nodes show the predicted association between the proteins detected in the samples of AML and non-leukemic healthy patients. The pathways involved by these differentially regulated proteins are presented in Table S3.

We further validated some of the deregulated proteins through ELISA, as it has the potential to accelerate validation of protein biomarkers for clinical use. HPT and all three chains of fibrinogen were not included for the validation because depletion column had affinity for HPT and fibrinogen, and most of them had depleted out of plasma sample in column, so these results seen in 2-DGE was not reliable.

SAA1 is a major acute phase reactant and is also found as apolipoprotein of the HDL complex. ELISA validation results of SAA1 show similar trend as 2-DGE results, and hence SAA1 may be used as a potential diagnostic biomarker for AML (Fig. 5). Plasminogen; Plasmin dissolves the fibrin of blood clots and acts as a proteolytic factor in a variety of other processes including, embryonic development, tissue remodelling, tumour invasion, and inflammation. Plasminogen was found to be down-regulated in ELISA validation.

ApoA1; participates in the reverse transport of cholesterol from tissues to the liver for excretion by promoting cholesterol efflux from tissues and by acting as a cofactor for the lecithin cholesterol acyltransferase (LCAT), and as part of the SPAP complex activates spermatozoa motility. The results from the analysis of ApoA1 protein from

Figure 6. Curated pathway of fifteen differentially expressed proteins in acute myeloid leukaemia acquired from online STRING database. Balls with structures show that their 3D structures are also available in database.
individual samples were in contradiction from the 2-DGE maps (Fig. 5). Therefore, these results of ApoA1 protein require further validation on large number of individual samples.

Complement component 7; constituent of the membrane attack complex (MAC) plays a key role in the innate and adaptive immune response by forming pores in the plasma membrane of target cells. ApoE mediates the binding, internalization, and catabolism of lipoprotein particles. It serves as a ligand for the LDL (ApoB/E) receptor, and for the specific ApoE receptor (chylomicron remnant) of hepatic tissues. Results of both complement C7 and ApoE were not in agreement with 2-DGE results.

During ELISA validation, we found that out of five differentiating proteins, only two proteins SAA1 and plasminogen showed potential of differentiation of AML from healthy group, during ELISA validation. These two proteins showed links to 4 Kyoto Encyclopedia of Genes and Genomes (KEGG) pathways including transcriptional misregulation in cancers, p53 signalling pathway, proteoglycans in cancers, and hypoxia-inducible factor 1 (HIF-1) signalling pathway (Fig. 7). These pathways regulate cell proliferation and apoptosis, inflammation, metabolism, angiogenesis, cell growth and migration, cell migration and metastasis, cell cycle arrest, cellular senescence, and DNA repair, which are properties of cancerous cell. Therefore, the relevant up-regulation of SAA1 and down-regulation of plasminogen may be due to the direct effect of these disturbed signalling pathways, and can be used for the diagnosis of acute myeloid leukaemia in future.

**Conclusion**

In conclusion, 5D proteomic strategy using immunodepletion, 2-DGE, ZOOM-IEF and MALDI-MS, and ELISA analysis has shown a promising approach for the detection of differentiated proteins in AML in comparison with the control. Fifteen proteins were found to be deregulated in comparison to healthy control. Some of these deregulated proteins were further validated through ELISA technique and the results suggest that SAA1 and plasminogen can be used as biomarkers for the diagnosis of AML patients. However, as the validation was performed on a small number of proteins, therefore validation of these deregulated proteins on a larger number of individual subjects is needed.

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Acknowledgements
The study was financially supported by the Higher Education Commission (No. 4493), Pakistan. Our special thanks to all AML patients and healthy participants, who voluntarily gave their blood samples for this research project.

Author Contributions
S.K.R. was involved in experimental work and manuscript writing. M.S. helped in the experiment. S.G.M. contributed to the study design, conceptual and technical guidance along with the laboratory equipment and expertise. T.S. provided the disease samples. A.R. and M.I.C. were involved in manuscript checking.

Additional Information
Supplementary information accompanies this paper at https://doi.org/10.1038/s41598-017-16699-2.

Competing Interests: The authors declare that they have no competing interests.

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