Identification of plasma Complement C3 as a potential biomarker for neuroblastoma using a quantitative proteomic approach

Patrick Y. Kim, Owen Tan, Sonya M. Diakiw, Daniel Carter, Eric O. Sekerey, Valerie C. Wasinger, Tao Liu, Maria Kavallaris, Murray D. Norris, Michelle Haber, Lou Chesler, Alla Dolnikov, Toby N. Trahair, Nai-Kong Cheung, Glenn M. Marshall, Belamy B. Cheung

Children’s Cancer Institute Australia for Medical Research, Randwick, NSW 2031, Australia
Bioanalytical Mass Spectrometry Facility, Mark Wainwright Analytical Centre, University of New South Wales, Sydney, NSW, Australia
Division of Cancer Biology, Institute for Cancer Research, Sutton, Surrey, UK
Kids Cancer Centre, Sydney Children’s Hospital, Randwick, NSW 2031, Australia
Department of Pediatrics, Memorial Sloan-Kettering Cancer Center, 1275 York Avenue, New York, NY, United States

ARTICLE INFO

ABSTRACT

The majority of patients diagnosed with neuroblastoma present with aggressive disease. Improved detection of neuroblastoma cancer cells following initial therapy may help in stratifying patient outcome and monitoring for relapse. To identify potential plasma biomarkers, we utilised a liquid chromatography–tandem mass spectrometry-based proteomics approach to detect differentially-expressed proteins in serum from TH-MYCN mice. TH-MYCN mice carry multiple copies of the human MYCN oncogene in the germline and homozygous mice for the transgene develop neuroblastoma in a manner resembling the human disease. The abundance of plasma proteins was measured over the course of disease initiation and progression. A list of 86 candidate plasma biomarkers was generated. Pathway analysis identified significant association of these proteins with genes involved in the complement system. One candidate, complement C3 protein, was significantly enriched in the plasma of TH-MYCN+/+ mice at both 4 and 6 weeks of age, and was found to be elevated in a cohort of human neuroblastoma plasma samples, compared to healthy subjects. In conclusion, we have demonstrated the suitability of the TH-MYCN+/+ mouse model of neuroblastoma for identification of novel disease biomarkers in humans, and have identified Complement C3 as a candidate plasma biomarker for measuring disease state in neuroblastoma patients.

Biological significance
This study has utilised a unique murine model which develops neuroblastoma tumours that are biologically indistinguishable from human neuroblastoma. This animal model has
effectively allowed the identification of plasma proteins which may serve as potential biomarkers of neuroblastoma. Furthermore, the label-free ion count quantitation technique which was used displays significant benefits as it is less labour intensive, feasible and accurate. We have been able to successfully validate this approach by confirming the differential abundance of two different plasma proteins. In addition, we have been able to confirm that the candidate biomarker Complement C3, is more abundant in the plasma of human neuroblastoma patient plasma samples when compared to healthy counterparts. Overall we have demonstrated that this approach can be potentially useful in the identification of biomarker candidates, and that further validation of the candidates may lead to the discovery of novel, clinically useful diagnostic tools in the detection of sub-clinical neuroblastoma.

© 2013 The Authors. Published by Elsevier B.V. Open access under CC BY license.
Sympathetic ganglia or tumours were dissected from age matched WT and TH-MYCN<sup>+</sup> mice respectively for later analysis by ELISA. Upon dissection, the isolated tissue was washed and cleaned of non-specific tissue in Hank's Balanced Salt Solution and snap-frozen for storage. The sympathetic ganglia (pooled superior cervical ganglia and coeliac ganglia) were isolated from WT mice for comparison with early and advanced-stage tumours from 4 week old and 6 week old TH-MYCN<sup>+</sup> mice respectively. For each age/genotype, ganglia or tumours were isolated from 4 different mice and later purified as protein lysates for ELISA.

Human plasma samples from healthy volunteers were obtained from the Sydney Red Cross (N = 4) and Sydney Cord Blood Bank (N = 3) under the ethics approval (#08/148) from the South Eastern Sydney Illawarra. The samples obtained from the Red Cross were from healthy adults and the samples from the Cord Blood Bank were from healthy newborns. These samples were used as a normal healthy control for this project. The plasma samples from neuroblastoma patients were collected by Dr Nai-Kong Cheung from the Memorial Sloan-Kettering Cancer Centre (New York, USA), under the clinical trial study, Molecular Characterisation of Neuroblastic Tumour: Correlation with Clinical Outcome (Clinical Trials identifier: NCT00588068) and with the IRB protocol number #00-109. These samples were from patients with MYCN-amplified tumours (n = 9), which were all stage VI and MYCN-non-amplified tumours (n = 6), where four of these were from stage I and one each from stages II and III. The average age of the patients from which the samples were collected from was 3.7 years and all of these samples were collected at time of diagnosis.

2.2 Sample, fractionation, purification and analysis by LC–MS/MS

Aliquots (200 μg plasma proteins for each sample) were pooled and an appropriate volume of protease inhibitor was added (Complete™, mini, EDTA-free, protease inhibitor tablet from Roche Diagnostics, Manheim, Germany) as per manufacturer recommendations. An in-solution electrophoretic device called ProteomeSep (NuSep, NSW, Australia) was used to analyse the plasma from each genotype and each time-point. Individual plasma samples were also stored for further analysis with 2D gel electrophoresis and ELISA. The plasma proteins were partitioned into five fractions based on protein mass, pl and surface charge using the ProteomeSep, a preparative electrophoresis instrument which causes charged molecules to migrate across hydro gel membranes into different molecular weight fractions when a voltage is applied. ProteomeSep fractionations were performed using 5-chamber assembly, where chamber boundaries were defined by 2.7 x 2.2 cm polyacrylamide membranes of different restrictive pore sizes. All buffers described for sample preparations were also used as circulating buffers. Plasma was diluted 1:1 to give a final concentration of 14 μg/ml plasma in 45 mM Tris/5 mM ε-aminoacproic acid (EACA) buffer in 2 M urea, pH 10.2. All fractionations were performed at 50 V for 30 min and then 250 V for 2 h at 15 °C. The fractions were recovered from the chambers using gel-loading tips. The fractions collected were: 1-5 kDa, 5-25 kDa, 25-50 kDa, 50-75 kDa and >75 kDa. The two low molecular mass fractions, 1-5 kDa and 5-25 kDa fractions were desalted using C18 Stage Tips (Proxeon, Odense, Denmark) as per manufacturer’s instructions.

The peptides were resuspended in 5 μl of buffer A 2% (v/v) acetic acid, 0.1% (v/v) FA and separated by nano-LC using an Ultimate 3000 HPLC and autosampler system (Dionex, Netherlands) and then concentrated and desalted onto a micro C18 pre-column (500 μm x 2 mm, Michrom Biosources, USA) with 0.05% (v/v) heptfluorobutryric acid (HFBA) at 20 μl/min. After a 4 min washing, the pre-column was automatically switched (Valco 10 port valve, Houston, Texas, USA) into line with a fritless nano column, as previously published [17]. The LTQ-FT (Thermo Scientific, San Jose, CA, USA) was operated as described previously [17]. Peak lists were generated using ‘Mascot Daemon/extract_msn’ (Matrix Science), default parameters and then submitted to the protein database search programme Mascot (Matrix Science, Boston, USA).

2.3 Label-free quantification of identified peptides using ion count

All LC-MS/MS spectra were searched against the mammalian non-redundant NCBI database (2010) using Mascot (www.matrixscience.com). In solution digests analysed on the LTQ-FT Ultra were searched with the following criteria: trypsin digestion, precursor and product ion tolerances ±6 ppm and ±0.6 Da, respectively; variable modifications of methionine oxidation, phosphorylation (S, Y, T), carbamidomethyl and acrylamide; and one missed cleavage allowed. Identifications were accepted based on the MOWSE scores with a score > 44 indicating significant homology (p < 0.05).

Acquired spectra run in triplicate for each time point and in each fraction files were loaded into the Progenesis LC-MS/MS software (version 2.5, Nonlinear) for label-free quantification. The datasets generated by the MS analysis of plasma proteins from 2, 4 and 6 week old TH-MYCN<sup>+</sup> and WT mice were transformed into peak lists representing a 2D map using retention time and m/z. Each size fraction from each time-point was analysed separately and aligned according to retention time to create a 2D LC feature map. Features selected for inclusion were charge state inclusive of MH2<sup>+</sup> to MH3<sup>+</sup>, retention time from 13 min, and non-matching features were filtered out from further analysis. The samples were then allocated to their experimental group, including positive group (TH-MYCN<sup>+</sup>) and negative group (WT). Statistical parameters (p < 0.05 and fold-change > 5) were added to the sample analysis criteria to identify significant differences in peptide quantification. Peptides differing in either TH-MYCN<sup>+</sup> or WT mice were followed closely during the progression of disease. Parent ion information that was used to quantitate the peptides, and then the daughter ions were used to search Mascot, after which the identifications imported and linked back to each parent ion “feature”.

2.4 Database search and protein identification

All LC-MS/MS spectra were searched using Mascot against the mammalian non-redundant NCBI database with the following criteria: the precursor tolerance and the product ion tolerances were at 6 ppm and ±0.6 Da respectively; variable modification of methionine oxidation and trypsin digest. Mascot generated Dat.files were also then accumulated in
Progenesis LC-MS/MS software to compare and assess for differential abundance based on ion counts.

2.5. Removal of plasma abundant proteins for validation assays

Mouse abundant plasma proteins were depleted from each sample using the ProteoSpin™ abundant plasma protein depletion kit (Norgen Biotek Corp., Ontario, Canada), according to the manufacturer’s protocol.

Eluted protein samples were measured for protein concentration using the BCA protein assay (Thermo Scientific) and then stored at −20 °C for further downstream experiments.

2.6. Two dimensional analysis of protein candidates

Abundant protein depleted plasma samples from 6 week old TH-MYCN+/+ transgenic and WT mice were separated and analysed on 2D gels. IPG strips (11 cm, pH 4–7, Bio-Rad) were rehydrated with 250 μl rehydration buffer (7 M urea, 2 M thiourea, 4% (w/v) CHAPS, 2% (v/v) carrier ampholytes pH 3–10, 10 mM Tris, 43 mM DTT and 0.01% (v/v) bromophenol blue) for 16 h using a gel reswelling tray (GE Healthcare, Australia). Depleted plasma samples (200 μg) were mixed with rehydration buffer to a final volume of 200 μl and incubated at room temperature for 30 min with orbital mixing (1000 rpm), to solubilise and denature the plasma proteins. For, IEF, each sample was loaded onto the rehydrated IPG strips using sample cups, which were placed at the acidic end of the strip. IEF was then performed at room temperature for 120 kVh using the IPGphor electrophoresis unit (GE Healthcare). After IEF, the IPG strips were equilibrated (50 mM Tris–HCl pH 8.8, 6 M urea, 30% (v/v) glycerol, 2% (w/v) SDS, 0.01% (v/v) bromophenol blue with trace amounts of DTT) for 20 min on an orbital shaker at low speed. Each of the IPG strips were laid on top of 4–20% Criterion gels (Bio-Rad) and sealed using a 0.5% (w/v) agarose sealing solution (Tris/Glycine/SDS buffer, 0.5% (w/v) agarose and 0.01% (v/v) bromophenol blue) for electrophoresis in the second dimension. Electrophoresis was performed on the gel using the buffer system of Laemmli [18] at 100 V until the bromophenol blue dye front migrated off the gel. After electrophoresis, the proteins on the 2D gel were transferred on to nitrocellulose membrane (GE Healthcare) by wet Western transfer. ZAG protein was detected on the membrane with rabbit polyclonal anti-ZAG antibody (Sigma-Aldrich, MO, USA); transfer. ZAG protein was detected on the membrane with biotinylated antibody was then labelled with streptavidin-HRP (Dako, K1016) for 45 min at room temperature. The sections were developed with 3, 3′-diaminobenzidine tetrahydrochloride (Dako, K3468) and counterstained with haematoxylin.

2.7. ELISA quantitation of protein candidates

Sandwich enzyme-linked immuno sorbent assay (ELISA) kits were used for the quantification of ZAG (Uscn Life Science Inc. Wuhan, China), complement C3, C-reactive protein (CRP) and Serum Amyloid A (SAA) (Genway Biotech Inc. CA). Concentrations of ZAG and complement C3, proteins in the plasma of TH-MYCN+/+ transgenic and WT mice as well as the human plasma samples were determined according to the manufacturer’s protocol. In brief, mouse plasma samples were diluted with dilution buffer, 1:1000 for ZAG and 1:50,000 for complement C3. Human plasma samples were diluted with dilution buffer, 1:30 for ZAG and 1:50,000 for complement C3. Plasma concentration levels of CRP and SAA were determined in the human samples using the aforementioned ELISA kits and by diluting the samples 1:200 for CRP and 1:50 for SAA. C3 protein extracted from mouse tissue was diluted at 1:500 and protein concentration was determined according to the manufacturer’s protocol.

2.8. Immunohistochemical analysis of TH-MYCN+/+ tissue samples

Paraffin-embedded section of TH-MYCN+/+ tissue at 6 weeks of age was harvested and fixed with formalin. Paraffin-embedded sections were dried in a 60 °C oven for 1 h. The tissue slides were then immersed in xylene to deparaffinise the section. Slides were then re-hydrated with 100%, 90%, and 70% ethanol sequentially. Antigen retrieval was done by immersing the slides in 0.01 M tri-sodium citrate buffer with 0.05% Tween-20, pH6, at 104 °C for 15 min. The endogenous peroxidases were inactivated by immersing slides in 3% hydrogen peroxide. 10% goat serum was used to block non-specific binding of immunoglobulin. The slides were incubated with rabbit anti-ZAG (Sigma Aldrich) at 1:150 dilution overnight at 4 °C. Rabbit IgG prepared to the same concentration was used as negative control. Secondary antibody used was goat anti-rabbit immunoglobulin/biotinylated (Dako, E0432) at 1 in 500 dilution for 1 h, room temperature. The biotinylated antibody was then labelled with streptavidin-HRP (Dako, K1016) for 45 min at room temperature. The sections were developed with 3, 3′-diaminobenzidine tetrahydrochloride (Dako, K3468) and counterstained with haematoxylin.

2.9. Bioinformatics, pathway analysis and statistical analysis

For relative protein expression analysis, fold change was calculated in TH-MYCN+/+ mice compared to WT littermates and log2 transformed for normalisation. Unsupervised hierarchichal clustering analysis was performed using Multi-Experiment Viewer (MEV–TM4 microarray analysis software suite) to generate evolutionary gene trees (Euclidean distance metric selection and average linkage clustering parameters) [19]. To identify functional protein groups and pathways the 86 candidate proteins were assessed using Gene Ontology (GO) terms via the Memorial Sloan-Kettering Cancer Centre database (http://cbio.mskcc.org/CancerGenes) and also the Broad Institute Molecular Signatures database (MSigDB — http://www.broadinstitute.org/ gsea/msigdb). Statistical significance was evaluated using the Mann–Whitney, unpaired, nonparametric t-test in GraphPad Prism 5 (p-values < 0.05 were considered statistically significant).

3. Results

3.1. Identification and quantification of differentially abundant plasma proteins from TH-MYCN+/+ and WT mice

The plasma collected from 10 TH-MYCN+/+ and 10 WT mice at 2, 4 and 6 weeks of age was pooled and prefractonated using the electrophoretic ProteomeSep device, to reduce the
complexity of the plasma proteome, prior to LC-MS/MS identification and quantification. Fig. 1A outlines the mass spectrometry and label-free approach designed for identification of differentially abundant candidate plasma proteins in TH-MYCN+/+ and WT mice and generation of the final list of candidate biomarkers for further analysis. Fig. 1B shows a representation of the peptides identified by LC-MS/MS separated by retention time and m/z. The datasets generated into peak lists and analysed by Progenesis LC-MS/MS identified > 200 proteins per size fraction and time-point to be differentially abundant in the plasma of TH-MYCN+/+ and WT mice (±5-fold relative peptide abundance and p < 0.05).

3.2. Profiling differentially abundant plasma proteins in TH-MYCN+/+ and WT mice

To validate the label free LC-MS/MS method we selected zinc-alpha2-glycoprotein (ZAG) as it displayed one of the highest concentrations in the plasma of TH-MYCN+/+ mice compared to controls at 6 weeks of age (408-fold higher concentration). For 2D Western immunoblot analysis, pooled plasma samples from ten TH-MYCN+/+ and ten WT mice were analysed following abundant plasma protein depletion. Three distinct isoforms of ZAG were detected in both the TH-MYCN+/+ and WT pooled samples (Fig. 2A), however, all three isoforms in the TH-MYCN+/+ sample were more intense than the isoforms in the WT sample. A similar finding was observed by 2D Western immunoblot analysis of plasma from 3 individual TH-MYCN+/+ and 3 individual WT mice (Fig. 2B). Importantly, plasma from 2 of the WT controls (B5767 and B5850) only displayed 2 of the 3 isoforms of ZAG, which may reflect different functional contributions of ZAG isoforms in the development of neuroblastoma [20]. Consistent with the label-free analysis and 2D Western immunoblot data, ZAG concentrations were also found to be significantly higher in the plasma of TH-MYCN+/+ mice using ELISA. ELISA quantitation of pooled plasma 6 week old mice (three pooled plasma sets, with each set containing ten mice), showed that the TH-MYCN+/+ mice had an average ZAG concentration of approximately 500 pg/ml compared to 120 pg/ml in the plasma of 10 WT mice (p-value: 0.0034) (Fig. 2C). Quantitation of ZAG in the sera from 4 individual mice showed median plasma concentrations of ZAG in TH-MYCN+/+ samples were approximately 300 pg/ml compared with 180 pg/ml for WT mice (p-value: 0.0003) (Fig. 2D). When plasma levels of ZAG were measured in human plasma samples, no significant difference could be detected between the average values of ZAG concentration in 7 normal healthy plasma samples compared with 14 neuroblastoma patient samples.

![Fig. 1](image1.png)

Fig. 1 – Overall study design implemented for neuroblastoma plasma protein biomarker identification, refinement and validation. (A) Plasma harvested from TH-MYCN+/+ and WT mice were pre-fractionated into five size fractions prior to LC-MS/MS and subsequent peptide quantification analysis was performed with ion count. A final list of 86 candidate proteins was generated based on differential abundance at more than one time-point, and a fold change of at least 5 at one time-point. (B) A representative of a reference runs selection for the peptides, which was separated by retention time and m/z ratio.
plasma samples (Fig. 2E). However, it should be noted that the ZAG concentration range in the healthy individual samples were in range of 120–276 pg/ml, whereas some of the neuroblastoma patient plasma samples exhibited a ZAG plasma concentration of more than 760 pg/ml. All concentrations were calculated using data extrapolated from a standard curve. To investigate whether ZAG was expressed in the neuroblastoma tumour, we performed immunohistochemical (IHC) analysis on tissue samples obtained from TH-MYCN+/– mice (Fig. 2F). Synthesis and secretion of ZAG has been reported to occur in both the gastrointestinal (GI) tract as well as the liver [21]. As indicated by the IHC staining of the
mouse tissue samples, the staining of ZAG can be observed in both the GI tract and the liver. However, a much lower level of ZAG expression could be detected in the neuroblastoma tissue samples, compared with the GI tract and the liver. This suggested that the greater abundance of ZAG in the plasma of the TH-MYCN+/− mice is not directly secreted from the tumour, but could be from another source that is affected by the initiation or the progression of neuroblastoma. Further studies will be required to determine the origin of ZAG in TH-MYCN+/− mice bearing neuroblastoma tumours.

3.3. Assessment of the top differentially abundant plasma proteins in TH-MYCN+/− and WT mice

From our initial analysis, we removed abundant proteins including albumin, immunoglobulins, transferrin, alpha-1-antitrypsin and haptoglobin. The list was further refined to include only proteins which were differentially expressed at more than one time point to generate a final list of 86 candidates. The top candidates were clustered into 4 main groups based on expression patterns over the time-course of the experiment in TH-MYCN+/− mice relative to WT mice (Fig. 3).

Two groups of proteins (group 2 and group 3) demonstrated higher average expression in plasma from TH-MYCN+/− mice compared to WT mice at the early 2 week time-point during tumour initiation. It is likely that these groups will contain suitable biomarkers for early detection of neuroblastoma at diagnosis or relapse. The average expression of group 1 by contrast was higher in TH-MYCN+/− mice at 4 weeks during early tumour formation, and expression of group 4 was high at both 4 weeks and 6 weeks suggesting proteins associated with early to advanced tumour formation. Interestingly, the majority of proteins identified have not previously been associated with neuroblastoma. We found a high proportion of proteins which are secreted extracellularly (12 biomarkers from the 86 total biomarkers), several of which are associated with other solid tumour types such as SerpinA6 (marker of resistance to therapy in breast cancer — [22]) and FN1 (protects against apoptosis in lung tumorigenesis — [23]).

Pathway analysis was performed on the 86 candidate protein set using MsigDB and the top 10 hits ranked on p-value are shown in Fig. 4A. The top pathway identified involves protein metabolism, specifically O-glycosylation, a post-translational modification which has known roles in the regulation of cellular nutrient metabolism, protein stability and ligand presentation for functional recognition [24]. Two additional pathways related to glycosylation and metabolism were identified in the top 50 pathway hits (ranked #10 and #16), and four of the 5 proteins in these data sets belonged to expression group 2 with high expression in TH-MYCN+/− mice at the early 2 week time-point (Fig. 4B). The second ranked pathway to be identified involves the complement and coagulation cascades, two linked proteolytic cascades with roles in inflammation, immunosuppression, and tumour development/metastasis [25]. A total of 5 protein sets were identified in the top 50 pathways relevant to the complement system, and 4 of the 5 proteins from our analysis in these data sets clustered in group 4 with enhanced abundance at 4 and 6 weeks in the TH-MYCN+/− (Fig. 4C). Interestingly, we found a total of 6 data sets related to liver development, hepatoblastoma, or liver cancer in our top 50 pathways (including two ranked within the top 10). These datasets comprised a total of 23 of our candidate proteins, with 10 of these clustered in expression group 4 (shown in Fig. 4D). The liver is a common site of metastasis for neuroblastoma [26], however our finding could also suggest some overlaps in the molecular mechanisms or the pathological processes of these two cancer types.

3.4. Complement C3 protein is more abundant in plasma samples from human neuroblastoma patients than healthy volunteers

Complement C3 protein (C3) was significantly elevated in TH-MYCN+/− mice compared to WT counterparts at both 4 and 6 weeks of age (140-fold and 50-fold higher respectively). The pathway analysis identified C3 as part of both the complement cascade and the liver cancer groups in our analysis (Fig. 4), and as C3 levels are elevated in other cancer types we chose to analyse it further in our mouse model and in human plasma samples. The proteomics data was validated using ELISA performed on pooled plasma from 10 mice of each genotype per time-point, which confirmed significantly higher C3 levels in TH-MYCN+/− mice at both 4 and 6 weeks of age (7-fold and 2-fold higher than in WT mice respectively, Fig. 5A). The pattern of expression of C3 was similar to the initial proteomics analysis, although the magnitude of difference in plasma protein concentration was reduced when detected using ELISA. To determine whether elevated plasma C3 was a consequence of higher C3 expression in tumour tissues, ELISA was used to detect C3 levels in ganglia from WT mice and in tumours from TH-MYCN+/− mice. C3 protein was significantly elevated in tumours from TH-MYCN+/− mice compared to WT counterparts at 6 weeks of age (Fig. 5B), however no difference was observed at the earlier 4 week time-point. Whilst this observation may reflect a reduced limit of detection for C3 protein using ELISA, it could alternatively indicate that accumulation of C3 in tumours is secondary to increased levels in plasma.

To determine if these experimental observations would translate to a similar finding in human disease, plasma samples obtained from healthy adults (N = 4) and newborns (N = 3) were compared with plasma from 15 patients with neuroblastoma at diagnosis for expression of C3 protein. The mean plasma level of C3 in healthy individuals was 20 ng/ml (range: 8–37 ng/ml) compared to 110 ng/ml (range: 73–176 ng/ml) in neuroblastoma patients (median age 3.7 years) as determined by ELISA (Fig. 5C). As C3 is an important component of the immune system, we also assessed levels of two other inflammation-associated proteins – acute phase response protein C-reactive protein (CRP) and Serum Amyloid A (SAA) – to determine if C3 levels in neuroblastoma plasma samples were non-specifically elevated due to an inflammatory response. We did not observe significant differences in plasma concentration of CRP or SAA between healthy individuals and neuroblastoma patients using ELISA (Supplementary Fig. 1), suggesting that elevated C3 may potentially be a tumour-specific biomarker. Further validation will be required in larger patient populations as well as age-matched controls to confirm the validity of C3 as a plasma biomarker of neuroblastoma. This study also demonstrates the feasibility of using the TH-MYCN+/− mouse model to identify
novel neuroblastoma biomarkers with relevance to human disease.

4. Discussion

Initiation of neuroblastoma is executed by a series of molecular events that is mediated by dysfunctional genes and proteins [1]. Postnatally persistent embryonal cells with tumorigenic capacity possess the capacity to resist cell death imposed during the final stages of neurodevelopment and undergo secondary changes that characterize later tumour promotion and progression [27]. At 2 weeks of age, \( TH-MYCN^{+/-} \) mice show the first signs of neuroblastoma tumour initiation with the appearance of neuroblast hyperplasia, which persists until tumour formation and progression [7]. Studies have shown that this aberrant survival is due to a diminished p53 response to conditions of cellular stress, such as serum starvation or withdrawal of trophic factor [28,29]. We hypothesised that identification of proteins differentially expressed at these early time-points, prior to the onset of overt neuroblastoma, may be clinically useful as indicators to predict the initiation and onset of neuroblastoma.

In this study, we employed a label-free approach based on a comparison of peptide intensities, and quantitative analysis by ion count, to assess differences in plasma protein concentrations from \( TH-MYCN^{+/-} \) and WT mice for the identification of candidate plasma protein biomarkers of neuroblastoma. This methodology has several advantages over traditional proteomic analysis techniques utilising 2D polyacrylamide gel electrophoresis (PAGE), including improved detection of low mass and basic proteins and enhanced reproducibility [30]. Using a combination of the label-free proteomics approach and a transgenic MYCN mouse model of neuroblastoma, we identified a list of 86 candidate protein biomarkers present in plasma with altered levels compared to WT mice. A total of 27 proteins from the candidate list were significantly higher in plasma of the \( TH-MYCN^{+/-} \) mice than in WT littermates at the early 2 week time-point, and 49 were higher at the 4 week
time-point, suggesting that these candidates should be considered as markers for early detection of neuroblastoma.

Cluster and pathway analyses identified a group of proteins involved in post-translational protein modification, specifically glycosylation, which demonstrated significantly higher expression in TH-MYCN+/+ mice at 2 weeks of age before steadily declining during disease progression. Aberrations in the glycosylation of proteins have been associated with almost all forms of human cancers [31]. Studies have also suggested that aberrations in protein glycosylation can result in the initiation of oncogenesis, or that oncogenic transformation can cause alterations in protein glycosylation [31]. Other studies have shown similarities in the glycosylation pattern of proteins in different human neuroblastoma. The glycopeptides extracted from neuroblastoma tumours exhibited a low percentage of fucose-containing glycosylation [32]. Thus, it appears that the potential biomarkers identified by our cluster and pathway analysis may have significant roles in neuroblastoma biology. Albeit, further functional studies of each candidate will be required to confirm this hypothesis.

Pathway analysis also identified the complement system as being significantly associated with our candidate protein list. From this data, C3 was further characterised as a plasma biomarker candidate for neuroblastoma in the TH-MYCN+/+ murine model and in primary human plasma samples from neuroblastoma patients. Activation of C3 is central to the three complement pathways, which collectively result in the inflammatory response and the elimination of self- and non-self-antigenic targets [33]. However, using assays that detect CRP and SAA plasma levels we have shown that elevated C3 levels are not due acute phase responses and are most likely linked to neuroblastoma tumour formation. C3 is elevated in sera from pancreatic adenocarcinoma patients compared to normal individuals [34], and high levels are also observed in serum from breast, colorectal and lung cancer patients [30]. According to The Royal College of Pathologists of Australasia manual, the normal range of C3 in human plasma is recorded to be 0.9 – 1.8 g/l. Complement proteins have a central role in innate immunity, where they destroy antibody-coated targets such as apoptotic cells [35]. Deregulation of these molecules has also been associated with autoimmune and non-immune diseases [36,37]. In the context of carcinogenesis, the complement C3 protein has been implicated in the production of VEGF and extracellular matrix reorganisation and disintegration which contributes to the tumourigenic hallmarks of angiogenesis, invasion and migration, respectively [38].

Complement C4-B and complement factor D, which are other members of the complement family were identified by ion count to be more abundant in the plasma of TH-MYCN+/+ compared to the WT mice at 4 and 6 weeks of age. Therefore, it is possible that as an immune response to the initiation and development of neuroblastoma, several members of the complement protein family have been found to be more abundant in the plasma of TH-MYCN+/+ mice, compared to their WT counterparts. It also should be noted that a recent study has suggested that complement protein members C3, C4 and C5a may aid tumour growth by a mechanism of immunosuppression [38,39]. Thus, the abundance of these complement proteins in the plasma of the TH-MYCN+/+ mice

Fig. 5 – Plasma concentration levels of complement C3 in the TH-MYCN+/+ mouse model and in human neuroblastoma patient plasma samples. (A) ELISA assays demonstrate a significant difference in the concentration of complement C3 in the pooled plasma of 4 week old TH-MYCN+/+ and WT mice. Asterisks (***') represent p-value < 0.001. (B) Proteins extracted from the neuroblastoma tumour and ganglia tissue from 4 and 6 week old TH-MYCN+/+ (N = 4/genotype/time-point) and WT mice were analysed with a complement C3 ELISA. Asterisk (*) represents p-value < 0.05. (C) ELISA assays were performed to detect C3 protein levels of plasma in a cohort of human neuroblastoma plasma samples (N = 15) and in plasma from healthy individuals (N = 3 from cord blood and N = 4 from peripheral blood). Asterisks (****') represent p-value < 0.0001.
may either be a marker of the immune response to the cancer or aid in the development and growth of neuroblastoma.

5. Conclusions

Early detection and disease recurrence monitoring are critical areas in cancer treatment in which specific biomarker panels are likely to be very important in these key areas. The MYCN transgene of the mouse model used in our studies accurately reflects the more clinically aggressive forms of neuroblastoma. The validity of our data is strengthened by the finding that ZAG and C3, chosen for validation, have been previously published as serum markers in cancer. Further characterisation of the candidate biomarkers in our neuroblastoma animal model, and in human plasma patient samples, will likely result in a panel of biomarkers, that could be utilised for early detection and prediction of relapse risk in childhood neuroblastoma.

Supplementary data to this article can be found online at http://dx.doi.org/10.1016/j.jprot.2013.10.032.

Acknowledgements

This research was supported by Australian Postgraduate Research Scholarship, University of NSW, Program Grants from the NHRMC Australia, Cancer Institute NSW, and Cancer Council NSW. The Children’s Cancer Institute Australia for Medical Research is affiliated with the University of NSW and Sydney Children’s Hospital.

REFERENCES


