Proteomic analysis of metabolic, cytoskeletal and stress response proteins in human heart failure

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Proteomic analysis of metabolic, cytoskeletal and stress response proteins in human heart failure

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Abstract

Human heart failure is a complex syndrome and a primary cause of morbidity and mortality in the world. However, the molecular pathways involved in the remodelling process are poorly understood. In this study, we performed exhaustive global proteomic surveys of cardiac ventricle isolated from failing and non-failing human hearts, and determined the regulatory pathway to uncover the mechanism underlying heart failure. Two-dimensional gel electrophoresis (2-DE) coupled with tandem mass spectrometry was used to identify differentially expressed proteins in specimens from failing (*n* = 9) and non-failing (*n* = 6) human hearts. A total of 25 proteins with at least 1.5-fold change in the failing heart were identified; 15 proteins were up-regulated and 10 proteins were down-regulated. The altered proteins belong to three broad functional categories: (i) metabolic [e.g. NADH dehydrogenase (ubiquinone), dihydrolipoamide dehydrogenase, and the cytochrome *c* oxidase subunit]; (ii) cytoskeletal [e.g. myosin light chain proteins, troponin I type 3 and transthyretin] and (iii) stress response [e.g. HSP27 and HSP20]. The marked differences in the expression of selected proteins, including HSP27 and HSP20, were further confirmed by Western blot. Thus, we carried out full-scale screening of the protein changes in human heart failure and profiled proteins that may be critical in cardiac dysfunction for future mapping.

Keywords: heart failure • proteomics • metabolism • cytoskeleton • heat shock protein

Introduction

Heart failure, a complex syndrome that leads to left ventricular systolic and diastolic dysfunction, is one of the main causes of morbidity and mortality in the world [1]. Despite considerable recent research effort regarding its genetic causes, heart failure prevention and treatment strategies still suffer significant limitations [2]. Proteins, the products of gene expression, are the essential
biological determinants of disease phenotype, and changes in endogenous and external stimulation result in changes in the proteomes of individual patients over time. It is the proteomics, not the genome, that reveal the details within a cell [3]. One goal of functional proteomics is to examine global protein expression to elucidate the functional role of proteins and provide new insights into cellular mechanisms. This establishes the link between a cellular/organ proteome and the genesis of a physiological/pathological phenotype, which presents the investigator with new opportunities.

The central goal of this study is to delineate changes in protein expression between failing and non-failing human hearts and define the regulatory pathway to uncover the mechanism. Two-dimensional gel electrophoresis (2-DE) profiling techniques, which promised to resolve several thousand proteins in a single myocardial tissue sample, were used to examine changes in protein expressions and to identify key molecular players involved during disease progression. At the same time, our establishment of the proteomic map of human myocardium further revealed and provided the fundamental basis for the mechanism of heart failure.

### Materials and methods

#### Patient samples

Tissues for proteomic analysis were collected from the left ventricular free wall which was obtained from explanted hearts of transplant recipients \((n = 9)\) with a diagnosis of dilated cardiomyopathy (DCM, LVEF \(\leq 35\%\)) and non-failing control hearts \((n = 6)\) which were obtained from unmatched donors whose hearts were not suitable to transplantation. The baseline characteristics of patients are displayed in Table 1. The hearts were arrested and transported in ice-cold, oxygenated cardioplegic solution. Once in the laboratory the tissue was flash frozen in liquid \(\mathrm{N}_2\) and stored at \(-80\,^\circ\mathrm{C}\). The study protocol was approved by the Medical Ethical Committee of the Chaoyang Hospital of Capital Medical University.

#### Whole protein extraction

Tissue was homogenized by a Polytron PT-MR2100 (Kinematica AG, Switzerland) in lysis buffer (7M urea, 2M thiourea, 4% CHAPS, 2% IPG buffer pH 4–7 linear, 40 mM Tris, 65 mM DTT). Homogenates were immediately placed on ice to inhibit proteolysis, and then centrifuged at 14,000 rpm at \(4\,^\circ\mathrm{C}\) for 30 min. The supernatant was collected and transferred to a new tube. The protein was stored in aliquots at \(-80\,^\circ\mathrm{C}\) until analysis. Protein samples were quantified using the Bradford protein quantitative method.

#### Two-dimensional gel electrophoresis

2-DE was performed according to the manufacturer’s instructions (GE Healthcare, Piscataway, NJ, USA). Briefly, each protein sample in the lysis buffer was diluted to 450 \(\mu\text{L}\) with rehydration solution (9M urea, 2% CHAPS, 30 mM DTT, 0.5% IPG buffer pH 4–7, 0.002% bromphenol blue). Immobiline DryStrip gels (pH 4–7, 24 cm; GE Healthcare) were rehydrated with 450 \(\mu\text{L}\) of mixture solution in 24-cm strip holders and electrofocused with an Ettan IPGphor Isoelectric Focusing System (GE Healthcare). The focusing protocol was performed as follows: 50 \(\mu\text{A/strip}\) at \(20\,^\circ\mathrm{C}\), 30 V for 6 hrs, 500 V for 1 hr, 1000 V for 3 hrs, 3000V for 2 hrs, 5000 V for 2 hrs and 10,000 V for 8.5 hrs. Equilibrated IPG strips were placed at self-cast large-format 13% gradient polyacrylamide gels (255 \(\times\) 196 \(\times\) 1 mm) and embedded with 0.5% agarose, 0.001% bromophenol blue in cathode-running buffer (192 mM glycine, 24.8 mM Tris, 0.1% SDS, pH 8.3). Proteins were separated in the second dimension by SDS-PAGE using Ettan DALT 12n electrophoresis system (Amersham LifeSciences, Arlington Heights, IL, USA) containing cathode-running buffer at a regulated temperature of \(16\,^\circ\mathrm{C}\) by a cooling unit. The gels were run at 2 \(\text{W/ gel}\) for the initial 30 m followed by 12 \(\text{W/ gel}\) until the bromophenol blue dye front reached the bottom of the gel (total duration 6 and 7 hrs). All samples were run in triplicate.

#### Protein visualization, imaging analysis and in-gel trypsin digestion

All protein spots in 2-D gels were visualized by Coomassie Blue R-350 staining (Amersham Biosciences). Other standard chemicals required were purchased from Sigma-Aldrich (St. Louis, MO, USA). Stained gels were scanned on an Image-Scanner (Amersham Biosciences), and images were analysed with ImageMaster 2D 7.0 software supplied by the manufacturer. A total of 45 gel images were used for analysis. On average, over 1000 spots were detected on each 2-D gel image. Among these, about 950 of the most reproducible spots were included in the data analysis. Protein spots separated on 2-D gels were quantitated in terms of their relative volume (spot volume/total spot volume). For comparison, one image was chosen and allocated as the reference gel to align all other images in the experiment. Following manual alignment, spot detection, background subtraction and normalization were performed. To normalize the expression data, the data from one sample were fixed and all other samples were calibrated to this reference. Only those spots with a twofold or greater changes in expression levels were considered significant and selected for spot picking, trypsin digestion and mass spectrometry analysis to identify their protein content.

Table 1 Clinical characteristics of patients with heart failure

<table>
<thead>
<tr>
<th>ID</th>
<th>Sex</th>
<th>Age</th>
<th>Disease</th>
<th>LVEF (%)</th>
<th>Drug therapy</th>
</tr>
</thead>
<tbody>
<tr>
<td>No. 1</td>
<td>M</td>
<td>33</td>
<td>DCM</td>
<td>14</td>
<td>DIG, ACEI</td>
</tr>
<tr>
<td>No. 2</td>
<td>F</td>
<td>45</td>
<td>DCM</td>
<td>35</td>
<td>ACEI, BB</td>
</tr>
<tr>
<td>No. 3</td>
<td>M</td>
<td>43</td>
<td>DCM</td>
<td>34</td>
<td>AMIO, ACEI</td>
</tr>
<tr>
<td>No. 4</td>
<td>M</td>
<td>39</td>
<td>DCM</td>
<td>35</td>
<td>BB, ACEI</td>
</tr>
<tr>
<td>No. 5</td>
<td>M</td>
<td>57</td>
<td>DCM</td>
<td>33.9</td>
<td>ACEI, AMIO</td>
</tr>
<tr>
<td>No. 6</td>
<td>M</td>
<td>38</td>
<td>DCM</td>
<td>31.7</td>
<td>DIG, ACEI</td>
</tr>
<tr>
<td>No. 7</td>
<td>M</td>
<td>48</td>
<td>DCM</td>
<td>30.4</td>
<td>ACEI, BB</td>
</tr>
<tr>
<td>No. 8</td>
<td>M</td>
<td>42</td>
<td>DCM</td>
<td>26.3</td>
<td>DIG, DOB</td>
</tr>
<tr>
<td>No. 9</td>
<td>F</td>
<td>50</td>
<td>DCM</td>
<td>35</td>
<td>ACEI, AMIO</td>
</tr>
</tbody>
</table>

F: female; M: male; LVEF: left ventricular ejection fraction measured prior to explant; DCM: dilated cardiomyopathy (pre-transplant diagnosis); Drug Therapy: DIG: digoxin; DOB: dobutamine; AMIO: amiodarone; ACEI: angiotensin-converting enzyme inhibitor (usually lisinopril); BB: \(\beta\)-adrenergic blocker (metoprolol or carvedilol).
Protein spots demonstrating different expression patterns compared with controls were excised from gels by hand, and processed for mass spectrometric analysis. Excised spots were digested (9 hrs) in situ with trypsin. Peptides were extracted by addition of a 50% acetonitrile, 5% trifluoroacetic acid (TFA) solution and extracted solutions were concentrated to 4 µl in a lyophilizer (LABCONCO Freezone Liter; Labconco Corporation, Kansas City, MO, USA). A protein-free gel piece was similarly processed and used as the control to identify autodigestion products derived from trypsin.

**Mass spectrometry analysis and database searching**

The digested samples were spotted onto a MALDI target with an equal volume of cyano-4-hydroxycinnamic acid (2 mg/ml; Bruker Daltonics, Bremen, Germany) saturated with 70% acetonitrile in 0.1% TFA. They were analysed by MALDI-TOF MS using an AutoFlex TOF/TOF II mass spectrometer (Bruker Daltonics). The instrument settings were positive ion reflection mode, an accelerating voltage of 20 kV, and 150-ns delayed extraction time. The spectrum masses were ranged from 800 to 4000 Da and acquired with laser shots at 100/spectrum. A Peptide Mixture-1 kit (Bruker Daltonics) was used for external calibration. The matrix and autolysis peaks of trypsin were used as internal standards for mass calibration.

Many of processed spectra from the AutoFlex MALDI-TOF/TOF II mass spectrometer were searched against the NCBlnonredundant protein database (updated on January 28, 2010), which contained 10,348,164 sequences. The search was restricted to Homo sapiens (human) as taxonomy, which contained 229,313 sequences. The other main search parameters for all peptide mass fingerprinting searches were as follows: type of search, peptide mass fingerprint; enzyme, trypsin; fixed modifications, carbamidomethylcysteine; variable modifications, oxidation (M); type of search, peptide mass fingerprint; enzyme, trypsin; fixed modifications, carbamidomethylcysteine; variable modifications, oxidation (M); tolerance index > 30. The known contaminant ions (human keratin and tryptic autodigest peptides) were excluded. For interpretation of the mass spectra, monoisotopic peptide masses were searched against Mascot (http://www.matrixscience.com/cgi/search_form.pl?FORMVER=2&SEARCH=PMF) for protein identity searching.

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**Ingenuity pathways analysis**

Ingenuity pathways analysis (IPA, Ingenuity Systems®, http://www.ingenuity.com) is a web-based application that enables the visualization, discovery and analysis of molecular interaction networks. The functional analysis identified the biological functions and/or diseases that were most significant to the data sets. Fisher’s exact test was performed to calculate a P value determining the probability that each biological function and/or disease assigned to the data set was due to chance alone. Networks of the altered proteins were algorithmically generated based on their connectivity.

**Western blot analysis**

Proteins were extracted from human hearts as described, and then Western blot analysis was performed. Protein expression was evaluated by incubating membranes in 1:1000 rabbit polyclonal antibody (Abcam, San Francisco, CA, USA), followed by incubation with a 1:5000 secondary antibody (anti-rabbit IgG, HRP linked). Proteins were detected using the Fuji 3000 Image Station system. Protein bands were quantitated by densitometry using the Image J software program.

**Gene expression microarray**

Total RNA was extracted from heart tissues using Trizol reagent (Invitrogen, Carlsbad, CA, USA), according to the manufacturer’s instructions. RNA samples were labelled using the Illumina labelling kit and hybridized on the Human WG-6v3 BeadChip Array. Scanning was performed with the Illumina’s iScan scanner. GenePix pro V6.0 was used to read the raw intensity of the image. Perfect match fluorescence intensities were background-corrected, mismatch-adjusted, normalized and summarized to yield log2-transformed gene expression data using the GCRMA algorithm. The threshold value we used to define up-regulated and down-regulated mRNAs was a fold change greater than 1.5, and P value < 0.05 calculated by t-test.

**Real-time reverse transcriptase-polymerase chain reaction (RT-PCR)**

Total RNA was extracted from tissues by using Trizol reagent (Invitrogen), and cdNA was synthesized with Superscript II (Invitrogen) in accordance with the manufacturer’s instructions. Selected mRNAs (HSP27 and HSP20) were quantified with TaqMan quantitative RT-PCR. Primers for HSP20: 5′-CGTGAGGTGTGCACGG-3′, 3′-GCCAGCTGGAACCTGC-5′, probe: FAM-ACGCCTGGGATGACGAG-BHQ1; Primers for HSP27: 5′-GATGAACGTGTCATCGAGGC-3′, 3′-CCATCTAGAGATTTGGATATGGT-5′, probe: FAM-TGTCCATGAGTGCAGTGCATCATTATC-BHQ1. The PCRs were carried out in a final volume of 20 µl using a 7500 Real-Time PCR System (Applied Biosystems). Reactions consisted of 1.33 µl cdNA, TaqMan Universal PCR Master mix (Applied Biosystems), 0.2 µM TaqMan Probe, 1.5 µM forward primer and 0.7 µM reverse primer. The PCRs were initiated with a 10-min. incubation at 95°C followed by 40 cycles of 95°C for 15 sec. and 60°C for 60 sec. The expression levels of mRNAs were normalized to internal control β-actin and were calculated utilizing the ∆∆Ct method.

**Statistical analysis**

In this study, more than 1.5-fold increase or decrease protein was cut-off to distinguish changes and analysis. Data are expressed as mean ± S.D. All statistical analyses were performed with SPSS version 13.0. Functional data were analysed using analysis of variance (with repeated measures and the Tukey test for pairwise comparisons) and the Student’s t-test (with Bonferroni correction), as appropriate. All statistical calculations utilized Microsoft Excel software. Statistical significance was set at P < 0.05.

**Results**

**Differential protein expression analysis of failing and non-failing human hearts**

A total of 45 2-D gels were used in the differential proteomic analysis. The protein profile between replicate gels and biological...
samples showed high similarities, and the spot pattern was observed to be comparable between two groups. Image analysis of 2-D gels from all samples detected a total of 1000 protein spots. A total of 25 protein spots showed at least 1.5-fold difference in the normal controls (non-failing human hearts) versus disease samples (Fig. 1). The total 25 protein spots (15 increased, 10 decreased; Tables 2 and 3) were also observed to have differentially expressed at least a 1.5-fold change in the three-way pairwise comparisons, which were obtained by liquid chromatography-mass spectrometry/mass spectrometry. Classification of these proteins using the LOCATE subcellular localization database [4] showed that the majority of differentially expressed proteins comprised macromolecular (40%), cytoskeletal (28%) and mitochondrial (28%) proteins (Fig. 2A). PANTHER classification [5] of proteins by biological process revealed that the majority of proteins were involved in protein binding (cell structure and muscle contraction), ion binding (calcium binding) and hydrolase activity (Fig. 2B). The spots of most interest were those observed to have expression-level changes in comparison of normal versus failing hearts. These different proteins participate in some important biological process categories, such as multi-cellular organismal processes, biological regulation, developmental processes, metabolic processes and response to stimulus (Fig. 2C). The distribution and function of these expressed proteins provide a link to molecular components and pathways that may be involved in the development of heart failure progression. The findings suggest that major protein changes in human heart failure pathogenesis can be grouped into three main categories: proteins involved in mitochondria and energy production, cytoskeletal proteins and binding proteins. Therefore, the pathology of the failing heart is characterized by reduced mitochondrial activity, disruption of energy production and loss of cardiomyocytic structural integrity.

The network of proteomics

To characterize the functions associated with the detected proteins, the protein list was uploaded into the IPA software server and analysed using the Core Analysis module per the manufacturer’s instructions (http://www.ingenuity.com). As seen in Figure 3, the proteins are most significantly related to cardiovascular diseases and organism injury and abnormalities, which suggests that the proteins detected in this study is mostly regulators/effectors of the progression of heart failure.

As shown in Figure 4, the proteins mentioned before constitute the main structure of the regulation network. They all participate in the process of heart failure connected by important modulators, such as NF-κB, Myc and P38-MAPK, in respective pathways [6–8]. Besides, several metabolism disease pathways, including Growth Hormone and PRDX6, were also highlighted [9]. Meanwhile, functional modules related to cell differentiation and proliferation, as well as apoptosis were centred in the interaction network including only experimentally validated targets. Moreover, some important modulators in immune/inflammatory response, cardiovascular development or angiogenesis, such as TTR, ANGPT2 and FABP can also be found in the interaction map [10–12]. Therefore, the altered expression of those key proteins may cause the acceleration or deference of the heart failure process.

Western blot analysis

Western blot analysis was used to confirm significant protein expression changes observed in the 2-D gel proteomic data. Immunoblotting for the 27-kD heat shock protein (HSP27) band confirmed a marked difference in protein expression in the context...
### Table 2  
Identification of up-regulated proteins (≥1.5-fold) in failing versus normal hearts

<table>
<thead>
<tr>
<th>Spot no.</th>
<th>Symbol</th>
<th>Gene name</th>
<th>Subcellular location</th>
<th>Molecular function</th>
<th>Biological process</th>
<th>Fold change</th>
</tr>
</thead>
<tbody>
<tr>
<td>2535</td>
<td>HSP20</td>
<td>Heat shock protein, α-crystallin-related, B6</td>
<td>Cytoplasm</td>
<td>Protein binding; protein homodimerization activity</td>
<td>Regulation of muscle contraction</td>
<td>4.65</td>
</tr>
<tr>
<td>93</td>
<td>COX5B</td>
<td>Cytochrome c oxidase subunit Vb</td>
<td>Mitochondrion</td>
<td>Cytochrome c oxidase activity; metal ion binding</td>
<td>Respiratory gaseous exchange</td>
<td>6.07</td>
</tr>
<tr>
<td>2537</td>
<td>CRYAB</td>
<td>Crystallin, αB</td>
<td>Cytoplasm/nucleus</td>
<td>Microtubule; cytoskeletal protein binding; protein homodimerization activity; unfolded protein binding</td>
<td>Glucose metabolic process; oxygen and reactive oxygen species metabolic process; muscle contraction; response to hydrogen peroxide</td>
<td>2/1</td>
</tr>
<tr>
<td>72</td>
<td>LGALS1</td>
<td>Lectin, galactoside-binding, soluble, 1</td>
<td>Extracellular space</td>
<td>Glycoprotein binding; signal transducer activity; protein binding; galactoside binding</td>
<td>Apoptosis; positive regulation of IκB kinase/NF-κB cascade; myoblast differentiation</td>
<td>2</td>
</tr>
<tr>
<td>126</td>
<td>MYL2</td>
<td>Myosin, light chain 2, regulatory, cardiac, slow</td>
<td>Cytoskeleton</td>
<td>Actin monomer binding; calcium ion binding; protein binding; structural constituent of muscle; myosin heavy chain binding</td>
<td>Regulation of striated muscle contraction; negative regulation of cell growth; cardiac myofibril assembly; ventricular cardiac muscle tissue morphogenesis; heart contraction</td>
<td>2.4</td>
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<tr>
<td>222</td>
<td>HSPB2</td>
<td>Heat shock 27 kD protein 2</td>
<td>Cytoplasm</td>
<td>Protein binding; enzyme activator activity</td>
<td>Response to unfolded protein; somatic muscle development</td>
<td>2</td>
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<tr>
<td>1037</td>
<td>MYL6</td>
<td>Myosin, light chain 6, alkali, smooth muscle and non-muscle</td>
<td>Cytoskeleton</td>
<td>Motor activity; calcium ion binding; structural constituent of muscle; actin-dependent ATPase activity</td>
<td>Muscle contraction; skeletal muscle tissue development; muscle filament sliding</td>
<td>3.07</td>
</tr>
<tr>
<td>874</td>
<td>DLD</td>
<td>Dihydrolipoamide dehydrogenase</td>
<td>Mitochondrion/cytoplasm</td>
<td>Dihydrolipoyl dehydrogenase activity; oxidoreductase activity; lipoamide binding; FAD binding; NAD or NADH binding</td>
<td>Acetyl-CoA biosynthetic process from pyruvate; mitochondrial electron transport, NADH to ubiquinone</td>
<td>1.5</td>
</tr>
<tr>
<td>1459</td>
<td>OXCT1</td>
<td>3-Oxoacid CoA transferase 1</td>
<td>Mitochondrion</td>
<td>Transferase activity; protein homodimerization activity</td>
<td>Metabolic process; cellular ketone body metabolic process; ketone body catabolic process</td>
<td>2.25</td>
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<tr>
<td>77</td>
<td>FABP4</td>
<td>Fatty acid binding protein 4, adipocyte</td>
<td>Cytoplasm/nucleus</td>
<td>Fatty acid binding; protein binding; transcription repressor activity</td>
<td>Negative regulation of protein kinase activity; fatty acid metabolic process</td>
<td>2.08</td>
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<tr>
<td>336</td>
<td>PRDX6</td>
<td>Peroxiredoxin 6</td>
<td>Cytoplasm/nucleus</td>
<td>Antioxidant activity; oxidoreductase activity; hydrolase activity</td>
<td>Response to oxidative stress; phospholipid catabolic process; lipid catabolic process</td>
<td>2</td>
</tr>
<tr>
<td>2534</td>
<td>HSPA2</td>
<td>Heat shock 70 kD protein 2</td>
<td>Cytoplasm/mitochondrion</td>
<td>Nucleotide binding; ATP binding; unfolded protein binding</td>
<td>Response to unfolded protein</td>
<td>2.53</td>
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<tr>
<td>354</td>
<td>TNNI3</td>
<td>Troponin 1 type 3 (cardiac)</td>
<td>Cytoplasm</td>
<td>Actin binding; calcium channel inhibitor activity; protein kinase binding; troponin C binding; troponin T binding; calcium-dependent protein binding</td>
<td>Regulation of systemic arterial blood pressure by ischaemic conditions; cellular calcium ion homeostasis; regulation of smooth muscle contraction; negative regulation of ATPase activity; cardiac muscle contraction</td>
<td>2</td>
</tr>
<tr>
<td>133</td>
<td>TTR</td>
<td>Transthyretin</td>
<td>Extracellular space</td>
<td>Transporter; hormone activity; protein binding</td>
<td>Regulation of systemic arterial blood pressure by ischaemic conditions; cellular calcium ion homeostasis; negative regulation of ATPase activity; cardiac muscle contraction</td>
<td>1.56</td>
</tr>
<tr>
<td>343</td>
<td>HSP27</td>
<td>Heat shock 27 kD protein 1</td>
<td>Cytoplasm</td>
<td>Protein binding; ubiquitin binding</td>
<td>Cellular component movement; response to unfolded protein; response to heat</td>
<td>2.1</td>
</tr>
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</table>
of heart failure. After normalization for loading differences, a four-fold change was detected ($P = 0.0012$; Fig. 5A and B). Immunoblotting for the 16-kD HSP20 band is also differently expressed in the context of heart failure; an approximately three-fold change was detected in the heart failure group compared with the non-failing group ($P = 0.0029$; Fig. 5C and D). Also, other six proteins were randomized selected for Western blot, including CRYAB, CKM, TNNI3, MYL2, COX5B and TTR. The results were coincident with proteomic analysis (Fig. 6).

### Table 3 Identification of down-regulated proteins (>=1.5-fold) in failing versus normal hearts

<table>
<thead>
<tr>
<th>Spot no.</th>
<th>Symbol</th>
<th>Entrez Gene name</th>
<th>Subcellular location</th>
<th>Molecular function</th>
<th>Biological process</th>
<th>Fold change</th>
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<tr>
<td>190</td>
<td>BCL2A1</td>
<td>BCL2-related protein A1</td>
<td>Cytoplasm</td>
<td>Protein binding</td>
<td>Regulation of apoptosis; anti-apoptosis</td>
<td>−2.3</td>
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<tr>
<td>730</td>
<td>CKM</td>
<td>Creatine kinase, muscle</td>
<td>Cytoplasm</td>
<td>Kinase</td>
<td>Creatine metabolic process; phospho-creatine biosynthetic process</td>
<td>−2</td>
</tr>
<tr>
<td>1317</td>
<td>IFIH1</td>
<td>Interferon induced with helicase C domain 1</td>
<td>Cytoplasm/nucleus</td>
<td>Nucleotide binding; helicase activity; protein binding; ATP binding; hydrolase activity</td>
<td>Regulation of apoptosis; innate immune response</td>
<td>−2</td>
</tr>
<tr>
<td>568</td>
<td>KRT9</td>
<td>Keratin 9</td>
<td>Cytoskeleton</td>
<td>Structural constituent of cytoskeleton; protein binding</td>
<td>Epidermis development; skin development; intermediate filament organization</td>
<td>−2.63</td>
</tr>
<tr>
<td>387</td>
<td>DCI</td>
<td>Dodecenoyl-Coenzyme A delta isomerase (3,2 trans-enoyl-Coenzyme A isomerase)</td>
<td>Mitochondrion</td>
<td>Isomerase activity</td>
<td>Lipid metabolic process; fatty acid metabolic process</td>
<td>−1.5</td>
</tr>
<tr>
<td>2536</td>
<td>NDUFS3</td>
<td>NADH dehydrogenase (ubiquinone) Fe-S protein 3, 30 kD (NADH-coenzyme Q reductase)</td>
<td>Mitochondrion</td>
<td>Protein binding; NADH dehydrogenase (ubiquinone) activity; electron carrier activity</td>
<td>Mitochondrial electron transport, NADH to ubiquinone; oxygen and reactive oxygen species metabolic process; transport; induction of apoptosis; electron transport chain; negative regulation of cell growth</td>
<td>−2</td>
</tr>
<tr>
<td>235</td>
<td>PRDX2</td>
<td>Peroxiredoxin 2</td>
<td>Cytoplasm</td>
<td>Antioxidant activity; oxidoreductase activity</td>
<td>Negative regulation of oxygen and reactive oxygen species metabolic process; negative regulation of lipopolysaccharide-mediated signaling pathway</td>
<td>−1.51</td>
</tr>
<tr>
<td>187</td>
<td>SRP19</td>
<td>Signal recognition particle 19 kD</td>
<td>Cytoplasm/nucleus</td>
<td>7S RNA binding</td>
<td>Cotranslational protein targeting to membrane; SRP-dependent cotranslational protein targeting to membrane</td>
<td>−2.36</td>
</tr>
<tr>
<td>296</td>
<td>MYL4</td>
<td>Myosin, light chain 4, alkali; atrial, embryonic</td>
<td>Cytoskeleton</td>
<td>Actin monomer binding; calcium ion binding; structural constituent of muscle; myosin II heavy chain binding; actin filament binding</td>
<td>Regulation of the force of heart contraction; muscle organ development; positive regulation of ATPase activity; cardiac muscle contraction</td>
<td>−1.5</td>
</tr>
<tr>
<td>310</td>
<td>UQCRFS1</td>
<td>Ubiquinol-cytochrome c reductase, Rieske iron-sulphur polypeptide 1</td>
<td>Mitochondrion</td>
<td>Oxidoreductase activity, metal ion binding; 2 iron, 2 sulphur cluster binding</td>
<td>Transport; electron transport chain; response to drug; oxidation reduction</td>
<td>−2.15</td>
</tr>
</tbody>
</table>

**Merging proteomics and gene expression signatures**

Integration of proteomics with gene expression profiling would be more informative than either platform alone in capturing differentially involved gene networks between heart functional subtypes. Hierarchical cluster analysis of 741 significantly differentially expressed genes successfully segregates heart failure disease from healthy controls (Fig. 7). To identify a core set of functionally relevant genes/proteins, we explored the overlap of the differentially
regulated gene/protein sets captured by gene expression (741 genes) and global proteomic surveys (25 proteins) and represented on both arrays. There was just one protein, HSPA2, presented in gene expression signature, suggesting that the regulation of the most of altered proteins occurred at the level of post-transcription (Table 4). Furthermore, we performed validation experiments to verify the expression data from the microarrays. There are no significant differences of the mRNA expression of HSP27 and HSP20, between failed and non-failed group by quantitative RT-PCR (Fig. 8).

Discussion

The strategies available to prevent or treat heart failure are ineffective, which is due to the lack of a fundamental understanding of the cause and pathophysiology of the disorder. This study provides several intriguing findings. First, we identified a novel protein pattern in the failed human hearts, which may become helpful for the diagnosis and prognosis of patients with heart failure. Furthermore, we evaluated the global protein expressions to elucidate the functional role of network proteins and provide new information for cellular mechanisms, complement to other investigations with cultured heart cells or animal studies [13–15]. In addition, a protein network was set up to show the relationship between pathological changes in protein expression and integrated interactions with other known functionally linked proteins that may precipitate heart failure, providing the basis for further research.

In this study, we have shown that 741 genes were dysregulated in failed hearts identified by mRNA microarray. Interestingly, dysregulated proteins identified by proteomics did not correlate well with dysregulated gene expression profile. Several possibilities may account for the discrepancy, for example, it could be due to the transcriptional efficiency, protein stability, post-transcriptional modulation and/or alteration of the ubiquitin–proteasome system [16–18]. It could also resulted from the time course differences in transcription and pathological processes, the dysregulated proteins in failed heart may be expressed at prior stage, suggesting it will be ideally to detect mRNA expression profile at early stage of heart failure in the future study.

The proteins expressed in failing hearts were first investigated over a decade ago in a mouse model using two-dimensional polyacrylamide gel electrophoresis (2-D PAGE) [13]. Other researchers have shown that main proteins that exhibited changes in expression in mouse heart failure models belong to specific functional categories, including myocardial proteins, signal transduction proteins, metabolism-related proteins, transcriptional and translational proteins, growth and proliferation proteins and proteins with unknown function [14–16]. Although several studies have identified specific pathways that may be involved in heart failure [17–18], few have taken a global approach to identify key components of heart disease pathogenesis, especially using proteins extracted from the failing human heart. Through proteomic
Fig. 3 Functional characterisation of proteins detected to display aberrant expression across the failing heart. The gene list was imported and analysed by the Core Analysis Module in IPA software to statistically determine the functions/pathways most strongly associated with the gene list.

Fig. 4 The network analysis of proteomics using IPA. The top list of several pathways by this enrichment analysis are highlighted, such as NF-κB, TGF-β and P38-MAPK signalling pathways demonstrate to play an important role in the mechanical signalling during heart failure. Besides, several metabolism disease pathways, including Growth Hormone and PRDX6 pathway, are also highlighted. Meanwhile, functional modules related to cell differentiation and proliferation, as well as apoptosis (caspases) are centred in the interaction network including only experimentally validated targets. Moreover, some important modulators in immune/inflammatory response, cardiovascular development or angiogenesis, including TTR, ANGPT2 and FABP can also be found in the interaction map. The arrows represent the regulation between two proteins.
analysis, we have carried out a full-scale screen of the protein changes in heart failure, which mainly refer to several functional protein groups: proteins involved in mitochondria and energy metabolism, proteins involved in cardiac muscle contraction and others.

Mitochondrial and energy metabolism proteins

The proteins located within the mitochondria or involved in processes of energy metabolism play an important role in the heart's metabolic function. The main way that myocytes compensate for decreasing mitochondrial function is by increasing the expression of glycolysis-related proteins [19]. Therefore, high rates of myocardial energy production are required to maintain the constant demand of the working heart for adenosine triphosphate (ATP) and key enzymes involved in the Krebs cycle [20].

A total of seven proteins involved in the tricarboxylic acid cycle and respiratory activities change. Pronounced changes in key enzymes that regulate oxidation processes have often been revealed in subunits of electron transport chain complexes [21, 22]. These structures are embedded in the inner mitochondrial membrane and facilitate the transfer of electrons, creating a proton gradient across the myocyte membrane, completing the generation of energy from adenosine diphosphate (ADP) to ATP. A recent study described a reduction in activity and expression during the development of cardiac hypertrophy in rats, supporting the relevance of mitochondrial membrane proteins as an early and persistent marker of oxidative stress-related mitochondrial alterations in the development of hypertrophy [23].

Our results (Tables 2 and 3) indicate that comparison of the proteome of the failing heart with that of the normal heart identified changes in expression of four protein subunits from the following four respiratory complexes: (i) Complex I: NADH dehydrogenase (ubiquinone) Fe-S protein 3; (ii) Complex II: dihydrolipoamide dehydrogenase; (iii) Complex III: cytochrome c oxidase subunit Vb and (iv) Complex IV: dodecenoyl-Coenzyme A delta isomerase (3,2 trans-enoyl-Coenzyme A isomerase). These data imply the membrane microdomains could be a platform to integrate chemiosmosis as well as oxidation/reduction reactions. Therefore, the increased proteins that are the pronounced enzymes in the respiratory process can be explained by an increased demand for energy by unsynchronized changes in dysfunctional myocytes [24]. This conclusion is in concordance with
studies of cardiac hypertrophy in non-failing hearts in pressure-overload-induced hypertrophy models [25].

**Cardiac muscle cell contraction**

Our study indicates proteins involved in cell structure and muscle contraction also represent a large proportion of the expression proteins in the failing human heart. Seven cytoskeletal proteins with marked increases in expression are grouped in this category, including myosin light chain proteins, troponin I type 3 and transthyretin. Myosin light chain proteins are essential components in generating actin-based contractile force, and alterations in the expression of these proteins could affect myocardial contractility [26, 27].

It has already been identified that the genetic ablation of the N-terminal myosin essential light chain extension induces reduced force per cross-sectional area of muscle and leads to age-dependent compensatory cardiac hypertrophy [28]. Specifically, changes in myosin light chain 2 protein expressions have been linked to contractile dysfunction in dilated cardiomyopathy [29], ischaemia/reperfusion injury [30] and heart failure [31]. The up-regulation of these proteins in the failing human heart may indicate compensatory mechanisms due to adaptive response to the decrease of cardiac actin. Moreover, changes in myosin light chain 2 expression observed in this study may represent a shift in phosphorylation or deamidation rather than an overall increase in expression. Recent research shows that 50% mutant myosin regulatory light chain causes mechanical dysfunction of muscle fibres by sufficiently perturbing the tension at subsaturating Ca\(^{2+}\) concentrations; calcium sensitivity and cooperativity of force generation of fibres may be directly linked to the mechanism for familial hypertrophic cardiomyopathy [32].

Another important protein is Troponin I isoform TNNI3 (TNNI3), a constituent protein of the troponin complex located on the thin filament of striated muscle. TNNI3 provides a calcium-sensitive switch for striated muscle contraction and has the ability to regulate systemic arterial blood pressure with respect to ischaemic conditions. TNNI3 takes part in the regulation of cardiac muscle contraction and ventricular cardiac muscle tissue.

Fig. 6 Confirmative Western blot assays for selected proteins. Sucrose gradient fractions were separated by 12% SDS-PAGE and transferred to the NC membrane, and hybridized with anti-CRYAB (A), anti-CKM (B), anti-TNNI3 (C), anti-TTR (D), anti-MYL-2 (E) and anti-COX5B (F) antibodies. The bar graph shows the relative spot densities. CRYAB: crystallin, αB; CKM: creatine kinase, muscle; TNNI3: troponin I type 3 (cardiac); TTR: transthyretin; MYL-2: myosin, light chain 2; COX5B: cytochrome c oxidase subunit Vb.
morphogenesis. It binds actin and inhibits actomyosin ATPase activity in the absence of activating levels of calcium [33]. Proteins related to changes in cardiac muscle contraction in response to cellular calcium ion homeostasis may be another important pathway in the development of heart failure.

Table 4 The expression levels of mRNAs of altered proteins by using microarray in failed group compared to control group

<table>
<thead>
<tr>
<th>Protein</th>
<th>Heart failure (n = 14)</th>
<th>Control (n = 8)</th>
<th>Fold change</th>
<th>P value*</th>
</tr>
</thead>
<tbody>
<tr>
<td>HSP20 (HSPB6)</td>
<td>5796.445</td>
<td>3005.389</td>
<td>1.928683</td>
<td>0.349</td>
</tr>
<tr>
<td>COX5B</td>
<td>8882.66</td>
<td>10863.76</td>
<td>0.817641</td>
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</tr>
<tr>
<td>CRYAB</td>
<td>20202.15</td>
<td>19992.46</td>
<td>1.010488</td>
<td>1</td>
</tr>
<tr>
<td>LGALS1</td>
<td>4213.337</td>
<td>4256.731</td>
<td>0.989805</td>
<td>1</td>
</tr>
<tr>
<td>MYL2</td>
<td>33942.31</td>
<td>36326.29</td>
<td>0.934373</td>
<td>1</td>
</tr>
<tr>
<td>HSPB2</td>
<td>646.0505</td>
<td>743.0538</td>
<td>0.869805</td>
<td>1</td>
</tr>
<tr>
<td>MYL6</td>
<td>5329.922</td>
<td>4815.298</td>
<td>1.106873</td>
<td>1</td>
</tr>
<tr>
<td>DLD</td>
<td>1058.641</td>
<td>922.835</td>
<td>1.147162</td>
<td>1</td>
</tr>
<tr>
<td>OXCT1</td>
<td>624.9647</td>
<td>436.0639</td>
<td>1.433195</td>
<td>1</td>
</tr>
<tr>
<td>FABP4</td>
<td>562.7787</td>
<td>395.8654</td>
<td>1.106873</td>
<td>1</td>
</tr>
<tr>
<td>PRX6</td>
<td>3072.933</td>
<td>2949.509</td>
<td>1.041846</td>
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<tr>
<td>HSPA2</td>
<td>396.8911</td>
<td>32.29702</td>
<td>4.776237</td>
<td>0.044</td>
</tr>
<tr>
<td>TNNI3</td>
<td>5090.322</td>
<td>5438.007</td>
<td>0.936064</td>
<td>1</td>
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<tr>
<td>TR</td>
<td>50.36996</td>
<td>40.90046</td>
<td>0.693634</td>
<td>1</td>
</tr>
<tr>
<td>HSP27 (HSPB1)</td>
<td>6059.404</td>
<td>5597.423</td>
<td>1.082535</td>
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</tr>
<tr>
<td>BCL2A1</td>
<td>2.169603</td>
<td>1.955042</td>
<td>1.109747</td>
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</tr>
<tr>
<td>CKM</td>
<td>15383.75</td>
<td>16266.23</td>
<td>0.945748</td>
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</tr>
<tr>
<td>IFIH1</td>
<td>85.44498</td>
<td>76.69858</td>
<td>1.114036</td>
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<tr>
<td>KRT9</td>
<td>25.62142</td>
<td>72.2167</td>
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</tr>
<tr>
<td>DDI</td>
<td>359.5333</td>
<td>464.1664</td>
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</tr>
<tr>
<td>NDUF3</td>
<td>2959.224</td>
<td>3386.83</td>
<td>0.873745</td>
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<tr>
<td>PRX2</td>
<td>1318.944</td>
<td>1411.478</td>
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<td>SRP19</td>
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<tr>
<td>UQCRFS1</td>
<td>9222.257</td>
<td>9154.988</td>
<td>1.007438</td>
<td>1</td>
</tr>
</tbody>
</table>

*P value was calculated by illumina custom error model implemented in GenomeStudio and corrected by Benjamini-Hochberg False Discovery Rate.
Our study investigates some highly expressed proteins that maintain both morphologic and functional integrity of cardiomyocytes and increase tolerance against various types of stress, including αB-crystallin, HSP27 and HSP20 [34–36].

HSPs fulfill a wide range of functions, including cytoprotection and the intracellular assembly, folding and translocation of oligomeric proteins, and also represent a rapid response to altered redox states [37]. Failure to increase the levels of these high molecular HSPs may represent a loss of compensatory responsiveness in the setting of heart failure [38]. The highly expressed small HSPs (smHSPs) may exert effects on the left ventricular remodelling process that are different from those of high molecular HSPs, and it may be important to investigate the expression levels of smHSPs to better understand the pathogenesis of heart failure [39]. αB-crystallin is an smHSP structurally related to other smHSPs including αA-crystallin, HSP27, HSP20, HSP22, myotonic dystrophy protein kinase binding protein and HSPB3 [40]. αB-crystallin has strong anti-apoptotic properties and, when fully induced, is the most abundant smHSP in the heart and may constitute as much as 5% of total cardiac myocyte protein. It has been proposed that chaperoning by αB-crystallin can stabilize myofilament proteins through selective interactions with actin, titin, nebulette and the intermediate filaments desmin and vimentin [41].

With the compromise of αB-crystallin function, abnormal desmin forms aberrant aggregates and disrupts the integrity of the desmin network, which causes a reduction in cardiac function [42, 43].

An interesting finding in our study is the down-regulation of BCL2A1 (often named bfl-1) in patients with heart failure. There is not much known about the role of BCL2A1 in cardiac function and the implication in heart failure. Like many members of BCL2 family, BCL2A1 contains both pro- and anti-apoptotic properties depending on its intracellular localization [44, 45]. In a mouse model that mimics the mitochondrial DNA mutation–induced dilated cardiomyopathy, it has been shown that BCL2A1 and HSP27 are up-regulated as a compensatory mechanism to protect against apoptosis [46]. Therefore, it may be possible that the low levels of BCL2A1 are associated with the increased cell death during the heart failure. More research needs to be done to fully understand the biological roles of BCL2A1 that influence the course and outcome of heart failure.

In this study, human heart tissues were used to find that major protein changes in heart failure pathogenesis can be grouped into proteins involved in mitochondria and energy production and cytoskeleton proteins. Further studies of these proteins and pathways during the end-stage heart failure are necessary to address their precise roles in the pathophysiology of heart failure and their potential as diagnostic tools or therapeutic targets. This study has raised numerous questions and opened several new avenues for the future study of this complex disease.

Acknowledgements

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Conflict of interest

The authors confirm that there are no conflicts of interest.

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