Chronic contractile dysfunction without hypertrophy does not provoke a compensatory transcriptional response in mouse hearts

Scot J. Matkovich
Washington University School of Medicine in St. Louis

David R. Grubb
Baker IDI Heart and Diabetes Institute, Melbourne

Julie R. McMullen
Baker IDI Heart and Diabetes Institute, Melbourne

Elizabeth A. Woodcock
Baker IDI Heart and Diabetes Institute, Melbourne

Follow this and additional works at: https://digitalcommons.wustl.edu/open_access_pubs

Recommended Citation
Matkovich, Scot J.; Grubb, David R.; McMullen, Julie R.; and Woodcock, Elizabeth A., "Chronic contractile dysfunction without hypertrophy does not provoke a compensatory transcriptional response in mouse hearts." PLoS One. 11,6. e0158317. (2016).
https://digitalcommons.wustl.edu/open_access_pubs/5160

This Open Access Publication is brought to you for free and open access by Digital Commons@Becker. It has been accepted for inclusion in Open Access Publications by an authorized administrator of Digital Commons@Becker. For more information, please contact vanam@wustl.edu.
Chronic Contractile Dysfunction without Hypertrophy Does Not Provoke a Compensatory Transcriptional Response in Mouse Hearts

Scot J. Matkovich¹*, David R. Grubb², Julie R. McMullen², Elizabeth A. Woodcock²

¹ Center for Pharmacogenomics, Department of Internal Medicine, Washington University School of Medicine, St. Louis, MO, United States of America, ² Baker IDI Heart and Diabetes Institute, Melbourne, Australia

* smatkov@dom.wustl.edu

Abstract

Diseased myocardium from humans and experimental animal models shows heightened expression and activity of a specific subtype of phospholipase C (PLC), the splice variant PLCβ1b. Previous studies from our group showed that increasing PLCβ1b expression in adult mouse hearts by viral transduction was sufficient to cause sustained contractile dysfunction of rapid onset, which was maintained indefinitely in the absence of other pathological changes in the myocardium. We hypothesized that impaired contractility alone would be sufficient to induce a compensatory transcriptional response. Unbiased, comprehensive mRNA-sequencing was performed on 6 biological replicates of rAAV6-treated blank, PLCβ1b and PLCβ1a (closely related but inactive splice variant) hearts 8 weeks after injection, when reduced contractility was manifest in PLCβ1b hearts without evidence of induced hypertrophy. Expression of PLCβ1b resulted in expression changes in only 9 genes at FDR<0.1 when compared with control and these genes appeared unrelated to contractility. Importantly, PLCβ1a caused similar mild expression changes to PLCβ1b, despite a complete lack of effect of this isoform on cardiac contractility. We conclude that contractile depression caused by PLCβ1b activation is largely independent of changes in the transcriptome, and thus that lowered contractility is not sufficient in itself to provoke measurable transcriptomic alterations. In addition, our data stress the importance of a stringent control group to filter out transcriptional changes unrelated to cardiac function.

Introduction

Heart failure, a condition wherein the pumping ability of the heart is severely compromised, develops as the end result of many different cardiac diseases. The prevalence of heart failure worldwide is increasing and it contributes substantially to the rising cost of health care. Despite this, current treatment options are largely limited to amelioration of symptoms. The failing
heart exhibits a number of characteristics that contribute to pump failure. Some of these involve non-myocytes, especially fibroblasts that are instrumental in remodelling the heart [1] but some changes occur within the myocytes themselves [2]. Myocytes from failing hearts exhibit structural changes as well as alterations in intracellular signaling pathways and calcium handling [3-5]. Importantly, the calcium content of the sarcoplasmic reticulum (SR) is lowered, partly due to heightened calcium leak into the cytosol mediated by ryanodine receptors and partly due to reduced SR calcium uptake via the sarcoendoplasmic reticulum calcium ATPase (SERCA) [2, 6]. Failing myocytes characteristically show lowered expression of the SR calcium pump SERCA2a encoded by \(Atp2a2\) [7] along with reduced phosphorylation of phospholamban (PLN), which further depresses SERCA2a activity [8]. PLN is phosphorylated by protein kinase A downstream of \(\beta\)-adrenergic receptor activation and also by calcium-calmodulin regulated kinase II (CaMKII\(\delta\)) at a neighbouring site [9]. These PLN phosphorylations disinhibit SERCA and enhance lusitropy; therefore the lowered \(\beta\)-receptor expression and activity that characterize the failing myocardium likely contribute to SR calcium depletion [10].

In addition to these considerations, recent studies in our laboratory have identified another factor that contributes to contractile depression in the failing hearts. Failed myocardium from humans and experimental animals showed heightened expression and activity of phospholipase C\(\beta\)1b (PLC\(\beta\)1b) [11], one of the splice variants of PLC\(\beta\)1 [12] that initiates signaling responses downstream of Gq-coupled receptor activation [13]. Our studies showed that increasing PLC\(\beta\)1b expression in adult mouse hearts by viral transduction was sufficient to cause rapidly developing, sustained contractile dysfunction [14], which lasted for several months in the absence of other pathological changes in the myocardium [14] (time course depicted diagrammatically in Fig 1).

Further evidence pointing to the significance of the contribution of heightened PLC\(\beta\)1b to pathology was provided by a more recent study [15], which showed that inhibiting PLC\(\beta\)1b selectively prevented or reversed contractile dysfunction following pressure overload induced by trans-aortic constriction. Along with the improved contractility, the PLC\(\beta\)1b-specific inhibitory mini-gene reduced hypertrophy and prevented premature death [15].

Chronically heightened PLC\(\beta\)1b expression results in compensatory hypertrophy and some fibrosis only after a prolonged period (36 weeks) [14]. In agreement with this, transcriptional activation of several ‘classical’ marker genes associated with hypertrophy and fibrosis was detected at 36 weeks, but not at earlier time points. In contrast to the signaling mechanisms responsible for the development of hypertrophy, it is not known whether transcriptional events can be initiated specifically by low contractility, either as causes or consequences of contractile dysfunction. PLC\(\beta\)1b-induced contractile dysfunction, prior to the onset of hypertrophy, provides the possibility of examining whether there are transcriptional responses associated with chronically depressed contractility in the absence of other changes related to the progression to heart failure. Importantly, transduction of the closely related splice variant PLC\(\beta\)1a had no effect on contractility [14], or on any other discernible response when expressed in mouse hearts at similar levels to PLC\(\beta\)1b [16] and thus PLC\(\beta\)1a provides a stringent control for all studies involving PLC\(\beta\)1b.

The literature contains many examples of transcriptome studies undertaken using hypertrophied or failed myocardium from humans and animal models [17-19]. Such studies have identified genes associated with cellular growth, fibrosis and remodelling as well as genes associated with calcium handling. Many of these identified genes could contribute to contractile depression, singly or in combination, but such studies raise the issue of which transcriptional changes are primarily causative and which are secondary to remodelling or to the loss of contractile function. Contractile dysfunction caused by PLC\(\beta\)1b does not induce other changes associated with the progression to failure until a much later time point, and thus provides a unique opportunity to examine transcriptional changes associated directly with low contractile function.
Such transcriptional changes could be either causative of dysfunction or secondary to the chronically low contractility. With this in mind, we undertook a comprehensive mRNA-sequencing study of mice with chronically low contractile function due to PLCβ1b overexpression, hypothesizing that reduced cardiac output alone would be sufficient to provoke at least a partly compensatory transcriptional response. We used a time point early in the response to avoid compensatory hypertrophy observed after several months of treatment [14] (Fig 1).

Methods
All experiments were conducted in accordance with the Australian Code of Practice for ‘The Care and Use of Animals for Scientific Purposes’ of the National Health and Medical Research Council.
Council of Australia and all studies were approved by the Alfred Medical Research and Education Precinct Animal Ethics Committee (#1362/2013). Mice used were C57BL/6 obtained from the Animal Resources Centre, Perth.

Expression of PLCβ1b or PLCβ1a in mouse hearts

rAAV6-PLCβ1b, rAAV6-PLCβ1a or rAAV6-blank (control virus with a promoter but no protein product [20]) were prepared as described previously [14]. Male mice, 8 weeks of age, were injected with rAAV6-FLAG-PLCβ1b, rAAV6-FLAG-PLCβ1a or rAAV6-blank intravenously (IV) via the tail vein at a vector dose of 3 x 10^{10} vg/g. This dose has been shown to produce heightened expression of FLAG-PLCβ1b or FLAG-PLCβ1a in all chambers of the heart in all animals [14]. Nine (9) weeks after viral delivery (8 weeks after PLCβ1b/PLCβ1a expression) the animals underwent echocardiographic studies to evaluate contractile function, followed by humane culling, removal of hearts and dissection of the left ventricle (LV) for RNA extraction and western blotting. Extracted RNA subsequently was used for mRNA sequencing (mRNA-Seq) analysis. Western blotting using anti-FLAG and anti-tubulin antibodies to confirm expression in all hearts used for sequencing was performed exactly as described previously (14).

Measurement of contractile function

In the current study, contractile function was assessed as ejection fraction determined by echocardiography. Anesthesia was maintained with 1.7% isoflurane. Echocardiography was performed using a Philips iE33 ultrasound machine and a 15 MHz linear-transducer. After a short-axis 2-D image of the LV at the level of the papillary muscles was obtained, 2-D guided M-mode images were acquired digitally at a sweep speed of 133 mm/s. Ejection fraction [EF = (LVIDd^3 – LVIDs^3)/LVIDd^3, where LVIDd/s is the left ventricular internal diameter at systole ’s’ or diastole ’d’].

RNA-Seq methods for AAV6-PLCβ1a/b cardiac analysis

Total RNA from mouse heart left ventricle was prepared by Trizol extraction from flash-frozen tissue. Six (6) biological replicates were used for each of rAAV6-blank, rAAV6-PLCβ1a and rAAV6-PLCβ1b conditions. Ribosomal RNA depletion along with RNA fragmentation and conversion to paired-end, strand-specific Illumina sequencing libraries were performed with Illumina TrueSeq preparation kits. Sequencing reads of 76 nt in length were obtained from multiplexed libraries on an Illumina NextSeq instrument at the Ramaciotti Centre for Genomics, Sydney, Australia with (3.02 ± 0.11) * 10^7 paired reads per heart (mean ± SEM). Reads were aligned to the mouse transcriptome represented in the Illumina iGenomes mm10 UCSC release using TopHat [21], achieving a read depth of (1.84 ± 0.07) * 10^7 paired reads per heart (mean ± SEM); this represents a mean alignment to the transcriptome of 61%. All mRNA-sequencing reads and transcriptome-aligned read counts have been deposited in the NCBI GEO with accession number GSE73909.

Statistical and informatic procedures

For ejection fraction measurements in Fig 1C, data were analyzed using a 2 way ANOVA for repeat measurements, as described previously [14]. For the newly performed RNAseq measurements in this study, the union mode of HTSeq [22] was used in a strand-specific manner to quantitate reads pertaining to each mRNA. These data were used as input to the DESeq package which adjusted for library depth, performed differential expression calculations, and derived false discovery rates (FDR). Similar findings were made when the HTSeq-quantitated reads were used as input to edgeR for calculating differential expression [23]. For previous
RNAseq studies [24–27] employing non-strand-specific mRNA reads, the union mode of HTSeq [22] was used to re-analyze raw RNAseq reads, newly aligned to the mm10 UCSC genome as above, without taking strand identity into account. DESeq was again employed for differential expression calculations.

Comparison of PLCβ1a / PLCβ1b RNA-Seq data with existing RNA-Seq data sets

Data sets for rAAV6-mediated adult cardiac expression of PLCβ1a and PLCβ1b were compared with existing RNA-Seq datasets of αMHC/Myh6-driven expression of Gaq [24–26] or of full-length PKCα cDNA in adult mouse hearts [27] commencing in the neonatal period. Although PKCα expression was under the control of the doxycycline-suppressible tet-off Myh6 promoter, doxycycline was never administered to dams or pups and thus transgene activity commenced at birth, as for Gaq. Only mRNAs with a cardiac abundance of FPKM ≥3 were included, in accordance with the design of the original studies. In total, 781 individual mRNAs exhibited significant difference between PLCβ1a/b and blank virus, between Gaq and wt controls (derived specifically from [24]), or between PKCα and tet-off (tTA; TO) controls at a false discovery rate (FDR)<0.1. The minimum fold-changes corresponding to these FDRs (up- or down-regulation) were 1.16-fold (16%) for PLCβ1a/b vs blank virus, and 1.2-fold (20%) for Gaq vs wt or PKCα vs tet-off. In order to compare expression changes for all 781 mRNAs across all three experiments, the fold-change of each mRNA between 'treatment' and appropriate 'control' groups was computed, the mean mRNA abundance value of control hearts was adjusted to 1 (log2 value = 0), and the resulting variation across individual hearts from each group was displayed in a heatmap (Partek Genomics Suite v6.6, Partek, St. Louis, MO).

Results

Functional and transcriptional responses to PLCβ1b or PLCβ1a expression

PLCβ1b or the splice variant that is functionally inactive in heart, PLCβ1a, were overexpressed in mouse hearts in vivo using rAAV6-mediated transduction (Fig 1A and 1B). Overexpression of PLCβ1b in mouse hearts for 8 weeks resulted in depressed contractile function, expressed as ejection fraction (EF%), without detectable hypertrophy. These data are similar to those reported previously and are typical of 5 different cohorts (Fig 1C, 1D and 1E; Table 1) [14]. In all cases, contractile dysfunction reached a plateau between 8–10 weeks after injection and was sustained for several months followed eventually by hypertrophy. For convenience, data from previous studies are depicted diagrammatically in Fig 1D, allowing the time frame of the current study to be seen in relation to phenotype progression. We hypothesized that low contractility and decreased cardiac output would be sufficient to trigger a compensatory transcriptional response. However, despite the low contractility in PLCβ1b-expressing mice, unbiased whole-transcriptome analysis identified only minimal differences between hearts from mice expressing PLCβ1b and hearts from mice transduced with a control vector that does not express a protein product (rAAV6-blank). Only nine (9) genes showed significant expression differences between these two groups at a false discovery rate (FDR) < 0.1 (which represents a rather lenient filter compared to typical FDRs < 0.05 or < 0.02) (Fig 2, S1 Table). Similar expression changes were observed when data from rAAV6-PLCβ1a expressing mice were compared with rAAV6-blank mice. A total of 15 genes showed a transcriptional response at FDR < 0.1 to either or both of PLCβ1b or PLCβ1a expression (Fig 2, S1 Table); these changes were all of relatively low magnitude and did not include any canonical genes known to be involved in the regulation of cardiac contractility or hypertrophy (Fig 3).
Since mRNA-sequencing studies utilizing genetically- and/or surgically-manipulated mouse hearts with similar numbers of biological replicates have typically observed hundreds of altered mRNAs [24–26, 28, 29], we evaluated the variance to read-count relationship for the mRNAs subjected to statistical comparison and found that variance declined for mRNAs with higher read counts in the expected manner (S1 Fig). This suggests that the extremely modest transcriptional response observed in rAAV6-PLCβ1a/b mouse hearts is not due to technical artefacts in the mRNA-sequencing procedure.

Comparison of PLCβ1b transcriptional responses to those resulting from neonatal induction of upstream and downstream signal transducers

Cardiac overexpression of either Gaq (upstream regulator of PLCβ1b [16]) or of PKCa (activated downstream of PLCβ1b [14]) beginning in the immediate post-natal period are known to induce widespread transcriptional responses, and in the case of Gaq a well-described phenotype of hypertrophy and dilation with severely blunted contractility occurs as a result [30–32]. As might be expected from the phenotypic outcomes, many more mRNAs are dysregulated in adult hearts overexpressing Gaq than PKCa, in which transgene expression (under the control of the αMHC / Myh6 promoter) began immediately after birth (see Methods) (Fig 4A) [24, 27]. We compared the transcriptomes of rAAV6-PLCβ1a and -PLCβ1b hearts (without filtering for statistical significance of differentially-expressed mRNAs) to published mRNA-seq profiles of adult hearts overexpressing Gaq or PKCa from the post-natal period [24, 27]. However, neither PLCβ1a nor PLCβ1b hearts showed any evidence of Gaq- or PKCa-mediated alterations in transcriptional profile (Fig 4B).

Discussion

Our previous studies showed that heightened expression of PLCβ1b is sufficient to cause chronic contractile dysfunction, whereas expressing the closely related splice variant PLCβ1a
had no discernible effect [14]. Furthermore, selectively inhibiting PLCβ1b activation prevented/reversed heart failure following pressure overload [15]. Hearts expressing PLCβ1b showed depressed contractile function without hypertrophy or fibrosis, at least until 36 weeks after virus delivery. This provided an opportunity to examine whether transcriptional responses occur in response to loss of contractility in isolation, without the difficulty of disentangling these from responses associated with hypertrophied or failed myocardium. PLCβ1b presented an especially attractive opportunity because PLCβ1a has no effect on contractility and could thus serve as a stringent control.

Despite chronically lowered contractility, we detected only minor changes in gene transcription associated with PLCβ1b expression and furthermore, similar changes followed expression of PLCβ1a, even though PLCβ1a did not alter contractile function. Thus, depressed contractility per se does not lead to widespread compensatory transcriptional changes. Our previous report noted changes in canonical transcriptional markers of cardiac hypertrophy, once hypertrophy finally ensued in PLCβ1b hearts, presumably as a result to prolonged contractile depression. After only
8 weeks of chronically lowered contractility, we found no changes in the expressions of ANP and α-skeletal actin, collagen-1A1, -3A1, α- or β-myosin heavy chain or phospholamban as measured by RT-qPCR [14], nor did we in the current RNA-sequencing study. After 32 weeks of depressed contractility, PLCβ1b-expressing hearts showed a modest degree of hypertrophy. At this time point, increased expression of ANP and α-skeletal actin was observed in PLCβ1b-expressing hearts compared with either PLCβ1a-expressing hearts or control hearts [14].

The finding that PLCβ1b-induced contractile dysfunction does not involve substantial transcriptional changes is compatible with our previous data, which showed that the contractile deficit could be fully reversed by treatment with a PKCα inhibitor for only 5 days [14, 33]. This rapid reversal argues against an involvement of major transcriptional changes in the contractile dysfunction. In addition, our studies in adult mouse ventricular myocytes and neonatal rat ventricular myocytes showed that PLCβ1b overexpression results in dephosphorylation of phospholamban and depletion of Ca²⁺ from the sarcoplasmic reticulum [14]. Neither of these responses would be expected to require altered gene transcription, as they are likely dependent on well-established phosphorylation cascades [34]. However, it is perhaps surprising that chronically low contractility and reduced cardiac output does not of itself induce any substantial transcriptional response to increase inotropy or lusitropy. Presumably the hypertrophy that eventually ensues is such a response.
Fig 4. Comparison of rAAV6-PLCβ1-mediated gene expression changes to those caused by neonatal overexpression of Gqα or PKCα. A. Venn diagram demonstrating limited overlap of 748 mRNAs regulated at FDR<0.1 in a comparison of Gqα transgenic mice vs WT [24] and 29 mRNAs regulated at FDR<0.1 in a comparison of PKCα transgenic mice vs tet-off (TO) controls [27]. B. 761 unique mRNAs are displayed, from a composite of 15 mRNAs regulated at FDR<0.1 in a comparison of PLCβ1a or PLCβ1b vs blank, 748 mRNAs regulated at FDR<0.1 in a comparison of Gqα transgenic mice vs WT [24] and 29 mRNAs regulated at FDR<0.1 in a comparison of PKCα transgenic mice vs tet-off (TO) controls [27]. Unsupervised hierarchical clustering of relative mRNA abundances (rows) and of individual hearts (columns) was performed using Euclidean distance with average linkage. Colors represent log2 ratios for each individual heart vs the mean of the appropriate controls (red is upregulated, blue is downregulated); the color range spans -2.0 to +2.0 in log2 scale (from 4-fold downregulation to 4-fold upregulation in linear scale).

doi:10.1371/journal.pone.0158317.g004
Whilst we conclude that chronically depressed contractility does not meaningfully alter transcription, it must be considered that the PLCβ1b-expressing mice were unchallenged, caged and fed and so had limited requirement for flight or fight responses. It remains possible that the combination of chronically low contractility induced by PLCβ1b and a stressor requiring increased cardiac performance might result in an altered transcription response compared with stressed but otherwise untreated mice. As a result, stressing PLCβ1b-overexpressing mice might result in a more rapid development of compensatory hypertrophy, together with its associated transcriptional changes.

In cardiac signal transduction, PLCβ1b is immediately downstream of Gαq [16] and upstream of PKCα [14]. Our previous data have provided evidence that PLCβ1b is specifically activated downstream of Gαq in heart and thus similar transcriptional responses might be expected. In neonatal rat ventricular myocytes, the phenotype caused by heightened expression of Gαq is similar to that caused by PLCβ1b [16], with either of these causing primarily a hypertrophic response. However, both the phenotype and the transcriptional response to Gαq in mouse hearts are dependent on the timing of Gαq overexpression. Use of the conventional αMHC promoter system for Gαq transgene expression beginning immediately after birth promotes hypertrophy, dilatation and failure [30–32] and this is associated with substantial changes in the transcriptome [26]. In marked contrast, conditional overexpression of Gαq initiated in adult mouse hearts (a condition more similar to the rAAV6-mediated delivery used in our studies) did not result in hypertrophy, dilatation or heart failure and the limited mRNA expression data provided showed no changes in canonical markers of cardiac hypertrophy [35]. Whilst comprehensive transcriptome data are unfortunately not available for this latter condition, it seems unlikely that widespread transcriptional changes would be present in hearts exhibiting normal function. As noted previously [14], functional responses of the mouse heart to heightened PLCβ1b are similar to those reported previously in PKCα-overexpressing mice [36] and our data are consistent with a signaling role for PLCβ1b upstream of PKCα in mouse heart [14]. A relatively small set of mRNA expression changes was observed in PKCα-expressing hearts even though PKCα transgene expression commenced at birth [5, 27, 36]. Thus, findings from transgenes expressed from the immediate post-natal period are not directly comparable with the current data where heightened expression was initiated in adult animals. This also speaks to the importance of evaluating phenotypes in which signaling has been altered concomitant with the typical post-natal enlargement of the heart and acquisition of adult metabolic and sarcomeric properties, in contrast to phenotypes in which signaling is only altered once the heart has fully matured.

In addition to showing that depressed contractility per se does not initiate major transcriptional responses, our study also stresses the importance of a stringent control group in studies of this type. Although transcriptional changes were detected in response to PLCβ1b expression, these do not appear intuitively connected to pathways known to cause, or to be affected as a consequence, of blunted contractility. Because these alterations were similar to those in PLCβ1a-expressing hearts, we were able to define these as events not specific to PLCβ1b overexpression, but caused by the overexpression of the common molecular domains of PLCβ1a and PLCβ1b, which are 97% homologous (Fig 1A) and differ only in membrane-targeting capacity [37].

We have developed a mini-gene inhibitor that specifically targets PLCβ1b activation by preventing its targeting to the sarcolemma and have shown that expressing this inhibitor in hearts protects from heart failure following pressure overload. Additionally, we have identified the binding interface responsible for sarcolemmal targeting, by showing that proline-rich sequences in the extreme C-terminal sequence of PLCβ1b bind to the SH3 domain of the scaffolding protein Shank3 [37]. This binding interface provides a potential therapeutic target for
the development of a new class of inotropic agent. The finding that PLCβ1b does not induce transcriptional changes makes targeting its activation a more attractive proposition for drug development.

**Supporting Information**

**S1 Fig.** Variance to read-count relationship for the mRNAs subjected to statistical comparison. Dispersion (variance) declines in accordance with increasing read depth-normalized number of counts per mRNA. (TIF)

**S1 Table.** Gene descriptions and assignment to Gene Ontology categories for the 15 mRNAs regulated at FDR <0.1 by rAAV6-PLCB1a or rAAV6-PLCB1b expression. Assignment to Gene Ontology ‘biological process’ categories was performed using the Cytoscape plugin BiNGO [1]. Chd6 (chromodomain helicase DNA binding protein 6) had no Gene Ontology entry and is therefore not listed. (XLS)

**Acknowledgments**

We thank Dr Helen Kiriazis for assistance with mouse echocardiography and for virus preparation we thank Drs Hong Wei Qian and Paul Gregorevic.

**Author Contributions**

Conceived and designed the experiments: SJM DRG EAW. Performed the experiments: SJM DRG. Analyzed the data: SJM. Contributed reagents/materials/analysis tools: JRM EAW. Wrote the paper: SJM EAW.

**References**

1. Moore-Morris T, Guimaraes-Camboa N, Yutzey KE, Puceat M, Evans SM. Cardiac fibroblasts: from development to heart failure. J Mol Med (Berl). 2015; 93(8):823–30. doi:10.1007/s00109-015-1314-y
2. Roe AT, Frisk M, Louch WE. Targeting cardiomyocyte Ca^{2+} homeostasis in heart failure. Curr Pharm Des. 2015; 21(4):431–48. PMID: 25483944
3. Wei S, Guo A, Chen B, Kutschke W, Xie YP, Zimmerman K, et al. T-tubule remodeling during transition from hypertrophy to heart failure. Circ Res. 2010; 107(4):520–31. doi: 10.1161/CIRCRESAHA.109.212324 PMID: 20576937
4. Ziolo MT, Houser SR. Abnormal Ca^{2+} cycling in failing ventricular myocytes: role of NOS1-mediated nitroso-redox balance. Antioxid Redox Signal. 2014; 21(14):2044–59. doi: 10.1089/ars.2014.5873 PMID: 24801117
5. van Berlo JH, Maillet M, Molkentin JD. Signaling effectors underlying pathologic growth and remodeling of the heart. J Clin Invest. 2013; 123(1):37–45. doi: 10.1172/JCI62839 PMID: 23281408
6. Marks AR. Calcium cycling proteins and heart failure: mechanisms and therapeutics. J Clin Invest. 2013; 123(1):46–52. doi: 10.1172/JCI62834 PMID: 23281409
7. Sipido KR, Vangheluwe P. Targeting sarcoplasmic reticulum Ca^{2+} uptake to improve heart failure: hit or miss. Circ Res. 2010; 106(2):230–3. doi: 10.1161/CIRCRESAHA.109.210740 PMID: 2033907
8. Haghighi K, Bidwell P, Kranias EG. Phospholamban interactome in cardiac contractility and survival: A new vision of an old friend. J Mol Cell Cardiol. 2014; 77:160–7. doi: 10.1016/j.yjmcc.2014.10.005 PMID: 25451386
9. Kranias EG, Hajjar RJ. Modulation of cardiac contractility by the phospholamban/SERCA2a regulome. Circ Res. 2012; 110(12):1646–60. doi: 10.1161/CIRCRESAHA.111.259754 PMID: 22679139
10. Lefkowitz RJ, Rockman HA, Koch WJ. Catecholamines, cardiac beta-adrenergic receptors, and heart failure. Circulation. 2000; 101(14):1634–7. PMID: 10758041
11. Woodcock EA, Grubb DR, Filtz TM, Marasco S, Luo J, McLeod-Dryden TJ, et al. Selective activation of the "b" splice variant of phospholipase Cbeta1 in chronically dilated human and mouse atria. J Mol Cell Cardiol. 2009; 47(5):676–83. doi: 10.1016/j.yjmcc.2009.08.020 PMID: 19729020

12. Bahk YY, Lee YH, Lee TG, Seo J, Ryu SH, Suh PG. Two forms of phospholipase C-beta 1 generated by alternative splicing. J Biol Chem. 1994; 269(11):8240–5. PMID: 7510682

13. Smrcka AV, Hepler JR, Brown KO, Sternweis PC. Regulation of polyphosphoinositide-specific phospholipase C activity by purified Gq. Science. 1991; 251(4995):804–7. PMID: 1846707

14. Grubb DR, Crook B, Ma Y, Luo JT, Qian HW, Gao XM, et al. The atypical 'b' splice variant of phospholipase C beta 1 promotes cardiac contractile dysfunction. J Mol Cell Cardiol. 2015; 85:95–103. doi: 10.1016/j.yjmcc.2015.04.016 PMID: 25918049

15. Grubb DR, Gao XM, Kiriazis H, Matsumoto A, McMullen JR, Du XJ, et al. Expressing an inhibitor of PLCbeta1b sustains contractile function following pressure overload. J Mol Cell Cardiol. 2016; 93:12–7. doi: 10.1016/j.yjmcc.2016.02.015 PMID: 26906633

16. Filtz TM, Grubb DR, McLeod-Dryden TJ, Luo J, Woodcock EA. Gq-initiated cardiomyocyte hypertrophy is mediated by phospholipase Cbeta1b. FASEB J. 2009; 23(10):3564–70. doi: 10.1096/fj.09-133983 PMID: 19564249

17. Rowell J, Koitabashi N, Kass DA, Barth AS. Dynamic gene expression patterns in early and late heart failure reveal biphasic-bidirectional transcriptional activation of signaling pathways. Physiol Genomics. 2014; 46(20):779–87. doi: 10.1152/physiolgenomics.00054.2014 PMID: 25159852

18. Chen J, Wang HY, Zeng CY. Transcriptome network analysis of potential candidate genes for heart failure. Genet Mol Res. 2013; 12(4):4687–97. doi: 10.4238/2013.October.18.7 PMID: 24222245

19. Chen HB, Wang L, Jiang JF. Re-analysis of expression profiles for revealing new potential candidate genes of heart failure. Eur Rev Med Pharmacol Sci. 2013; 17(7):903–11. PMID: 23640436

20. Gregorevic P, Blankinship MJ, Allen JM, Crawford RW, Meuse L, Miller DG, et al. Systemic delivery of genes to striated muscles using adeno-associated viral vectors. Nat Med. 2004; 10(8):828–34. doi: 10.1038/nm1085 PMID: 15275747

21. Trapnell C, Roberts A, Goff L, Pertea G, Kim D, Kelley DR, et al. Differential gene and transcript expression analysis of RNA-seq experiments with TopHat and Cufflinks. Nat Protoc. 2012; 7(3):562–78. doi: 10.1038/nprot.2012.016 PMID: 22383036

22. Anders S, Pyl PT, Huber W. HTSeq—a Python framework to work with high-throughput sequencing data. Bioinformatics. 2015; 31(2):166–9. doi: 10.1093/bioinformatics/btu638 PMID: 25260700

23. Anders S, McCarthy DJ, Chen Y, Okoniewski M, Smyth GK, Huber W, et al. Count-based differential expression analysis of RNA sequencing data using R and Bioconductor. Nat Protoc. 2013; 8(9):1765–86. doi: 10.1038/nprot.2013.099 PMID: 23975260

24. Westenbrink BD, Ling HY, Divakaruni AS, Gray CBB, Zambon AC, Dalton ND, et al. Mitochondrial reprogramming induced by CaMKII delta mediates hypertrophy decompensation. Circ Res. 2015; 116(5):e28–e39. doi: 10.1161/CIRCRESAHA.116.304682 PMID: 25605649

25. Zhang Y, Matkovich SJ, Duan X, Gold JI, Koch WJ, Dorn GW 2nd. Nuclear effects of G-protein receptor kinase 5 on histone deacetylase 5-regulated gene transcription in heart failure. Circ Heart Fail. 2011; 4(5):659–68. doi: 10.1161/CIRCHEARTFAILURE.111.962563 PMID: 21768220

26. Matkovich SJ, Zhang Y, Van Booven DJ. Dorn GW 2nd. Deep mRNA sequencing for in vivo functional analysis of cardiac transcriptional regulators: application to Galphaq. Circ Res. 2010; 106(9):1459–67. doi: 10.1161/CIRCRESAHA.109.217513 PMID: 20360248

27. Zhang Y, Matkovich SJ, Duan XJ, Diwan A, Kang MY, Dorn GW. Receptor-independent PKCa signaling by calpain-generated free catalytic domains induces HDAC5 nuclear export and regulates cardiac transcription. J Biol Chem. 2011; 286(30):26943–51. doi: 10.1074/jbc.M111.234757 PMID: 21642422

28. Matkovich SJ, Edwards JR, Grossenheider TC, de Guzman Strong C, Dorn GW 2nd. Epigenetic coordination of embryonic heart transcription by dynamically regulated long noncoding RNAs. Proc Natl Acad Sci U S A. 2014; 111(33):12264–9. doi: 10.1073/pnas.1410622111 PMID: 25071214

29. Song M, Matkovich SJ, Zhang Y, Hammer DJ. Dorn GW 2nd. Combined cardiomyocyte PKCd and PKCe gene deletion uncovers their central role in restraining developmental and reactive heart growth. Sci Signal. 2015; 8(373):ra39. doi: 10.1126/scisignal.aaa1855 PMID: 25900833

30. Dorn GW 2nd, Brown JH. Gq signaling in cardiac adaptation and maladaptation. Trends Cardiovasc Med. 1999; 9(1–2):26–34. S1050-1738(99)00004-3 [pii]. PMID: 10189964

31. Mende U, Kagen A, Cohen A, Aramburu J, Schoen FJ, Neer EJ. Transient cardiac expression of constitutively active Galphaq leads to hypertrophy and diluted cardiomyopathy by calcium-intrinsically dependent and independent pathways. Proc Natl Acad Sci U S A. 1998; 95(23):13893–8. PMID: 9811897
32. Adams JW, Sakata Y, Davis MG, Sah VP, Wang Y, Liggett SB, et al. Enhanced Gaq signaling: a common pathway mediates cardiac hypertrophy and apoptotic heart failure. Proc Natl Acad Sci USA. 1998; 95:10140–5. PMID: 9707614

33. Hambleton M, Hahn H, Pleger ST, Kuhn MC, Klevitsky R, Carr AN, et al. Pharmacological- and gene therapy-based inhibition of protein kinase Calpha/beta enhances cardiac contractility and attenuates heart failure. Circulation. 2006; 114(6):574–82. doi:10.1161/CIRCULATIONAHA.105.592550 PMID: 16880328

34. Florea S, Anjak A, Cai WF, Qian J, Vafiadaki E, Figueria S, et al. Constitutive phosphorylation of inhibitor-1 at Ser67 and Thr75 depresses calcium cycling in cardiomyocytes and leads to remodeling upon aging. Basic Res Cardiol. 2012; 107(5):279. doi:10.1007/s00395-012-0279-z PMID: 22777184

35. Syed F, Odley A, Hahn HS, Brunskill EW, Lynch RA, Marreez Y, et al. Physiological growth synergizes with pathological genes in experimental cardiomyopathy. Circ Res. 2004; 95(12):1200–6. doi:10.1161/01.RES.0000150366.08972.7f PMID: 15539635

36. Braz JC, Gregory K, Pathak A, Zhao W, Sahin B, Klevitsky R, et al. PKC-alpha regulates cardiac contractility and propensity toward heart failure. Nat Med. 2004; 10(3):248–54. doi:10.1038/nm1000 PMID: 14966518

37. Grubb DR, Luo JT, Woodcock EA. Phospholipase C beta 1b directly binds the SH3 domain of Shank3 for targeting and activation in cardiomyocytes. Biochem Biophys Res Comm. 2015; 461(3):519–24. doi:10.1016/j.bbrc.2015.04.060 PMID: 25911318