Transcriptional profiling of foam cells in response to hypercholesterolemia

Young-Hwa Goo a,⁎, Vijay K. Yechoor b, Antoni Paula a,⁎

a Department of Molecular and Cellular Physiology, Albany Medical College, Albany, NY, United States
b Division of Diabetes, Endocrinology & Metabolism, Department of Medicine, Baylor College of Medicine, Houston, TX, United States

ABSTRACT

Hypercholesterolemia is a main risk factor for atherosclerosis development. Arterial macrophages, or foam cells, take-up and process lipoprotein particles deposited in arteries, and store much of the cholesterol carried by these particles in their cytoplasm. However, the effects of exposure to different cholesterol levels on foam cells remain poorly understood. Given the remarkable plasticity of macrophages in response to environmental variables, studies on macrophage biology should ideally be performed in the environment where they exert their physiological functions, namely atherosclerotic lesions in the case of foam cells. We used a mouse model of atherosclerosis, the apolipoprotein E-deficient mouse, to study in vivo the transcriptional response of foam cells to short- and long-term elevations in plasma cholesterol, induced by feeding mice a western type diet. The microarray data sets from this study have been deposited in NCBI's Gene Expression Omnibus under the accession number GSE70619. Here we provide detailed information on the experimental set-up, on the isolation of RNA by laser capture microdissection, and on the methodology used for RNA amplification and analysis by microarray and quantitative real-time PCR.

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2.2. Laser capture microdissection of macrophages

Like in humans, atherosclerotic lesions of apoE−/− mice are cellularly heterogeneous, and the small size of murine arteries represents an added challenge for the isolation of specific cell populations. Laser capture microdissection (LCM) is a technique that allows to identify and isolate specific cells from sections of heterogeneous tissues, and can therefore be used to harvest individual cell populations within mouse atheroma [2]. To this end, we performed consecutive 7-µm sections through the aortic sinuses of mice in our 3 experimental groups. Sections were sequentially mounted on 3 slides. The first and third slides, which were used for LCM, were immediately fixed through a battery of cells, and nuclei were stained with toluidine blue with the HistoGene LCM Frozen Section Staining Kit (Applied Biosystems), and stored O/N at room temperature in a slide box containing fresh desiccant. The second slide was stained for macrophages with anti-Lamp2/Mac3 antibody (Santa Cruz Biotechnology), and used as a template to identify macrophages within lesions. This slide was also used to quantify lesion area, and the total and relative areas of macrophages within lesions. LCM of macrophages was performed with a Veritas Microdissection System (Applied Biosystems). We used an infrared laser to perform ~2000 shots on macrophage-rich areas of approximately 20 sections of each sample. While the duration and intensity of the laser pulses were adjusted for each individual sample, normally the laser power was set to ~65 mV and the pulse duration was ~2500 µs, which typically resulted in a spot size of between 15 and 20 µm. Individual cells or small clusters were collected on CapSure HS LCM Caps (Applied Biosystems).

2.3. RNA purification and amplification

Immediately after LCM, RNA from captured macrophages was purified with the PicoPure RNA Isolation Kit (Applied Biosystems) and stored at −80 °C. While this RNA is suitable for analysis of gene expression by quantitative polymerase chain reaction (qPCR), the amount of RNA that can be isolated considerably limits the number of genes that can be analyzed. Thus, we used the RiboAmp HS RNA Amplification Kit (Applied Biosystems) to perform two rounds of RNA amplification by in vitro transcription (IVT). Each round consisted of a reverse transcription using oligo(dT) primers tagged with the T7 promoter sequence and synthesis of double-stranded DNA, followed by an IVT reaction with a T7 RNA polymerase to yield amplified complementary RNA (cRNA). The quality of the cRNA was assessed by A260/A280 ratios and by an electropherogram with the Agilent 2100 Bioanalyzer. Of note, the IVT reactions do not amplify ribosomal RNA. Thus, standard quality controls such as 28S/18S ribosomal RNA ratio or the RNA integrity number (RIN) cannot be used to characterize amplified RNA. Instead, upon electrophoresis the amplified cRNA typically appears as a broad band, and we confirmed that the intensity distribution of the bands were similar among samples. Additional quality controls were performed using qPCR to confirm the enrichment in macrophage markers in the LCM populations with respect to RNA isolated and amplified from whole aortic sinuses, as well as depletion in endothelial and smooth muscle cell markers in the same samples [1,2].

2.4. Gene expression analysis

For broad gene expression profiling the LCM-RNAs from foam cells in the different dietary groups were hybridized to Mouse Genome 430 2.0 Arrays from Affymetrix. A common step when using the Affymetrix platform is the use of an IVT reaction to generate biotinylated cRNA. Given that the LCM RNA was already submitted to two rounds of IVT, for cRNA labeling we employed the TURBO Labeling Biotin Kit (Applied Biosystems) to biotinylate 15 µg of each cRNA. This system uses a chemical reaction to form a coordinate bond between the base guanine and a biotin complex, and does not involve further RNA amplification. Labeled cRNA was then fragmented and hybridized to microarray chips, and subsequent steps such as washing, staining and scanning were performed following standard Affymetrix protocols at the Genomic and RNA Profiling Core at Baylor College of Medicine. The microarray data were filtered with dChip software to only include in the analysis the probes with ≥50% presence call and with expression levels of ≥25 in ≥50% samples. The filtered data were transferred to MeV software [3], log2 transformed, and analyzed by ANOVA. Pairwise comparisons were performed with the Welch’s t-test, assuming unequal variances. Differences were considered significant when P < 0.01. To determine the main biological pathways affected in foam cells in response to WD feeding for 2 or 14 weeks, data were analyzed with Pathway Express [4]. A P value was generated based on the number of genes differentially
expressed in each pathway with respect to the number expected to change by chance. Pathway Express was also used to generate a gamma \( P \) value that, in addition to classical statistics, takes into consideration relevant biological parameters such as the fold-change and the topology of the genes in the pathways. A detailed description of the results of the analyses can be found in [1].

Quantitative-PCR was used to confirm the enrichment in macrophages in the LCM populations and to validate the microarray data. Reverse transcription was performed with SuperScript III (Invitrogen), using random hexamers as primers for the cRNA. The classical antisense RNA amplification by IVT is usually 3′ biased, i.e. the product is shorter than the parent mRNA. This is because oligo(dT) primers are used to insert the T7 polymerase promoter and, as a result, transcription is more robust in mRNA regions closer to the 3′ end, where the reaction is initiated. Thus, primers were designed at the 3′-terminus region of the mRNA, including the 3′-untranslated region. Other than these considerations, standard PCR protocols and quality controls were used for analysis of gene expression by qPCR.

3. Discussion

Foam cells are a hallmark of atherosclerotic lesions, and play central roles in the regulation of both lipid homeostasis at the arterial wall and local inflammation. Given the high plasticity of macrophages in response to environmental variables, it seems most reasonable to study their function under the environment where they exert their physiological role. This may be quite challenging when we study atherosclerosis in mice, as the small size of their arteries and the cellular heterogeneity of atherosclerotic lesions make it difficult to isolate specific cell populations. Here we describe the use of LCM for isolation of foam cells resident within murine atherosclerotic lesions that were exposed to different cholesterol levels, as well as the methodology and quality controls used for RNA amplification and gene expression analysis. Given the high relevance of inflammation to the pathogenesis of atherosclerosis, the emphasis of our initial study was on the analysis of genes that regulate inflammatory and immune responses [1]. By thoroughly describing the experimental design and procedures, this article may facilitate the work of other researchers who might be interested in analyzing the data from a different perspective.

Conflict of interest

The authors declare no conflict of interest.

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