Metabolites Interrogation in Cell Fate Decision of Cultured Human Corneal Endothelial Cells

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PURPOSE. Aiming to clarify the metabolic interrogation in cell fate decision of cultured human corneal endothelial cells (cHCECs).

METHODS. To analyze the metabolites in the culture supernatants (CS), 34 metabolome measurements were carried out for mature differentiated and a variety of cHCECs with cell state transition through a facility service. Integrated proteomics research for cell lysates by liquid chromatography−tandem mass spectrometry (LC-MS/MS) was performed for 3 aliquots of each high-quality or low-quality chCEC subpopulations (SP). The investigations for the focused genes involved in chCEC metabolism were performed by using DAVID and its options “KEGG_PATHWAY.”

RESULTS. The clusters of metabolites coincided well with the distinct content of CD44−/+ SPs. Both secreted pyruvic acid and lactic acid in the CS were negatively correlated with the content of high-quality SPs. Lactic acid and pyruvic acid in the CS exhibited the positive correlation with that of Ile, Leu, and Ser, whereas the negative correlation was with glutamine. Platelet-derived growth factor-ββ in the CS negatively correlated with lactic acid in CS, indicating indirectly the positive correlation with the content of CD44−/+ SPs. Upregulated glycolytic enzymes and influx of glutamine to the tricarboxylic acid cycle may be linked with a metabolic rewiring converting oxidative metabolism in mature differentiated CD44−/+ SPs into a glycolytic flux-dependent state in immature SPs with cell state transition.

CONCLUSIONS. The findings suggest that the cell fate decision of cHCECs may be dictated at least partly through metabolic rewiring.

Keywords: CD44, c-Myc, p53, lactate, pyruvic acid, oxidative phosphorylation, glycolysis, mitochondria

Corneal endothelial (CE) disorders are caused by pathologic conditions such as bullous keratopathy and Fuchs’s endothelial corneal dystrophy.1 Recently, we reported our development of a novel clinically effective therapeutic modality to regenerate monolayer CE tissues by injection into an anterior chamber of cultured human CE cells (cHCECs).2–8 Recently, cHCECs have been recognized to be heterogeneous not only in their biochemical features but also in their key functions, including the metabolic plasticity.9

Corneal endothelium is one of the most metabolically active tissues in the human body.10 Dissecting the biologic functions of metabolites and mitochondria in chCEC cell fate decision will provide essential cues to further improve cell-based therapy. Mitochondria are poised to play an essential role.11,12 Glycolysis and mitochondrial respiration are the most relevant pathways for producing adenosine triphosphate (ATP) indispensable to HCE cell functions.13–19 However, the requirement for specific metabolic reprogramming in the maturation/differentiation of chHCECs or in cell fate decision remains elusive.20 Previously, we preliminary described20 that metabolomic profiling segregated mature differentiated-chCECs from immature-chCECs.9 Cultured HCECs have a tendency for cell-state transition (CST) into a dedifferentiation state, a senescent phenotype, epithelial-mesenchymal transition, and transformed fibroblastic cell morphology, indicating the presence of vast heterogeneity among these chHCECs with CST. Among chCECs with CST, immature chCECs switched to a glycolytic metabotype, whereas mature differentiated chCECs became more oxidative and elicited reduced amount secretion of lactate, suggesting a possible application of metabolomics for...
quality control or their usefulness as biomarkers for diagnosis, prognosis, and therapeutic efficacy of cHCEC cell therapy for CE disorders.

The secreted metabolites were assigned to the cHCEC subpopulations (SPs) distinct in the expression levels of surface CD44 antigens, which expression is regulated by c-Myc. The link between bioenergetics and the cell fate decision of cHCECs remains largely unknown. In other cell systems, CD44 functions to regulate metabolic flux to mitochondrial respiration and induces entry into glycolysis. C-Myc reportedly regulates lactate production and is an important regulator for glycolysis. It has been suggested that c-Myc can addict cells to specific bioenergetic substrates. In this context, unraveling the plasticity of the mitochondrial response to in vitro cell culture environments would provide critical insights into how cHCEC cell fate decisions are established and may support the cell-based regenerative medicine.

Given the distinct expression profiles of CD44, cMyc and p53, also known as a major metabolic regulator, among cHCEC SPs, here we had tried to detail the inclination of p53, also known as a major metabolic regulator, among medicine.

Metabolites in Cell Fate Decision of cHCECs

**Materials and Methods**

**Human Corneal Tissue and Endothelial Cell Donors**

The human tissue used in this study was handled in accordance with the tenets set forth in the Declaration of Helsinki. HCECs were obtained from human donor corneas obtained from CorneaGen Inc. (Seattle, WA) eye bank, and were cultured before the experimental analysis. Informed written consent for eye donation for research was obtained from the next of kin of all deceased donors. All tissues were recovered under the tenets of the Uniform Anatomical Gift Act of the particular state in which the donor consent was obtained and the tissue was recovered. All donor corneas were preserved in Optisol-GS (Chiron Vision, Inc., Irvine, CA) corneal storage medium and then shipped by international air transport for research purposes. Donor consent was obtained and the tissue was recovered. All chemical, biological, and physical quality specifications were measured in the positive and negative modes of CE time-of-flight mass spectrometry, and anionic compounds were measured in the positive mode of CE tandem MS (CE-MS/MS). Hierarchical cluster analysis (HCA) was performed by use of our proprietary software, that is, “PeakStat.” cHCECs from donors between the ages of 7 and 29 were used to obtain reproducible cultures producing high-quality cHCECs suitable for cell-injection therapy. Thus we used cHCECs from 3 donors of nearly the same age (age: 17, 14, and 18 years) to clarify the influence of the fine differences of the culture conditions. Differences between the values were statistically analyzed by use of the Student’s t test or 1-way analysis of variance with Bonferroni post hoc tests (GraphPad Prism 6.0, after harvest with 10× TrypLE Select (Thermo Fisher Scientific, Inc., Waltham, MA) treatment at 37° C for 12 minutes when they reached confluence. The HCECs at passages 2 to 3 were used for all experiments.

**Flow Cytometry Analysis of cHCECs**

HCEC cells were collected from the culture dish by TrypLE Select treatment, as described above, and provided to Fluorescence Activated Cell Sorter (FACS) analysis according to the procedures described previously. The antibodies used were as follows: FITC-conjugated anti-human CD90 mAb, phycoerythrin (PE)-conjugated anti-human CD166 mAb, PerCP-Cy 5.5 conjugated anti-human CD24 mAb, PE-Cy 7–conjugated anti-human CD44 and FITC-conjugated antihuman CD90 mAb (all from BD Biosciences, San Jose, CA), and allophycocyanin APC-conjugated anti-human CD105 (eBioscience, Inc.). After washing with FACS buffer, the HCECs were analyzed by use of the BD FACS Canto II (BD Biosciences) fluids-system flow cytometer.

**Reagents**

Rho-associated protein kinase (ROCK)-inhibitor Y-27632 (Y) and epidermal growth factor was purchased from Wako Pure Chemical Industries, Ltd. (Osaka, Japan) and SB203580 (SB2) was obtained from Cayman Chemical (Ann Arbor, MI). Dulbecco’s modified Eagle’s medium—high glucose (DMEM-HG) and fetal bovine serum were obtained from Gibco Industries Inc. (Langley, OK) and plastic culture plates were obtained from Corning. Unless indicated differently, all other chemicals were purchased from Sigma-Aldrich, Inc. (St. Louis, MO).

**Measurement of Metabolites in Culture Medium**

For the measurement of metabolites in the culture supernatants (CS), 20 μL of the CS and 80 μL Milli-Q (Merck KGaA, Darmstadt, Germany)—purified water containing Internal Standard Solution 1 (H3304-1002; Human Metabolome Technologies, Inc., Yamagata, Japan) were thoroughly mixed. The procedures followed those described previously. Briefly, cationic compounds were measured in the positive mode of CE time-of-flight mass spectrometry, and anionic compounds were measured in the positive and negative modes of CE tandem MS (CE-MS/MS).
Integrated Analysis of Phosphoprotein by Bio-Plex

The culture lysates of cHCECs were harvested after 4-days cultivation and then immediately frozen and stored at −80°C until analysis. The phosphoprotein levels of the cHCEC lysates were analyzed by Luminex Corporation (Austin, TX) xMap Technology (Bio-Plex 200; Bio-Rad Laboratories, Inc., Hercules, CA) with the Bio-Plex Human 27-Plex Panel Kit (Bio-Rad Laboratories) according to the manufacturer’s instructions. Among the multiple phosphoproteins, only phosphorylated platelet-derived growth factor-ββ receptor-B was used for this study. On the other hand, CS were harvested from the corresponding culture of cHCECs and spun in a centrifuge at 1580g at room temperature for 10 minutes to remove detached cells. The CS were collected and filtered through 0.220-μm filters (Milllex-GV; EMD Millipore Corporation, Temecula, CA). Enzyme-linked immunosorbent assay was performed by use of a PDGF-ββ human enzyme-linked immunosorbent assay kit (Abcam Plc., Cambridge, UK). The measurement of metabolites in the CS was carried out as described above.

GraphPad software). A P value <0.05 was considered statistically significant.

LC/MS Data-Set Analysis

The LC/MS data set, composed of 4641 proteins in total, was obtained by use of Proteome Discoverer 2.2 software. After removal of the data in which the abundance ratio could not be calculated, we analyzed the remaining data by means of a web-based program, DAVID v6.8 (The Database for Annotation, Visualization and Integrated Discovery; https://david.ncifcrf.gov). Finally, it ended up with 4315 genes, each with a unique DAVID Gene ID, for the subsequent analyses.

As for the gene expression analysis, we calculated the statistical P value and fold-change between 2 groups and drew the volcano plot to extract genes differentially expressed in HQ and LQ cHCECs. Further investigations for the focused genes, as well as the related genes/pathways that were suggested to be involved in cHCEC metabolism, were performed by using DAVID and its options “BIOCARTA” and “KEGG_PATHWAY.” We also referred their original databases of BioCarta (https://cgap.nci.nih.gov/Pathways/BioCarta_Pathways) or KEGG (Kyoto Encyclopedia of Genes and Genomes; https://www.genome.jp/kegg/) and showed the referred genes/pathways in the figures with slight modifications.

As for the gene ontology (GO) analysis, we divided the data into 3 groups based on the range of abundance ratio (LQ/HQ), and analyzed in DAVID with a “GOTERM_DIRECT” option by each group. The results of GO were sorted by the P value and top ten-ranked GO terms in each group are shown. In LC/MS data analysis, the significance of difference between HQ and LQ cHCECs was assessed by Student’s or Welch’s t-test after the confirmation by F test.

Results

Variations of Metabolites in CS of HCECs

Aiming to develop a practical and noninvasive method for monitoring the conformity of cHCEC quality for regenerative therapy, we comprehensively surveyed the secretory metabolites in CS precisely assigned to the cHCEC SPs distinct in the expression levels of surface CD44 antigens. Cultured HCECs subjected to the analysis were C32 (P2, 17Y, ECD=3347), C33 (P2, 14Y, 3554), and C23 (P2, 18Y, 3280), all produced for clinical setting of cell infusion therapy. Unsupervised HCA of the metabolites in the CS among cHCECs, C32, C33, and C23 turned out to identify 4 metabolite clusters. Despite the superficially similar microscopic features of these 3 cHCECs (data not shown), the HCA differed greatly, between C32, 33, and C23 (Fig. 1).

The distinction of these clustering coincided well with the distinct content of CD44−/± SPs between these 2 groups; the former 2 were almost mature differentiated SPs with E-ratio, 94.6 and 99.3%, respectively, whereas the latter intermediary mature one was 76.3%. The clusters were divided into 3 metabolite clusters (Fig. 1): (1) increased metabolites in mature differentiated CD44−/± cHCECs, namely glyoxylate, glycerol 3-phosphate acid, 2-phosphoglyceric acid, glycolic acid, folic acid, guanine, His, Thr, Pro, Phe, Trp, Tyr, and Lys, (2) decreased metabolites in mature differentiated CD44−/+ cHCECs, namely β-Ala, pyruvic acid, Ser, branched chain amino acids (BCAA) Ile, Leu, and Val, and citrulline, (3) increased metabolites in both cHCECs, namely TCA cycle related metabolic intermediates: lactic acid, malic
Figure 1. Hierarchical cluster analysis identified 3 metabolite subsets. Hierarchical clustering of the metabolites in the CS among cHCECs, C32, C33, and C23. Unsupervised HCA showed clear separation among these 3 lots of cHCECs. Cultured HCECs subjected to the analysis were C32 (male donor, P2, 23Y, ECD3884), C33 (male donor, P2, 29Y, ECD3309), and C23 (female donor, P2, 18Y, 3280). ECD, endothelial cell density in donor corneal tissues; P2, passage 2; Y, year of the corneal tissue donors.

Table 1. Variations of Secreted Metabolites between High- and Low-Quality cHCECs

| Decreased Secretion in High-Quality cHCECs | Increased Secretion in High-Quality cHCECs | Increased Secretion in Low-Quality cHCECs |
|-------------------------------------------|-------------------------------------------|-------------------------------------------|
| β-Ala | Glyoxylate | Lactic acid |
| Pyruvic acid | Glyceraldehyde 3-phosphate | Malic acid |
| Ser | 2-Phosphoglyceric acid | Argininosuccinic acid |
| BCAA (Ile, Leu, Val) | Glycolic acid | Fumaric acid |
| Citrulline | Folic acid | cis-Aconitic acid |
| | Guanine | Citric acid |
| | His | Succinic acid |
| | Thr | Ornithine |
| | Pro | Uric acid |
| | Phe | Urea |
| | Trp | |
| | Tyr | |
| | Lys | |

High-Quality cHCECs corresponds to culture lots C32 (P2, 17Y, 3347) and C33 (P2, 14Y, 3554), and Low-Quality cHCECs to culture lot C23 (P2, 18Y, 3280) described in the text. Typical metabolites are listed.

In the previous finding we had confirmed that CD44−/+ mature-differentiated cHCECs are disposed to mitochondria-dependent oxidative phosphorylation, whereas CD44++/+++ immature cHCECs with CST switched to a glycolytic metabotype. In this context, we next investigated, using 28 lots of cHCECs different in the content of SPs, the association of secreted lactic acid or pyruvic acid with BCAA, Ser and Gln, which were confirmed to be selectively up- or down-regulated in HCA analysis described above (Table 1). Pyruvic acid in the CS of cHCECs exhibited the positive correlation with Ile, Leu, and Ser (R² = 0.675, 0.710, and 0.716, respectively), whereas the negative correlation was with Gln (R² = 0.308) (Fig. 2a). The correlation of lactic acid in the CS of cHCECs exhibited almost the same tendency; positive correlation was with Ile, Leu and Ser (R² = 0.689, 0.712 and 0.541, respectively), whereas the negative correlation was with Gln (R² = 0.548) (Fig. 2b).

Association of Secreted Pyruvic Acid and Lactic Acid with E-Ratios

Both secreted pyruvic acid and lactic acid in the CS of 28 different lots of cHCECs were negatively correlated (R² = 0.470 and 0.627, respectively) with the content of CD44− CD90− mature differentiated SPs in cHCECs, whereas those were positively correlated (R² = 0.817 and 0.880, respectively) with that of CD44++ CD90+ immature SPs in cHCECs (Fig. 3). The content of CD44− CD90−
Figure 2. Correlation of secreted lactic acid or pyruvic acid with BCAA, Ser, and Gln in the culture supernatants among 28 different lots of chCECs. The metabolites were analyzed by CE-MS, as described in the text. (a) Association of secreted BCAA, Ser, Gln, and lactic acid. (b) Association of BCAA, Ser, Gln, and Pyruvic Acid

Figure 3. Association of secreted pyruvic acid and lactic acid with effector ratios (E-ratios). Both of secreted pyruvic acid and lactic acid in the CS were analyzed by CE-MS similarly as other metabolites. The chCEC SP with surface expression of CD166+, CD105-, CD44-, CD26-, CD90-, and CD24- was designated as effector cells and the ratio of that SP was designated among heterogeneous chCECs as an E-ratio.

Association of Phosphorylation of PDGFR-β with Secreted Products
To investigate the roles of cytokines extracellularly secreted, we studied the effect of PDGF-ββ in modulating metabolic
FIGURE 4. Diagram illustrating the inverse correlation of PDGF-ββ ligand and lactic acid, Ile, and Ser in the CS of cHCECs. The intensity of phosphorylation of PDGFR-B was measured as detailed in the text. Twelve different cultures, n=3 for each with one culture additive deleted, namely, no deletion control, –epidermal growth factor, –SB2, or –Y, were extended either 34 or 35 days after cell seeding, and the cell lysates and corresponding CSs were then analyzed. (a) Association of phosphorylation of PDGFR-B with lactic acid or PDGF-ββ in the CS. (b) Association of phosphorylation of PDGFR-B with Ile or Ser in the CS.
dynamics of cHCECs. Considering that PDGF-ββ is one of the most abundant cytokines secreted selectively from mature differentiated cHCECs (Hamuro, unpublished data), the amount of free PDGF-ββ ligand in CS of cHCECs, cultivated under different culture conditions producing divergent cHCEC SPs, might be greatly distinct. To confirm the hypothesis we prepared 12 different cultures from 3 separate donor corneal tissues. Actually as illustrated in Figure 4, the intensity of phosphorylation of PDGFR-B was inversely correlated with the amount of free PDGF-ββ in CS (R² = 0.594) among 12 different cultures, n = 3 for each with one culture additive deleted, namely epidermal growth factor, SB2, or Y. All of the culture was extended either 34 or 35 days after cell seeding, and the cell lysates and the corresponding CS were analyzed.

Of note, the amount of phosphorylated PDGFR-B was positively correlated with the amount of lactic acid in the CS (R² = 0.832) (Fig. 4a), indicating the critical role of PDGF-ββ in rewiring of energy metabolism in cHCECs to dispose to glycolysis producing higher amount of lactic acid extracellularly. Accordingly, the amount of phosphorylated PDGFR-B was also positively correlated with the amount of Ile and Ser in the CS (R² = 0.470 and 0.569, respectively) (Fig. 5b), indicating the critical role of these 2 amino acids in reducing PDGFR-B phosphorylation and in the increase of resulting PDGF-ββ in the CS, which in turn correlated with the elevation of E-ratios. This observation is also in consistent with the above-mentioned positive correlation of the amount of lactic acid in the CS with those of Ile and Ser (Fig. 2a).

**Integral Proteome Analysis: Gene Ontology Analysis**

After calculating the abundance ratio for all proteins detected either in the immature LQ or mature differentiated HQ cHCEC SP, we analyzed the data by means of a web-based program, DAVID v6.8 for 4,315 genes. The numbers of identified proteins from these SPs were classified by Venn diagram. Total identified proteins (3781), immature LQ cHCEC dominant proteins (160), and mature differentiated HQ cHCEC dominant proteins (253) were characterized (the complete procedures are diagrammed in Fig. 5). Biologic characteristics of HQ and LQ cHCECs were clarified. Top ten-ranked biologic processes (BP) and cellular components (CC) for each group are shown as bar charts (Fig. 6).

As illustrated in BP classification, cell-to-cell adhesion, glycolytic process, and glucogenesis-related proteins were enriched in the group with the abundance ratio > 2.0, namely in LQ cHCECs, whereas metabolic process, TCA cycle, oxidation-reduction process, fatty acid β-oxidation, and branched-chain amino acid metabolism–related proteins were enriched in the group with the abundance ratio < 0.5, namely in HQ cHCECs. These kinds of segregated distributions of distinct functional proteins were observed also in CC classification. Mitochondria and lysosome related proteins were enriched in HQ cHCECs, whereas cytoplasmic proteins, extracellular exosomes, stress fiber or actin cytoskeleton related proteins were enriched in LQ cHCECs. Some of the observations were fully consistent with our previous observation describing the metabolomic profiling segregated differentiated from dedifferentiated-cHCECs, the linkage of
the dedifferentiation state with epithelial-mesenchymal transition and transformed fibroblastic cell morphology, and the observation that cHCECs SPs with CST secreted higher amount of exosomes than the mature differentiated cHCECs SPs.

### Elevated Expression of Anaplerotic Metabolic Enzymes in Immature cHCECs

Further investigations for the focused genes and the related genes/pathways involved in cHCEC metabolism, were performed. The part of the referred genes/pathways were shown in the figures with slight modifications for glycolysis (a) and alanine, aspartate and glutamate metabolism (b) (Figs. 7a, b).

It is of interest that most of the enzymes involved in the earlier metabolic stage of glycolysis were upregulated in immature LQ cHCEC SPs than in mature differentiated HQ cHCECs SPs. Typical enzymes are aldolase A, B, and C, enolase 1, 2, glyceraldehyde-3-phosphate dehydrogenase, glucose-6-phosphate isomerase, hexokinase 2, lactate dehydrogenase A, phosphoglycerate dehydrogenase 1, 2, and 4, phosphoglucomutase 1, 2, and pyruvate kinase. On the other hand, pyruvate dehydrogenase (conversion of pyruvate to acetyl-CoA) and aldehyde dehydrogenase 2 family (mitochondrial localized) were selectively upregulated in HQ cHCEC SPs, whereas other isoforms of aldehyde dehydrogenase were isoform specifically upregulated either in HQ or LQ cHCEC SPs.

In regard to alanine, aspartate, and glutamate metabolism (Fig. 7b), argininosuccinate synthase 1 (ASS1), known as the catalytic enzymes in urea cycle, and glutamate-ammonia ligase were upregulated in the LQ cHCEC SPs, whereas, to the contrary, the reduced expression was evident for glutamate dehydrogenase 1, 2 and mitochondrial glutaminase in those SPs (Fig. 7b).

### DISCUSSION

We previously reported the presence of heterogeneous cHCEC SPs and identified 1 specified mature differentiated SPs, sharing the surface phenotypes with mature HCECs in fresh CE tissues and with the expression of CD166+, 133−, 105−, 44−, 26−, and 24−, as a candidate cHCEC SP for the cell therapy against CE dysfunctions.

The homeostatic cellular identity of cHCECs may require passive adaptation to environmental culture conditions. Upon exposure to divergent stimuli in cultures, cHCECs dynamically regulate the transcriptional response that results in either differentiation or CST including dedifferentiation. In our culture system, the presence of Y induced the differentiation of cHCECs with reduced expression of CD44.

Consistent with the observation, CD44++/+ immature cHCECs displayed segregated profiles of extracellularly secreted metabolites; the increased amount of lactic acid,
while the decreased amount of pyruvic acid, Ser, and BCAA was evident in mature differentiated cHCEC SPs (Table 1). Not only lactic acid, but also other TCA cycle intermediates, such as malic acid, fumaric acid, cis-aconitic acid, isocitric acid, citric acid, and succinic acid, were secreted increasingly by CD44+++/+++ immature cHCEC SPs, compared with CD44−/+ mature differentiated SPs (Table 1). The increased secretion of TCA cycle intermediates by the former SPs is seemingly incompatible with our previous argument that the latter SPs are disposed to mitochondria dependent oxidative phosphorylation. However, and of interest, most enzymes with mitochondrial-localized reductive activity were upregulated in the differentiated SPs, whereas most with cytosolic reductive activity were upregulated in immature SPs (Fig. 7a), similar to what we found in a recent study on enzymes involved in TCA cycle (Numa et al., unpublished data).

It is of note that the amounts of secreted BCAA and Ser were positively associated with those of secreted lactic acid and pyruvic acid (Figs. 1 and 2). This implies that CD44−/+ SPs catabolize BCAA and Ser more actively than CD44+++/+++ SPs, which may be conceptually rationalized by recent findings.26,27 The elevated BCAA metabolism is more active in differentiated leukemia than in undifferentiated leukemia.26 In actual, BCAT2, a mitochondrial amino-transferase for BCAA and BCKDH, branched chain keto acid dehydrogenase, were upregulated in CD44−/+ SPs (Supplementary Fig. S2). In a fuel-efficient mode more pyruvate is diverted to the mitochondria and more glucose-derived carbon is channeled into serine biosynthesis to support cell proliferation.27 To the contrary negative association of the amount of Gln in the CS with those of secreted lactic acid and pyruvic acid (Fig. 2) may imply that CD44+++/+++ SPs catabolize Gln more actively than CD44− SPs.

Myc-induced alterations in glucose (Glc) and Gln metabolism and overexpressed c-Myc resulted in the concurrent conversion of Glc to lactate and the oxidation of Gln via the TCA cycle28 and these were the case also in our present study in the context of elevated expression of anaplerotic metabolic enzymes in immature SPs (Fig. 7; Table 2). c-Myc
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FIGURE 7. Continued.

reportedly regulates lactate production through transcriptional regulation and is an important regulator for glycolysis.21,22–25 In this regard, it is noteworthy that c-Myc is also regulating CD44 expression,21 which is the key in distinguishing differentiated cHCECs from either immature SPs or SPs with CST. These findings can be interpreted into the negative association of effector (differentiated SP) ratios (E-ratios) of heterogeneous bulk cHCECs with secreted lactic acid and pyruvic acid (Fig. 2). It should be noted that CD44 might be repressed by wild-type p5329 in consistent with our previous finding of upregulated p53 in CD44+ differentiated SPs.30 It should be noted that mutant p53 promotes aerobic glycolysis.29

Given the recent finding that PDGF-β stimulated Glc uptake, lactate production, and ATP production through lactate dehydrogenase A (LDHA)31 and the segregated upregulation of LDHA in CD44++/+/+ differentiated SPs (Fig. 7a), it is conceivable that the amount of PDGF-β in the CS of CD44+ SPs may have the close association with secreted lactate as shown in Figure 4a. The decreased signaling through PDGF-β/PDGFR-B binding (PDGFR-B phosphorylation) resulted in the increased amount of residual PDGF-β and the reduction of secreted lactate in the CS of CD44− SPs (Fig. 4a). The similar association was also confirmed between the amount of residual PDGF-β in the CS and the secreted Ile or Ser in the CS (Fig. 4b). We are currently using PDGF-β and lactic acid in CS as noninvasive biomarkers to qualify the culture process and the content of mature differentiated cHCEC SPs product for cell therapy in clinical settings.

Our present data may indicate that dissecting metabolic processes in cHCEC differentiation will provide essential cues to understand the dynamics of cell fate modulation in cHCECs, because chemical fluxes can affect both transcriptional and epigenetic mechanisms.32 It is a shift from oxidative phosphorylation to aerobic glycolysis characteristic of cancer cells (the Warburg effect in pre-differentiated cells).33–36 Myc is depicted to regulate genes involved in Glc metabolism (e.g., hexokinase 2, pyruvate kinase M2, LDHA, and pyruvate dehydrogenase kinase 1, favoring the conversion of Glc to lactate.37 Enhanced expression of these glycolytic enzymes (Fig. 7a) is a hallmark of oncogenic signaling and transcriptional networks38 or the dedifferentiation into pluripotency with proliferating capability. In glycolysis, most glucose carbon is converted to pyruvate and NADH-dependent reduction of pyruvate to lactate by LDH.39 This reductive reaction recycles the NAD+ reduced to NADH during glycolysis.40

Anaplerosis via glutaminolysis or pyruvate carboxylation, as well as other sources such as fatty acids, contribute to the additional supplies of biosynthetic precursors not met by Glc metabolism.39–41 Gln is converted to α-KG for catabolism through the TCA cycle to malate, which is transported into the cytoplasm and converted to pyruvate and then to lactate.41 Recent report described that glutaminolysis mainly sustains ATP production in terminally-differentiated HCE and accelerated the CE pump function,10 although the authors described that relative contribution of 2 energy sources Glc and Gln into the TCA cycle will require further investigation.
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Table 2. Segregated Expression of Anaplerotic and Cataplerotic Metabolic Enzymes

| Pathway                                      | Gene Symbol | Description                                      | HQ | LQ |
|----------------------------------------------|-------------|--------------------------------------------------|----|----|
| Glycolysis                                   | ALDOA       | Aldolase, fructose-bisphosphate A                 | +  |    |
|                                              | ALDOB       | Aldolase, fructose-bisphosphate B                 | +  |    |
|                                              | ALDOC       | Aldolase, fructose-bisphosphate C                 | +  |    |
|                                              | ENO1        | Enolase 1                                        | +  |    |
|                                              | ENO2        | Enolase 2                                        | +  |    |
|                                              | GAPDH       | Glyceraldehyde-3-phosphate dehydrogenase         | +  |    |
|                                              | GPI         | Glucose-6-phosphate isomerase                    | +  |    |
|                                              | HK2         | Hexokinase 2                                     | +  |    |
|                                              | LDHA        | Lactate dehydrogenase A                          | +  |    |
|                                              | PGAM1       | Phosphoglycerate mutase 1                        | +  |    |
|                                              | PGAM2       | Phosphoglycerate mutase 2                        | +  |    |
|                                              | PGAM        | Phosphoglycerate mutase family member 4          | +  |    |
|                                              | PGM1        | Phosphoglucomutase 1                             | +  |    |
|                                              | PGM2        | Phosphoglucomutase 2                             | +  |    |
|                                              | PKLR        | Pyruvate kinase, liver and RBC                   | +  |    |
|                                              | PKM         | Pyruvate kinase, muscle                          | +  |    |
| Alanine, aspartate, and glutamate metabolism | ABAT        | 4-Aminobutyrate aminotransferase                 | +  |    |
|                                              | ADSL        | Adenylosuccinate lyase                           | +  |    |
|                                              | ADSS        | Adenylosuccinate synthase                        | +  |    |
|                                              | ADSSL1      | Adenylosuccinate synthase-like 1                 | +  |    |
|                                              | ALDH5A1     | Aldehyde dehydrogenase 5 family member A1        | +  |    |
|                                              | ASS1        | Argininosuccinate synthase 1                     | +  |    |
|                                              | CAD         | Carbamoyl-phosphate synthetase 2, aspartate      | +  |    |
|                                              | GFPT1       | Glutamine-fructose-6-phosphate transaminase 1    | +  |    |
|                                              | GFPT2       | Glutamine-fructose-6-phosphate transaminase 2    | +  |    |
|                                              | GLS         | Glutaminase                                      | +  |    |
|                                              | GLS2        | Glutaminase 2                                    | +  |    |
|                                              | GLUD1       | Glutamate dehydrogenase 1                        | +  |    |
|                                              | GLUD2       | Glutamate dehydrogenase 2                        | +  |    |
|                                              | GLUL        | Glutamate-ammonia ligase                         | +  |    |
|                                              | GOT1        | Glutamic-oxaloacetic transaminase 1              | +  |    |
|                                              | GOT2        | Glutamic-oxaloacetic transaminase 2              | +  |    |
|                                              | PPAT        | Phosphoribosyl pyrophosphate amidotransferase    | +  |    |

Genes involved in both Glc and Gln metabolism are regulated by c-Myc and c-Myc overexpressing cells appear to become addicted to Gln.37 A Glc-independent TCA cycle solely supported by Gln involved the supply of acetyl-CoA and oxaloacetic acid (OAA) to the TCA cycle.28 Most of genes involved in Glc metabolism were elevated in immature SPs; however, those involved in Gln were segregated between mature differentiated and immature SPs (Table 2). To dissect the role of anaplerosis via glutaminolysis (TCA cycle or urea cycle) among cHCECs SPs can provide critical insights into how cell fate decisions are established either in vitro or in vivo pathogenesis of HCE disorders. To establish a possible application of the metabolomics included in this study as biomarkers for diagnosis, it would require a very precise analysis of the correlation of the in vitro metabolomics reported in this study with the metabolites in the aqueous humor after the infusion of the cHCECs.

Divergent energetic requirements among heterogeneous cHCECs may implicate either a segregated metabolic profile or distinct mitochondria biogenesis between mature differentiated and immature cHCECs. Further intensive researches in this field will provide the new molecular target for the therapy of CE disorders beyond cell therapy.

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