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Insufficient Activity of MAPK Pathway Is a Key Monitor of Kidney-Yang Deficiency Syndrome

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Abstract

Objective: To explore the genetic characteristics and molecular regulator of Kidney-Yang Deficiency Syndrome (KDS).
Design: A typical KDS family was collected using a questionnaire of cold feeling and a 40-item scoring table of KDS based on Traditional Chinese Medicine (TCM), by single-blind method repeated annually over three years. Their transcriptomes were assayed by microarray and validated by RT-PCR and ELISA. Simultaneously, 10 healthy volunteers were recruited as controls and the same protocols were performed.
Results: This typical KDS family has 35 members, of whom 11 were evaluated as having severe KDS and 6 as having common KDS. Results of the cDNA microarray revealed that there were 420 genes expressed sequence tags differentially expressed in KDS transcriptomes, indicating a global functional impairment in the mass-energy-information carrying network of KDS patients, involving energy metabolism, signal transduction, development, cell cycle, and immunity. Pathway analysis by gene set enrichment assay (GSEA) and other tools demonstrated that mitogenic activated protein kinase (MAPK) is among the most insufficiently activated pathways, while the oxidative phosphorylation and glycolysis/gluconeogenesis pathways, the two main pathways relevant to ATP synthesis, were among the most excessively activated pathways in KDS patients. Results of RT-PCR and ELISA confirmed the status of insufficient activity of the MAPK pathway.
Conclusion: KDS patients undergo overall attenuated functions in the mass-energy-information carrying network. The marked low level of energy output in KDS may be primarily attributed to the insufficient activity of the MAPK pathway, which may be a key monitor for the abnormal energy metabolism and other impaired activities in KDS.

Introduction

Kidney Yang deficiency syndrome (KDS) is one of the primary concepts in Traditional Chinese Medicine (TCM). Although it affects millions of people, little research has been done to understand it. There are relatively few studies of the diversity of body temperature homeostasis, or the relationship between morphology and thermoregulation patterns. The definition of KDS is still rather vague in terms of conventional Western medicine, let alone its genetic background. However, we now have the tools to scan the global features of TCM syndromes with the extensive application of high throughput and genome scale tools such as microarray. TCM by its sheer nature is based on the analysis of various syndromes, and its holistic features and diagnosis methods can be verified by well-described protocols. Therefore, we believe that microarray may be an adequate tool to explore TCM syndromes such as KDS.

Some microarray experiments and relevant analysis designed to discover the basic features of KDS have been reported. Pan and colleagues evaluated a warming and tonic herb treatment for sibling patients from a KDS family with both clinical and molecular approaches to probe a clinical assessment for therapeutic efficacy of TCM. Wang and colleagues studied KDS and cold syndrome through family investigation and suggested that the formation and development of these two syndromes have not only genetic but also environmental factors. Wu and colleagues established a discriminative mathematical model for the diagnosis of deficiency-cold syndrome, a major subtype of KDS, based on a
support vector machine (SVM), and demonstrated that the differentially expressed genes can be used to identify KDS. Yang et al. analyzed the KDS transcriptome module and indicated that KDS might be caused by physiologic imbalance and/or disorder of metabolite processes. However, we still have not discovered the key monitor or critical molecular pathway that can control the extensive disorders of energy and relative mass metabolism in KDS. Such studies will be important to understand the molecular regulators of KDS. In this paper we analyze a typical KDS family and attempt to assay the differentially expressed genes beyond microarray to uncover the molecular monitor of KDS patients.

Materials and Methods

Epidemiological study and primary recruitment of KDS subjects

We first performed a mid-scale epidemiological survey in order to identify research subjects who suffered from KDS in Chengdu, China. Because there is no Western medical definition of KDS, we chose the potential subjects based on a 10-question interview (Table 1). This questionnaire consists of the most frequent complaints of those suffering from unusual coldness, the major symptom of KDS. Subjects were chosen from those who answered “yes” to at least seven of the ten questions in the interview.

Scoring table and quantitative criteria

Potential subjects were diagnosed for KDS by five independent TCM practitioners according to a specific criteria system (GB/T15657-1995) promulgated by the Health Department of China. It includes the main results of TCM diagnosing techniques—interrogation, observation, auscultation, olfaction, and pulse palpation—with clinical manifestation, olfaction, and pulse palpation—with clinical presentations as: none, light, medium, or heavy symptoms. The four main symptoms of KDS are: cold pain of lumbar and knee; profuse urination at night; long voiding of clear urine; and insufficient cubit pulse. Subjects were classified based on their total scores: ≤5 points, healthy individual; 6–11 points, cold constitution; 12–18 points, common KDS; and ≥19 points, severe KDS. Such a standard is consistent with the clinical practice of treating KDS. People with common KDS and severe KDS were chosen for further tests.

Isolation of total RNA

As described previously, 10 mL samples of blood were collected from each subject. Total RNA was extracted using TRIzol reagent (Invitrogen Life Technologies, Carlsbad, CA) and the recommended protocol. All RNA samples were quantified by measuring the absorbance of A260 and A280 nm and followed by gel electrophoresis to ascertain the integrity of total RNA. The purified RNA was then stored in RNAsecure (Ambion, Austin, TX) at −80°C prior to use.

Microarray experiments

Microarray assays were performed with Qiagen chips with 21,400 human 70mer Oligo DNA immobilized by the National Engineering Research Center for Beijing Biochip Technology, China, according to the protocols set by the ScanArray 5000 apparatus, (Packard BioScience) and analyzed using ImAge3.0 software, (BioDiscovery, Inc.).

We designed three kinds of microchips to reciprocally confirm and effectively identify the key genes that may be attributed to KDS. The first chip was to hybridize both the transcriptomes of patients with severe KDS and healthy people, using the latter as the control group. The second was to compare patients with common KDS with the controls. The third chip was to compare a patient with severe KDS with her healthy husband, in order to filter out artifacts coming from differences in age, environment, and nutritional conditions. We conducted all hybridizations twice, with the interchange of both fluorescence Cy5 and Cy3 to decrease experimental error. Genes were identified as differentially expressed if the ratios were ≥2.0 or ≤0.5. To minimize artifacts arising from low expression values, only genes with raw intensity values for bothCy3 and Cy5 of >2000 counts were chosen for further analysis.

To functionally annotate the abnormally expressed genes, we retrieved and integrated information from these public databases: Kyoto Encyclopedia of genes and genomes (www.genome.jp/); UniGene (www.ncbi.nlm.nih.gov/UniGene/) and Gene Ontology (www.godatabase.org/dev/database/archive). At first we linked the differentially expressed genes to the corresponding UniGene clusters. We then retrieved these databases of the biochemical pathways involved in the over- or under-expressed genes. Finally, disorders or diseases were summed and analyzed by means of gene set enrichment assay (GSEA) and other tools.

Validation of the differentially expressed genes by RT-PCR

Semiquantitative RT-PCR was performed with sequence-specific primers to validate the results observed in the microarray experiments. The primer pair for MEK1 was...
forward 5'-CGCTG TAGAAGCACC CATAGAA-3' and back-
ward 5'-TGGGAA ACTGA AGGA T GACGA-3'. The primer 
pair for extracellular signal-regulated kinase (ERK) 2 was 
forward 5'-AGGCGA ATGCCAGTTAC-3' and backward 
5'-GCATA GATATCCCTTTACCA-3'. The PCR produc-
tions were displayed by 1.8% gel electrophoresis and ob-
served in the Gel-Doc system (Bio-Rad, Hercules, CA) and 
the data were analyzed by the attached software.

Assay activity of two representative members of MAPK pathway by ELISA

Two ELISA kits, MEK1 (pSer218/222) (CBA030) and 
ERK1/2 (Thr202/Tyr204) (CBA032) (Calbiochem, San Diego, CA) were used. Following the manufacturer’s protocols, we assayed the activities of two important MAPK members of every sample. The relative concentration of phosphorylated amino acid in specific sites of certain enzymes represent its relative activity. Statistical differences were measured by t test.

Results

Questionnaire and TCM diagnosis

Using the questionnaire mentioned above, we obtained 82 potential research subjects, 22 of whom belong to two families. One of the KDS pedigrees live in Xingxin town of Peng Zhou City, Sichuan Province, China, and have been inten-
sively studied.6,8–10 Another family, which has 17 KDS patients and lives around Chengdu, Sichuan, China, was chosen as the subject for this present study (Fig. 1). Simul-
taneously, we recruited 10 healthy volunteers as control subjects. The control subjects’ economic condition, living environment, age, and gender ratio matched the members of the KDS family.

Reliability of diagnosis results

In order to improve the reliability of TCM diagnosis, we invited 5 experts in TCM differentiation to independently evaluate all subjects in this family. Almost all of the results recorded by the 5 physicians were consistent (Fig. 2). The third physician may be stricter than the first and fifth physician, but there was no statistically significant difference among them. As the result, 11 individuals dispersed in generations II, III, and IV were evaluated as having severe KDS; 6 as having common KDS; and 6 as having cold constitution or as healthy individuals. The rest of the subjects were not evaluated due to death, nonavailability, or being too young.

We repeated the diagnosis on the same participants once a year for three years. Diagnosis in all three years was con-
ducted on the first Saturday of December. Results revealed that every family member can be classified into the same KDS subtype in each year. For instance, in 2005 the average score of severe KDS within the pedigree was as high as 40.63 points, which was markedly high than that of the common KDS (15.38 points), cold constitution (6.89 points), or control group (3.24 points). Furthermore, the scores of every subtype of the KDS family were rather stable from 2003 through 2005, indicating the reliability of the subjects involved and the validity of the employed TCM criteria.

Abnormally expressed genes related to energy metabolism as the major contributors in KDS transcriptome

Results of the cDNA microarray experiments revealed that there were 420 differentially expressed genes expressed sequence tags (EST) between KDS patients and the control group. Fully one third of these differentially expressed genes are related to energy metabolism. The remaining differentially expressed genes are associated with signal transduction, transport, development, cell cycle, and response to stress (Fig. 3). Concomitantly with the abnormally low level of energy output observed in KDS subjects, many other major functions such as signal transduction, mass transport, development, reproduction, metabolism, and immu-

nity were found to be disturbed. In short, the results of our bioinformation analysis suggest the disturbance of the KDS transcriptome affects the mass-energy-information carrying network.
A higher ratio of reappeared genes abnormally expressed among different microchips was observed in the pedigree.

It is interesting that the family data obtained from microarray analysis revealed an extraordinarily high frequency of reappeared genes abnormally expressed among various chips. Reappeared genes (those observed at ≥2 chips as being abnormally expressed) and abnormally expressed genes among various microchips made up 12% of all differentially expressed genes. For example, two genes, NM_006158 and NM_005252, were all insufficiently expressed in all of the tested chips while two genes, NM_053025 and NM_002619, were all excessively expressed in all chips. This is in contrast to similar microarray studies on non-pedigree KDS patients, where there were few such reappeared genes observed in different chips. This indicates that pedigree can strongly facilitate KDS research.

Importance of the MAPK pathway in KDS patients

Pathway analysis showed that MAPK was the most common pathway to be insufficiently expressed in the KDS transcriptome (Table 2). Specifically, there were as many as 15 genes involved in the MAPK pathway observed among the abnormally expressed genes from KDS transcriptomes. Unexpectedly, genes associated with ATP synthesis pathways were almost all excessively expressed in KDS patients. For instance, genes associated with the oxidative phosphorylation pathway, playing a central role in ATP synthesis, were the most excessively expressed genes in our survey. The up-regulation of oxidative phosphorylation together with glycolysis/gluconeogenesis might act as a kind of feedback mechanism prompted by the low level of energy production in KDS.

We then compared the differentially expressed gene profiles using GSEA. The results show that only subjects with severe KDS demonstrated different gene sets than those of the healthy control group. The MAPK gene set was among those that displayed differences. MAPK is a highly conserved signal-transduction pathway, which plays an important role in the regulation of energy metabolism and thermogenesis. Therefore, the extensive abnormalities of the MAPK pathway may be a factor in the low energy level exhibited by KDS patients (Fig. 4). Pathway analysis suggests that it is not the ATP synthesis system itself that is responsible for the low level of energy output in KDS patient; rather the main monitors, such as MAPK, are to blame.

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<th>Table 2. Pathways Related to the Abnormally Expressed Genes of the Kidney-Yang Deficiency Syndrome Pedigree</th>
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<td><strong>P Insufficiently expressed pathways</strong></td>
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<td>MAPK signaling pathway</td>
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<td>Amyotrophic lateral sclerosis</td>
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<td>Complement and coagulation cascades</td>
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<td>Purine and pyrimidine metabolism</td>
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<td>Starch and sucrose metabolism</td>
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<td>N-glycans biosynthesis</td>
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<td>Folate biosynthesis</td>
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<td>Nicotinate and nicotinamide metabolism</td>
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Results of RT-PCR and ELISA assays

RT-PCR results showed that both ERK2 and MEK1 genes were insufficiently represented in KDS patients (Fig. 5). Similarly, results of ELISA assay showed that the activities of MEK1 and ERK1 were significantly lower than in the control group. This confirms that in the microarray experiments at both the transcript and protein levels, insufficient activity of the MAPK pathway in KDS were ubiquitous.

Discussion

TCM diagnosis can distinguish KDS subtypes effectively

Classifying TCM syndromes (Zheng) under the criteria of evidence-based medicine is an important problem. It is characterized by extremely complex elements such as genetics, environment, and disease history. Even though techniques and approaches from multiple disciplines have
been brought to bear on the subject, classifying TCM syndromes remains a challenge to practitioners of alternative medicine.

We designed our studies based on the principle rules of evidence-based medicine. We used the questionnaire described earlier as a primary screening tool to obtain 82 potential research subjects. Next, we employed 5 experts to identify the KDS patients via a single-blind method. Then, we performed the TCM differentiation three times on the same subjects in order to confirm the validity and reliability of our TCM diagnosis. Finally, we recruited healthy control subjects whose living environment, economic condition, age, and gender ratios were similar to those of the KDS subjects. The results of this research indicate that the time-honored TCM criteria for KDS not only effectively distinguishes KDS patients from healthy people, but also classifies them into various levels of KDS (severe KDS, common KDS, and KDS constitution). The 40-item criteria of KDS diagnosis not only reveals the chronic chill, the major symptom of KDS, but also elicits impaired physiological functions associated with KDS, such as circulation, respiration, digestion, mass and water metabolism, and immunological functions. Therefore, TCM diagnosis for KDS puts chronic chill and low body temperature as revealed by our questionnaire in a much more integrated framework. This serves as the basis for detailed investigation of KDS at a molecular level. Also, the procedures we used for subject recruitment provide us with a substantial number of subjects for the genetic study of KDS.

**Pedigree: An important and convenient approach to the study of physical constitution**

The pedigree approach is well accepted among researchers of human genetics. Most disease genes have been directly isolated from pedigrees. Genetics can also be the key factor responsible for myriad TCM syndromes. Likewise, the pedigree method is useful for further understanding TCM syndromes. Compared with similar studies published by our group, we report here on a much more typical KDS family. It has a higher ratio of KDS patients, more family members, and shares more symptoms than those described in previous studies. Furthermore, almost all KDS patients in the family can be effectively treated by You-Gui-Wan, a classical TCM formula for the treatment of KDS (data not shown). Our research strongly suggests that inheritance plays a key role in KDS. Therefore, the pedigree approach to researching KDS could be effective in understanding this TCM syndrome.

Microarray results of the KDS family also confirmed our assumption that pedigree is a useful approach for revealing
the traits of TCM syndromes. For instance, compared to the microarray studies derived from non-pedigree KDS patients, the frequency of reappeared genes was markedly higher among pedigree KDS patients. This might account for the fact that characteristic genes are heaped up and the expression pattern of certain genes are repeatedly displayed among KDS patients within a family. In our opinion, TCM syndromes may share some features with complex diseases (such as hypertension and diabetes) in their genetic background: all are attributed to multiple genes, show diverse etiopathogenesis, and also develop through various molecular pathways. To this point we have obtained less success in complex diseases than with single gene diseases. However, the pedigree approach is a useful way to study complex diseases. And the pedigree method can also facilitate our further exploration of KDS biomarkers.

KDS transcriptome focuses on energy metabolism

The results of the microarray tests showed that abnormally expressed genes involved in energy metabolism are significant in the KDS transcriptome and in the pathways related to mass and energy metabolism. Similar results have been observed by others. Shen and colleagues studied the gene networks and signal transduction pathways in a KDS rat model, and found KDS covered the neuroendocrine-immune (NEI) network as a whole-genome impairment. They discovered significant down-regulation of various neurotransmitters of the hypothalamus–pituitary–adrenal–thymus (HPAT) axis, followed by those of the growth and sex hormone related genes. We also observed many abnormally expressed genes whose products include regulation factors involved in energy metabolism, including a dozen transcription regulators, interleukin 8 (IL-8) mediated inflammatory regulator, and tissue factor pathway inhibitor for blood coagulation.

Although a defined pattern of dysfunction transcriptome has not yet been obtained, we can assume that the marked low level of energy metabolism in KDS may be the leading indication of an overall imbalance of the KDS transcriptome, which gradually intensifies the cold feeling and attenuating physiological functions as a whole. Consequently, KDS transcriptome obtained from our experiments revealed that KDS patients suffer from extensive impairments involving large parts of the mass-energy-information carrying network, while the low energy expenditure is merely one of the distinctive features, yet a marked symptom, in KDS patients.

The MAPK pathway in energy metabolism

The most important discovery presented in this paper might be the universal abnormal expression of certain members of the MAPK pathway. MAPK, one of the critical joint points for many signal transduction pathways (i.e., PKC, RPTK, JAK-STAT), is a key monitor of energy metabolism by regulating the concentration of blood glucose, manipulating the glucose and lipid metabolism, controlling mitochondrial function, and regulating hormonal response for energy output. As described above, abnormally low levels of energy output is the distinguishing symptom of KDS. Hence, in theory, MAPK should suffer some kind of impairment in KDS. Results of pathway analysis indicate the universally abnormal expression of MAPK might be the critical mechanism of insufficient energy output in KDS. The GSEA results confirmed that MAPK is among the signal transduction pathways significantly down-regulated in KDS. Results of both RT-PCR and ELISA assay on MEK1 and ERK2 reconfirmed the fact that some members of the MAPK pathway in KDS patients were insufficiently active.

Shen and colleagues found in a KDS rat model that the NF-kappaB signal transduction pathway exerts a hinge role in the molecule regulating network. The intensive interaction among mitochondria, MAPK, and NF-kappaB has been reported. As a pre-inflammation cytokine, NF-kappaB may react more quickly than MAPK in the process of acute modeling of KDS. MAPK, as a predominant hinge and signal transduction highway within all kinds of cells, may play a much more important role in lengthy clinical pathogenesis and TCM syndromes (such as KDS) than does NF-kappaB. Therefore, in our opinion, both MAPK and NF-kappaB are major monitors for energy metabolism, yet they alternatively monitor different TCM syndromes or diverse stages of identical syndromes.

Thermoregulation in human beings and animals is a process with high complexity, integrating a large number of mechanisms and pathways. The extensive insufficient activities of MAPK, and the universal excessive expression of genes involved in ATP synthesis pathways led to our hypothesis that it is not ATP synthesis itself but the critical monitor of energy metabolism such as MAPK that may be responsible for the abnormally low level of energy metabolism in KDS patients.

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Disclosure Statement

No competing financial interests exist in this article.

References


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