

Research Article

Bioinformatic analysis of key pathways and genes involved in pediatric atopic dermatitis

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The initiation of atopic dermatitis (AD) typically happens very early in life, but most of our understanding of AD is derived from studies on AD patients in adult. The aim of the present study was to identify gene signature specific to pediatric AD compared with adult AD. The gene expression profiles of four datasets (GSE32924, GSE36842, GSE58558, and GSE107361) were downloaded from the GEO database. Gene ontology (GO) and Kyoto Encyclopedia of Genes and Genomes pathway (KEGG) enrichment analyses were performed, and protein–protein interaction (PPI) network was constructed by Cytoscape software. Total 654 differentially expressed genes (DEGs) (394 up-regulated and 260 down-regulated) were identified in pediatric AD samples with adult AD samples as control. The up-regulated DEGs were significantly enriched in the migration and chemotaxis of granulocyte and neutrophil, while down-regulated DEGs were significantly enriched in biological adhesion. KEGG pathway analysis showed that up-regulated DEGs participated in chemokine signaling pathway while down-regulated DEGs participated in adherens junction, focal adhesion, and regulation of actin cytoskeleton. The top 10 hub genes GAPDH, EGFR, ACTB, ESR1, CDK1, CXCL8, CD44, KRAS, PTGS2, and SMC3 were involved in chemokine signaling pathway, cytokine–cytokine receptor interaction, interleukin-17 signaling pathway, and regulation of actin cytoskeleton. In conclusion, we identified DEGs and hub genes involved in pediatric AD, which might be used as therapeutic targets and diagnostic biomarkers for pediatric AD.

Introduction

Atopic dermatitis (AD) is the most common inflammatory skin disease with an estimated prevalence of around 20% in children and 7–10% in adults [1–4]. AD is predominantly a Th2/Th22 polarized disease with Th1 polarization in the chronic phase and the impairment of Th17 pathway [5]. The initiation of AD typically happens very early in life, but most of our understanding of AD is derived from studies on AD patients in adult. Therefore, the molecular mechanism underlying pediatric AD initiation and progression is elusive, resulting in a lack of specific treatment for this disease.

Bioinformatics analysis of microarray data is increasingly valued as a promising tool in gene expression profiling in inflammatory diseases to identify differentially expressed genes (DEGs) that play important role in the diseases [6–8]. However, comparative analysis of the DEGs between pediatric AD and adult AD remains to be elucidated.

The aim of the present study was to explore gene signature of pediatric AD and identify differentially expressed genes involved in pediatric AD compared with adult AD. In present study, we download the original data (GSE32924, GSE36842, GSE58558, and GSE107361) from Gene Expression Omnibus and compared gene expression profiles of pediatric AD with those in adult AD. The DEGs were identified and analyzed by gene ontology (GO) and pathway enrichment analysis.

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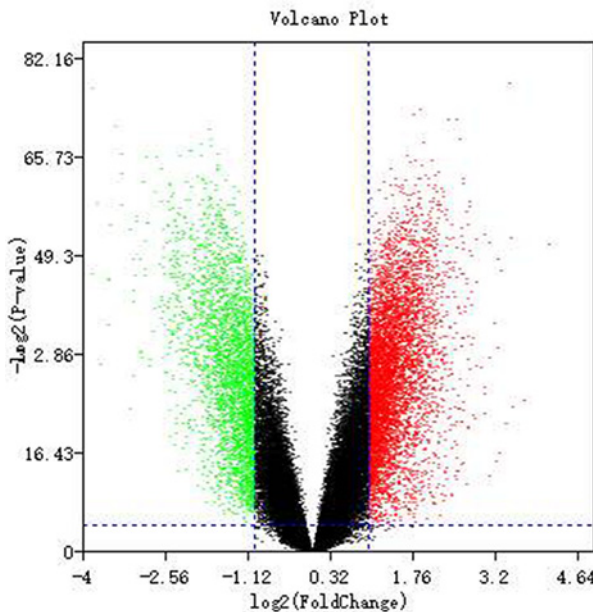


Figure 1. Volcano plots of genes that are significantly different between pediatric and adult controls

The X-axis indicates the *P* values (log scale), whereas the Y-axis shows the fold change (log scale). Each symbol represents a different gene, and the red/green color of the symbols categorize the up-regulated/down-regulated genes falling under different criteria (*P*-value and fold change threshold). *P*-value <0.01 is considered as statistically significant, whereas fold change = 2 is set as the threshold.

Materials and methods

Identification of DEGs

From the Gene Expression Omnibus (<http://www.ncbi.nlm.nih.gov/geo/>), four gene expression profiles (GSE32924, GSE36842, GSE58558, and GSE107361) were selected because they were on gene expression profiling of AD samples (total 49 adult AD samples versus 19 pediatric AD samples) based on Affymetrix GPL570 platform [9–12]. The original probe-level data were converted into gene-level data using Robust multi-array average (RMA) approach for background correction and normalization. Next, limma package in R language was used to identify the DEGs between pediatric and adult samples. Subsequently, a between-subjects *t*-test was performed to identify DEGs of each AD group with the cutoff criteria of log₂ fold change (FC) >2 and FDR <0.01. Volcano plots were generated to visualize the distribution of DEGs between pediatric and adult samples of AD patients.

Gene ontology and pathway enrichment analysis of DEGs

Bioinformatics analysis of the DEGs was performed as described previously [13].

Gene ontology (GO) and Kyoto Encyclopedia of Genes and Genomes (KEGG) analysis were performed by employing an online software DAVID Database (<https://david.ncifcrf.gov/>). *P*<0.05 was considered statistically significant.

Integration of protein–protein interaction network

STRING online database (<http://string-db.org>) was used for analyzing the protein–protein interaction (PPI) information. The cut-off criteria were a combined score of > 0.4 for a PPI network and a node degree of > 10 for screening hub genes. Cytoscape MCODE plug-in was used for searching clustered sub-networks. The default parameters were as follows: degree cutoff ≥10, node score cutoff ≥0.2, *K*-core ≥2, and max depth = 100.

Results

Identification of DEGs

A total of 654 genes (394 were up-regulated and 260 were down-regulated) special to pediatric AD samples were identified after the analyses in all four independent cohorts with adult AD samples as control (Supplementary Tables

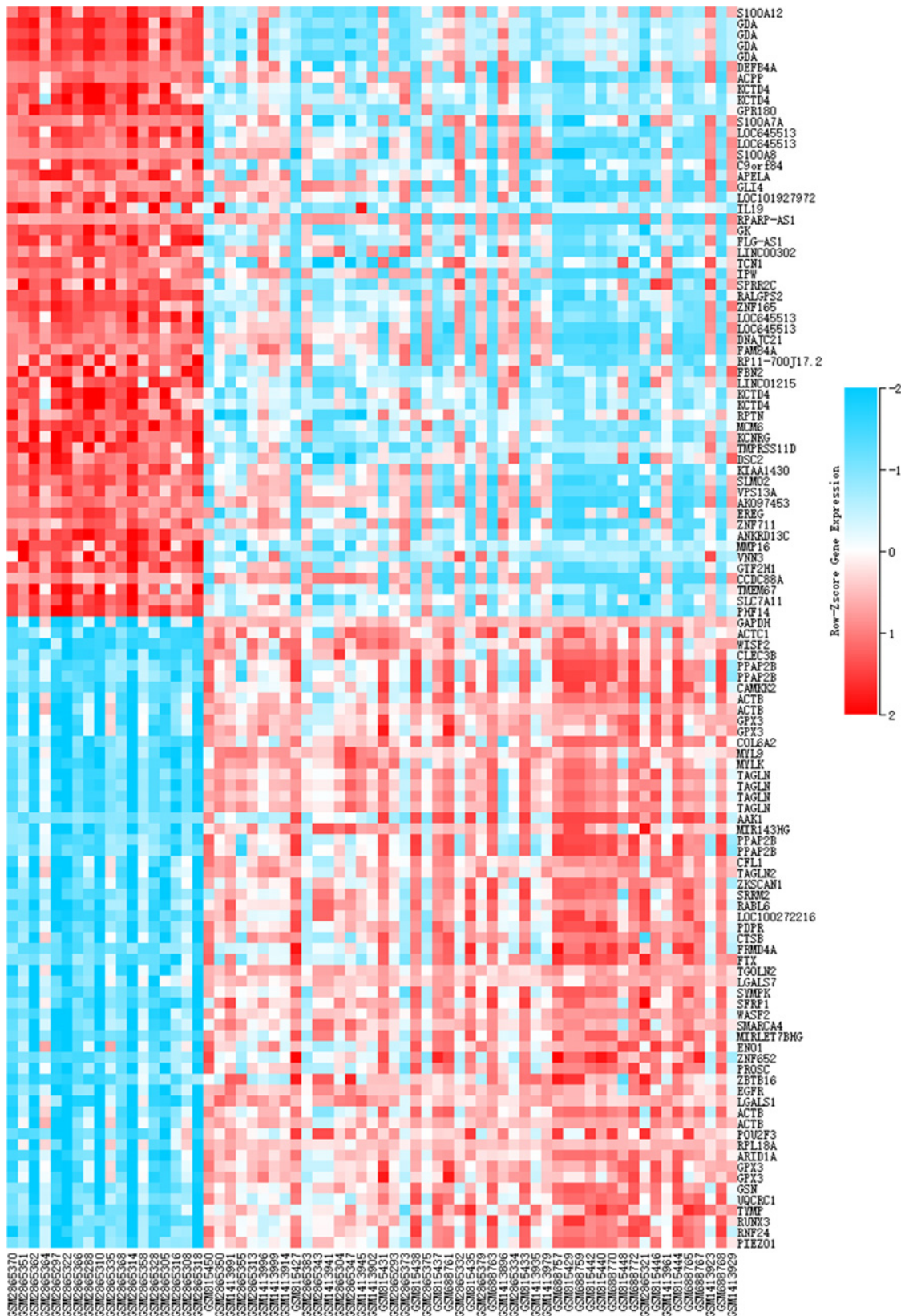


Figure 2. Heat map of the top 100 differentially expressed genes

Shown were 50 up-regulated genes and 50 down-regulated genes. Each column represented a biological sample and each row in the heat map represents a gene. Red: up-regulation; Blue: down-regulation.

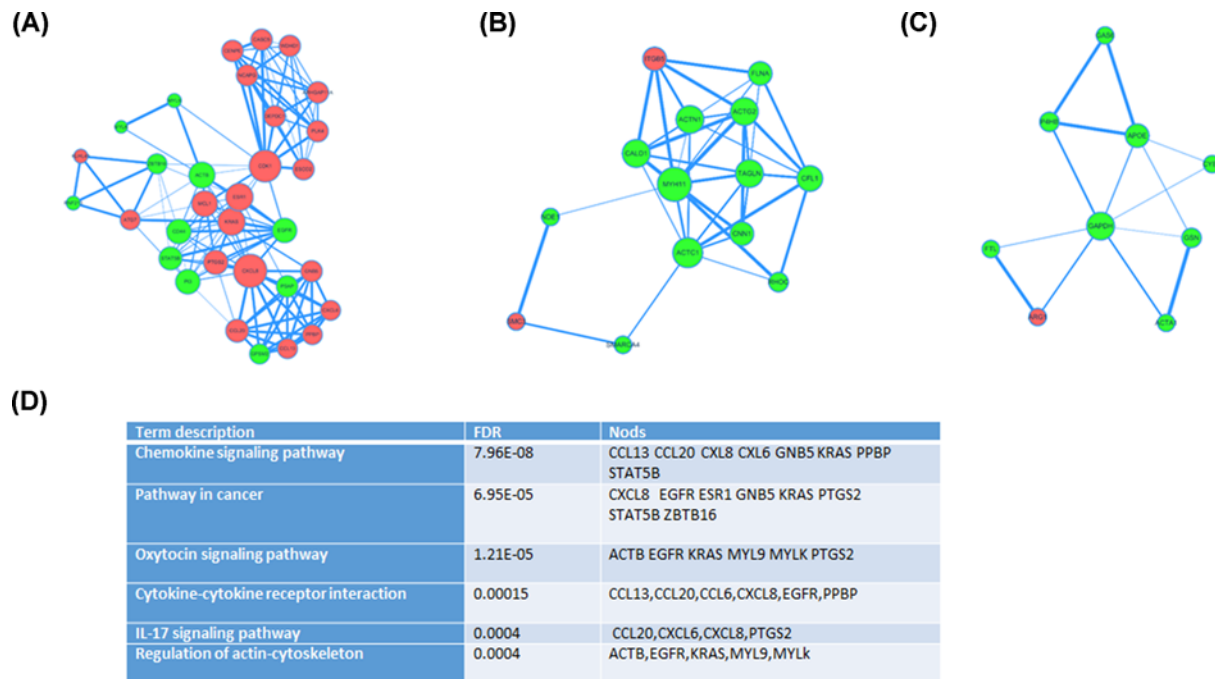


Figure 3. Top 3 modules from the protein-protein interaction network

(A) module 1, (B) module 2, (C) module 3. Red: up-regulation; green: down-regulation. (D) the enriched pathways of the three modules (FDR < 0.0005).

S1 and S2). Red or green dots in the volcano plots represented significantly up- or down-regulated genes, respectively (Figure 1). The top 50 up- and down-regulated genes were shown in the heat map (Figure 2).

Functional and pathway enrichment analyses

We uploaded all DEGs to the online software DAVID to identify over-represented GO categories and KEGG pathways. GO term enrichment analysis showed that up-regulated DEGs were significantly enriched in the migration and chemotaxis of granulocyte and Neutrophil, while down-regulated DEGs were mainly involved in a multi-organism process. In addition, molecular function analysis showed that up-regulated DEGs were mainly associated with chemokine activity, while down-regulated DEGs were involved in protein binding (Table 1). Furthermore, KEGG pathway analysis showed that up-regulated DEGs participated in the chemokine signalling pathway while down-regulated DEGs participated in adherens junction, focal adhesion, and regulation of actin cytoskeleton (Table 2).

Protein–protein interaction network construction and analysis of modules

Based on the information in the STRING database, the top 10 hub nodes with higher degrees were screened (Table 3). Among these nodes, GAPDH showed the highest degree. A total of 594 nodes and 1651 edges were analyzed using plug-ins MCODE. The top 3 significant modules were selected, the functional annotation of the protein involved in the modules was summarized. Enrichment analysis showed that the proteins in modules 1–3 were mainly associated with the chemokine signalling pathway, Pathway in cancer, Oxytocin signalling pathway (Figure 3).

Discussion

Understanding of the molecular mechanism of pediatric AD might help develop approaches that can prevent atopic diathesis [14]. Previous studies have compared gene expression profiling of pediatric AD samples with adult AD samples or normal healthy samples, respectively, but the sample size of the individual study was limited and the conclusion was controversial [9–12]. Therefore, in the present study we retrieved gene expression data of 19 pediatric AD samples and 49 adult AD samples from previous studies and identified 654 DEGs in pediatric AD samples, among which 394 were up-regulated and 260 were down-regulated. Cumulative evidence has demonstrated that the co-expressed genes

Table 1 Gene ontology analysis of DEGs associated with pediatric AD

Category	Term	Involved in	n*	%	P
Up-regulated					
GOTERM_BP_FAT	GO:0097530	granulocyte migration	9	2.3	1.32E-03
GOTERM_BP_FAT	GO:0006275	regulation of DNA replication	9	2.3	1.56E-03
GOTERM_BP_FAT	GO:1990266	neutrophil migration	8	2.0	1.86E-03
GOTERM_BP_FAT	GO:0071621	granulocyte chemotaxis	8	2.0	3.13E-03
GOTERM_BP_FAT	GO:0030593	neutrophil chemotaxis	7	1.8	4.78E-03
GOTERM_CC_FAT	GO:0005615	extracellular space	40	10.2	9.77E-03
GOTERM_CC_FAT	GO:0098687	chromosomal region	13	3.3	2.74E-02
GOTERM_MF_FAT	GO:0005125	cytokine activity	13	3.3	1.03E-03
GOTERM_MF_FAT	GO:0042379	chemokine receptor binding	7	1.8	1.16E-03
GOTERM_MF_FAT	GO:0008009	chemokine activity	6	1.5	2.20E-03
GOTERM_MF_FAT	GO:0016791	phosphatase activity	12	3.1	1.71E-02
GOTERM_MF_FAT	GO:0016810	hydrolase activity, acting on carbon–nitrogen bonds	8	2.0	2.12E-02
Down-regulated					
GOTERM_BP_FAT	GO:0016032	viral process	32	12.3	4.97E-06
GOTERM_BP_FAT	GO:0044764	multiorganism cellular process	32	12.3	5.75E-06
GOTERM_BP_FAT	GO:0022610	biological adhesion	46	17.7	5.76E-06
GOTERM_BP_FAT	GO:0044403	symbiosis, encompassing mutualism through parasitism	32	12.3	9.55E-06
GOTERM_BP_FAT	GO:0044419	interspecies interaction between organisms	32	12.3	9.55E-06
GOTERM_CC_FAT	GO:0005912	adherens junction	37	14.2	8.19E-12
GOTERM_CC_FAT	GO:0070161	anchoring junction	37	14.2	1.64E-11
GOTERM_CC_FAT	GO:0070062	extracellular exosome	73	28.1	5.49E-08
GOTERM_CC_FAT	GO:1903561	extracellular vesicle	73	28.1	6.76E-08
GOTERM_CC_FAT	GO:0043230	extracellular organelle	73	28.1	6.86E-08
GOTERM_MF_FAT	GO:0008092	cytoskeletal protein binding	31	11.9	1.30E-06
GOTERM_MF_FAT	GO:0032403	protein complex binding	29	11.2	1.70E-06
GOTERM_MF_FAT	GO:0050839	cell adhesion molecule binding	19	7.3	5.91E-05
GOTERM_MF_FAT	GO:0044877	macromolecular complex binding	36	13.8	7.47E-05
GOTERM_MF_FAT	GO:0098641	cadherin binding involved in cell-cell adhesion	14	5.4	1.70E-04

*Number of enriched genes in each term. If there were more than five terms enriched in this category, the top five terms based on P value were chosen.

normally consist of a group of genes with similar expression profiles and participate in parallel biological process. To better understand the interactions of DEGs, we further performed GO, KEGG pathway, and PPI network analysis.

GO analysis showed that DEGs mainly participated in extracellular space, anchoring junction and adherens junction, involved in granulocyte and neutrophil migration, performed functions of cytokine activity, chemokine receptor binding, chemokine activity, and cytoskeletal protein binding. Furthermore, enriched KEGG pathways of up-regulated DEGs included chemokine signaling pathway and cytokine–cytokine receptor interaction, and those of down-regulated DEGs included adherens junction, focal adhesion, and regulation of actin cytoskeleton. Therefore, all these pathways could contribute to the pathogenesis of pediatric AD.

The analysis based on PPI networks indicated that GAPDH, EGFR, ACTB showed the highest betweenness and belonged to crucial modules of the PPI network. GAPDH is a classic glycolytic enzyme involved in membrane transport and membrane-fusion, microtubule assembly, nuclear RNA export, protein phosphotransferase/kinase reactions, and translational control of gene expression [15]. The β-actin cytoskeleton functions in cellular shape and anchorage where transmembrane glycoproteins link fibronectin in the extracellular matrix with actin microfilaments on the

Table 2 KEGG pathway analysis of DEGs associated with AD

Category		Term	Count*	%	P
Up-regulated					
KEGG_PATHWAY	hsa04062	Chemokine signaling pathway	8	2.0	0.044
KEGG_PATHWAY	hsa04060	Cytokine–cytokine receptor interaction	9	2.3	0.061
KEGG_PATHWAY	hsa04064	NF-kappa B signaling pathway	5	1.3	0.065
KEGG_PATHWAY	hsa04012	ErbB signaling pathway	5	1.3	0.065
KEGG_PATHWAY	hsa05323	Rheumatoid arthritis	5	1.3	0.067
Down-Regulated					
KEGG_PATHWAY	hsa04520	Adherens junction	6	2.3	0.004
KEGG_PATHWAY	hsa04510	Focal adhesion	9	3.5	0.013
KEGG_PATHWAY	hsa04810	Regulation of actin cytoskeleton	9	3.5	0.015
KEGG_PATHWAY	hsa04530	Tight junction	5	1.9	0.044
KEGG_PATHWAY	hsa04512	ECM–receptor interaction	5	1.9	0.044

*Count: the number of enriched genes in each term. If there were more than five terms enriched in this category, the top five terms based on P value were chosen.

Table 3 The top 10 hub nodes in protein–protein interaction network

Hub node	Information	Degree
GAPDH	Glyceraldehyde-3-phosphate dehydrogenase	89
EGFR	Epidermal growth factor receptor	69
ACTB	Actin, cytoplasmic 1	51
ESR1	Estrogen receptor	46
CDK1	Cyclin-dependent kinase1	44
CXCL8	Interleukin-8	43
CD44	CD44 antigen	41
KRAS	GTPase Kras	36
PTGS2	Prostaglandin G/H synthase 2	33
SMC3	Structural maintenance of chromosomes protein 3	27

cytoplasmic side of the membrane [16]. While GAPDH and β-actin are regarded as housekeeping genes, accumulating evidence has suggested their mRNA levels vary with cellular proliferation [17–21]. Moreover, their transcription is up-regulated rapidly in response to mitogenic stimuli including epidermal growth factor, transforming growth factor-β, and platelet-derived growth factor [22–24]. We hypothesized that β-actin and GAPDH expression levels in AD were variable and not suitable for normalizing mRNA levels. Our results were similar to some studies in asthma, which was part of the atopic march [25].

Epidermal growth factor receptor (EGFR) is a large transmembrane glycoprotein with ligand-induced tyrosine kinase activity [26]. Inhibition of EGFR signaling leads to decreased expression of cytoskeleton proteins such as actin-binding protein ACTN1 (actinin-1), increased keratinocyte adhesion, resulting in the inhibition of the migration of keratinocytes from the basal layer to the stratum corneum [27–30]. Blockade of EGFR signaling can regulate the expression of CCL26/eotaxin-3 in primary keratinocytes in AD [31,32].

In summary, we identified genes differentially expressed in pediatric AD compared with adult AD and explored their potential function and relevant pathways in the pathogenesis of pediatric AD. Moreover, our study suggested that chemokine pathway and cytoskeletal protein binding play a vital role in the molecular mechanism of pediatric AD. However, the present study has limitation because it is based on bioinformatic analysis of online datasets and the differentially expressed genes in pediatric AD should be validated by real-time PCR analysis and function assay. In particular, further studies are needed to validate GAPDH, EGFR and ACTB, which can be considered as crucial genes involved in pediatric AD, with the potential to be used in the diagnosis and therapy.

Data Availability

All data are available upon request.

Competing Interests

The authors declare that there are no competing interests associated with the manuscript.

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Author Contribution

C.Z. designed the study. T.W., B.Z., D.L., and X.Q. collected and analyzed the data. All authors read and approved the manuscript.

Ethics Approval

No ethics statement was required because this study involved no human or animals.

Abbreviations

AD, atopic dermatitis; DEG, differentially expressed gene; EGFR, epidermal growth factor receptor; GO, Gene ontology; KEGG, Kyoto Encyclopedia of Genes and Genomes; PPI, protein–protein interaction; RMA, Robust multi-array average.

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Supplemental table 1. Downregulated 260 genes.

Gene Name	ENTREZ_GENE_ID	AFFY_ID
AGPAT3	56894	223182_S_AT
ABHD17A	81926	221267_S_AT
ASMTL	8623	36554_AT
ACTB	60	AFFX-HSAC07/X00351_M_AT AFFX-HSAC07/X00351_5_AT
ACTA1	58	203872_AT
ACTC1	70	205132_AT
ACTG2	72	202274_AT
ACTN1	87	208637_X_AT
AP2S1	1175	208074_S_AT
ADD3	120	201753_S_AT
ADGRA2	25960	221814_AT
AKAP1	8165	201674_S_AT
AKAP8L	26993	218064_S_AT
ALDH3A2	224	210544_S_AT
ALKBH7	84266	223318_S_AT
ANKRD10-IT1	100505494	226663_AT
AAK1	22848	205434_S_AT
APOE	348	203381_S_AT/203382_S_AT
AQP5	362	213611_AT
ASS1	445	207076_S_AT
ABCC3	8714	230682_X_AT
ATP5G2	517	208764_S_AT
ATP2A3	489	207522_S_AT/213036_X_AT
ARID1A	8289	210649_S_AT/212152_X_AT
ATG16L2	89849	229389_AT
BCL6	604	228758_AT
BCL9L	283149	227616_AT
BMPR2	659	231873_AT
BAP1	8314	1555735_A_AT
BRD2	6046	208685_X_AT/214911_S_AT
BRD3	8019	203825_AT
BRD4	23476	202102_S_AT
BTG1	694	200920_S_AT
BTNL9	153579	229985_AT
CAMKK2	10645	213812_S_AT
CALD1	800	215199_AT
CNN1	1264	203951_AT
CA12	771	210735_S_AT/203963_AT
CARMN	728264	1558828_S_AT
COMP	1311	205713_S_AT
CSNK1A1	1452	240221_AT
CTSB	1508	227961_AT
CCZ1B	221960	215024_AT
CD248	57124	219025_AT
CD44	960	210916_S_AT
CD81	975	200675_AT
CD82	3732	203904_X_AT
CD99	4267	201028_S_AT
CLN6	54982	1567080_S_AT
CBX6	23466	202047_S_AT
CHD4	1108	201183_S_AT

C10orf10	11067	209183_S_AT
C12orf57	113246	224719_S_AT
CLOCK	9575	204980_AT
CFL1	1072	1555730_A_AT
CCDC85B	11007	204610_S_AT
COL1A2	1278	229218_AT
COL6A2	1292	209156_S_AT
COL7A1	1294	217312_S_AT
COL17A1	1308	204636_AT
CFDP1	10428	210701_AT
CTDSPL	10217	201904_S_AT
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CXCL14	9547	237038_AT
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DLEU2	8847	1556821_X_AT
DES	1674	214027_X_AT/202222_S_AT
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DIXDC1	85458	214724_AT
DLGAP4	22839	1557394_AT
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ENO1	2023	217294_S_AT
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EIF5	1983	241843_AT
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FASN	2194	212218_S_AT
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FBXO17	115290	220233_AT
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FTL	2512	213187_X_AT
FGFR1	2260	210973_S_AT
FNDC3B	64778	218618_S_AT
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FLII	2314	222065_S_AT
FTX	100302692	1558515_AT
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LGALS1	3956	201105_AT
LGALS7	3963	206400_AT
GATAD1	57798	214718_AT
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HSPB7	27129	218934_S_AT
HGSNAT	138050	1557064_S_AT
HCRP1	387535	216176_AT
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LYRM9	201229	1560703_AT
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LAPTM5	7805	201721_S_AT
MIF	4282	217871_S_AT
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ZBTB16	7704	205883_AT
ZNF551	90233	211721_S_AT
ZNF652	22834	205594_AT
ZNF91	7644	206059_AT
ZKSCAN1	7586	214670_AT
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Supplemental table 2. Upregulated 394 genes.

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AP4S1	11154	210277_AT
ADGRF1	266977	236489_AT
ARL14	80117	220468_AT
ARL5B	221079	242727_AT
AKR1B10	57016	206561_S_AT
AGPS	8540	225114_AT
AREG	374	205239_AT
ANGEL2	90806	217630_AT
ANKRD13C	81573	1556361_S_AT
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APOL6	80830	1557236_AT
APOOL	139322	222269_AT
APLF	200558	241379_AT
ARG1	383	231662_AT
ARG2	384	203946_S_AT
ARMC8	25852	1555281_X_AT 1555279_AT
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ATP11B	23200	1554557_AT 1564063_A_AT 1554556_A_AT
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ATG7	10533	1569827_AT
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BMP2K	55589	214716_AT
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BROX	148362	241908_AT
CDH26	60437	232306_AT
CALCRL	10203	234996_AT
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CAPN6	827	202965_S_AT
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CATSPERB	79820	220293_AT
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CCL20	6364	205476_AT
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C12orf54	121273	240353_S_AT
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C15orf65	145788	243309_AT
C3orf33	285315	1554176_A_AT
C4orf47	441054	236915_AT
C7orf57	136288	1557636_A_AT
C9orf84	158401	233504_AT
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DEFB4B	100289462	207356_AT
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DDIAS	220042	228281_AT
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EGR1	1958	227404_S_AT
ENTPD6	955	234946_AT
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ESCO2	157570	235178_X_AT
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HYAL4	23553	220249_AT
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IL24	11009	206569_AT
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IFT74	80173	61732_R_AT
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KLHL7	55975	220239_AT 220238_S_AT
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KRIT1	889	229785_AT
KYNU	8942	217388_S_AT
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SPINK7	84651	223720_AT
STYX	6815	235180_AT
STYK1	55359	220030_AT
SERPINB3	6317	209719_X_AT
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