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Transcriptomic landscape of lncRNAs in inflammatory bowel disease

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Abstract

Background

Inflammatory bowel disease (IBD) is a complex multi-factorial inflammatory disease with Crohn's disease (CD) and ulcerative colitis (UC) being the two most common forms. A number of transcriptional profiling studies have provided compelling evidence that describe the role of protein-coding genes and microRNAs in modulating the immune responses in IBD.

Methods

In the present study, we performed a genome-wide transcriptome profiling of lncRNAs and protein-coding genes in 96 colon pinch biopsies (inflamed and non-inflamed) extracted from multiple colonic locations from 45 patients (CD = 13, UC = 20, Controls = 12) using expression microarrays platform.

Results

In our study, we identified widespread dysregulation of lncRNAs and protein-coding genes in both inflamed and non-inflamed CD and UC compared to the healthy controls. In case of inflamed CD and UC, we identified 438 and 745 differentially expressed lncRNAs, respectively, while in case of the non-inflamed CD and UC, we identified 12 and 19 differentially expressed lncRNAs, respectively. We also observed significant enrichment (p-value < 0.001, Pearson's Chi-squared test) for 96 differentially expressed lncRNAs and 154 protein-coding genes within the IBD susceptibility loci. Furthermore, we found strong positive expression correlations for the intersecting and *cis*-neighboring differentially expressed lncRNA-protein-coding gene pairs. The functional annotation analysis of differentially expressed genes revealed their involvement in immune response, pro-inflammatory cytokine activity and MHC protein complex.

Conclusions

The lncRNA expression profiling in both inflamed and non-inflamed CD and UC, successfully stratified IBD patients from the healthy controls. Taken together, the identified lncRNA transcriptional signature along with clinically relevant parameters suggests their potential as biomarkers in IBD.

Background

Inflammatory bowel diseases (IBD) are idiopathic chronic relapsing inflammatory conditions of the gastrointestinal (GI) tract. Crohn's disease (CD) and ulcerative colitis (UC) are two most common forms of the IBD. IBD is emerging as a global disease with its incidence and prevalence differentially increasing geographically around the world. Accumulating evidence suggests that IBD result from the complex interplay between genetic, immunologic, and modifiable environmental factors [1], in a genetically susceptible host against a subset of gut commensal microbiota [2-4].

CD is characterized by intestinal inflammation in a discontinuous fashion and involves any part of the GI tract, although in majority of the cases terminal ileum and/or colon is affected. Transmural pattern of inflammation is a hallmark of CD accompanied with other pathophysiological complications like thickened submucosa, intestinal fibrosis, fissuring ulceration in highly active disease, non-caseating granulomas, strictures, abscesses and fistulas [3]. By contrast, UC involves only rectum and colon, and is characterized by superficial inflammation that is restricted to the mucosa and submucosa with presence of cryptitis and crypt abscesses. Disease activity in both CD and UC is typically relapsing and remitting and both conditions are often difficult to diagnose because of idiosyncrasies in the presentation of overlapping and distinct clinical and pathological features [2,3]. Characteristically, diagnosis of either CD or UC is based on number of findings including: clinical symptoms, endoscopic features, radiologic tests, and biopsy histology.

According to a recent meta-analysis of IBD genome-wide association studies (GWAS) data, the number of confirmed genetic loci associated with risk for IBD has increased to 163, with 110 shared between CD and UC, 30 CD-specific and 23 UC-specific. Interestingly, an overwhelming majority of these IBD loci are located in the noncoding intergenic and intronic regions [5]. Most are overlapping the regulatory elements and consequently are likely to influence gene regulation. Findings from our recent studies have demonstrated that large number of annotated long non-coding RNAs (lncRNAs), including novel evolutionarily conserved structured RNA motifs with regulatory potential ([6], unpublished observations (Seemann et al.)), overlap with the IBD loci. Consistent with our findings, another recent study, elegantly revealed that IBD loci overlaps with active regulatory regions in primary intestinal epithelium and immune cells and also IBD loci was found significantly enriched within these active regulatory regions [7].

Several transcriptome profiling studies have provided compelling evidence describing the role of protein-coding and non-coding RNAs (ncRNAs), such as microRNAs, in modulating immune responses in IBD [8-15]. In murine models, loss of endogenous intestinal microRNAs is known to cause impairment of epithelial barrier function that result in acute inflammation [16]. Several studies have explored clinical differences between CD and UC based on transcriptional regulation [17,18]. Recently, Granlund et al. demonstrated lack of major differences between CD and UC based on protein-coding gene expression profiling in IBD [9]. In contrast, expression profiling of colon biopsies from IBD patients allude to differential diagnosis of CD and UC based on transcriptional signatures associated with intestinal inflammation [19].

Long non-coding RNAs (lncRNAs) have emerged as important regulators of gene expression, with accumulating body of evidence linking lncRNAs to a plethora of human pathologies including inflammatory diseases [20]. Although, the precise role of lncRNAs in intestinal diseases remains poorly understood, evidence from recent studies indicates that lncRNAs might be playing a crucial role in inflammatory cascades. Indeed, preponderance of emerging evidence from number of studies, demonstrates important roles of lncRNAs in regulating gene expression within the immune system. Nevertheless, identification of IBD susceptibility loci has afforded limited success in translating results from gene expression studies to advance our knowledge and understanding of the IBD pathophysiology. In particular, the details about the initiation, propagation and maintenance of the lingering inflammation in IBD remains unclear. Furthermore, earlier transcriptomic studies in IBD have mostly been focused on the protein-coding genes with limited profiling studies focusing on microRNAs. However, no study has explored the genome-wide expression profile of lncRNAs in IBD.

In the present study, transcriptomic profiling of lncRNAs and protein-coding genes from colon pinch biopsies of IBD patients was performed using expression microarray platform. Our results identified widespread dysregulation of lncRNAs and protein-coding gene expression in both CD and UC. Notably, differential transcriptomic signature of lncRNAs and protein-coding genes in inflamed-CD (iCD) and inflamed-UC (iUC) enabled clear stratification of the CD and UC phenotype. These data indicate that lncRNAs could potentially be used as predictive biomarkers in IBD.

Methods

Samples collection (Patients and control)

All the patient samples were collected from an IBD cohort at North Zealand Hospital, Hillerød, Denmark. Subjects were required to meet the Copenhagen criteria for Crohn's disease or ulcerative colitis. Participants recruited for the study were patients admitted to the Department of Gastroenterology for colonoscopy who were diagnosed either with CD or UC, or were admitted to the clinic for diagnostic colonoscopy because of symptoms unrelated to the IBD. Written informed consent from all the participants in the study was acquired prior to the collection of samples and medical history. In total, 90 biopsies were collected from 45 individuals (13 CD, 20 UC patients and 12 healthy controls). Subjects were included as normal controls only after all clinical examinations had concluded no signs of autoimmune or inflammatory disease. In case of the IBD groups (CD and UC), one to five endoscopic pinch biopsies were extracted from macroscopically most inflamed mucosa (iCD/iUC) and adjacent non-inflamed (niCD/niUC) mucosa within colon (transverse, descending, sigmoid), ileum, transverse ileum and rectum for the CD, and colon sigmoid and rectum for the UC patients. Whereas, for the control group, one to five biopsies were taken from the same locations as in the CD group, except for the two samples (G3 G1 and 60 G1) that were extracted from duodenal bulb. All biopsies were placed in RNA*later* solution (QIAGEN, Hilden, Germany), and stored for the later downstream use. The study was approved by the Regional Ethical Committee (H-4-2012-030). The inflammation status of biopsies was confirmed by the histologic examination and features of chronic intestinal inflammation for each patient were scored using a previously described scoring system for UC [21] and CD [22]. The pathologists were blinded to the status of inflammation. Additionally, we also tested expression of a panel of twenty-six pro-inflammatory markers (cytokines, interleukins, metalloproteases) using qPCR (Fluidigm platform) to confirm the inflammation status of biopsies (data not shown) prior to the microarray analysis.

RNA extraction and quality control

Total RNA was extracted from biopsies stored in RNA*later* using RNeasy Mini Kit (QIAGEN, Hilden, Germany) according to the manufacturer's instructions. Briefly, the biopsy samples were homogenized in lysis buffer with 1.4 mm ceramic beads (MO BIO Laboratories) using Thermo Savant FastPrep FP120 Homogenizer for 30 seconds at a speed of 4 m/sec. All the remaining steps of the protocol were performed according to the manufacturer's recommendations. To remove traces of genomic DNA, samples were treated with DNase I (QIAGEN, Hilden, Germany). RNA was finally eluted with nuclease-free water supplied with the kit. The quantity and purity of isolated RNA was determined by UV absorbance using NanoDrop 2000 Spectrophotometer (Thermo Scientific, DE, USA), and the integrity of RNA was assessed by analysis of rRNA band integrity on an Agilent 2100

Bioanalyzer RNA 6000 LabChip kit (Agilent Technologies). Only RNA samples with RNA integrity number (RIN) >7 were used for the microarray experiments.

Microarray hybridization

100 ng of total RNA was labeled using LowInputQuick Amp Labeling kit v6.5 (Agilent 5190–2305) following manufacturer instructions. Briefly, mRNA was reverse transcribed in the presence of T7-oligo-dT primer to synthesize cDNA. The cDNA was then in-vitro transcribed with T7 RNA polymerase in the presence of Cy3-CTP to generate labeled cRNA. The labeled cRNA was hybridized to the Agilent Custom 8x60K format lncRNA expression microarray (AMADID 047718, based on Gencode v15 catalog of human long noncoding RNAs, probe length = 60 nt) according to the manufacturer's protocol. Finally, the arrays were washed, and scanned on an Agilent G2565CA microarray scanner at 100% PMT and 3 μ m resolution. Intensity data was extracted using the Feature Extraction software (Agilent). A more detailed and general information about the array can also be found on GENCODE website [23]. The raw microarray data reported in this manuscript have been deposited in the Gene Expression Omnibus (GEO) database with the accession number GSE67106.

Statistical analyses

Raw data was corrected for background noise using normexp method [24]. To assure comparability across samples we used quantile normalization [unpublished observations (Bolstad B. Probe Level Quantile Normalization of High Density Oligonucleotide Array Data. 2001. [25]). Median intensity was taken between technical replicates after checking pairwise Pearson correlation coefficients ($r^2 \ge 0.98$). Differential expression analysis was carried out on non-control probes with an empirical Bayes approach on linear models (LIMMA) [26]. Principal Component Analysis (PCA) method was employed for the initial interpretation of the data. In total, we made seven contrasts to identify differentially expressed genes (iCD vs control, iUC vs control, iCD vs niCD, iUC vs niUC, niCD vs control, niUC vs control and iCD vs iUC) (Additional file 1: Table S1, S2, S3, S4).. P-values were adjusted for multiple comparisons using the False Discovery Rate (FDR) correction [27]. Differentially expressed genes were identified using the double-filtering criterion: adjusted p-value (FDR) < 0.05 and an absolute fold change (abs FC) > 1.5. For transcripts targeted by two probes, only those probes that were changing in the same direction and the probes with highest FC values were retained for further analysis. All statistical analyses were performed with Bioconductor in R statistical environment [28].

Validation of differentially expressed genes by quantitative real-time PCR (qPCR)

The expression of differentially expressed genes from microarray experiments was validated by the quantitative real-time PCR (qPCR) using hydrolysis probe based inventoried and custom designed PrimeTime qPCR 5' Nuclease assays procured from Integrated DNA Technologies (IDT). The Double-Quenched hydrolysis probes with 5' FAM fluorophore, a 3' IBFQ quencher, and an internal ZENTM quencher was used for all the assays.. From the list of top differentially expressed genes from different contrasts, 6 up- and 6 down-regulated genes were selected for their expression validation by qPCR in a subset of samples used for the microarray experiments. In case of protein-coding, 3 up-regulated (DUOXA2, CHI3L1, DST), and 3 down-regulated (PCK1, KCNK10, and SERPINB3) genes were validated. Whereas, for the lncRNAs, 3 up-regulated (*MMP12, RP11-731 F5.2, AC007182.6*), and 3 down-regulated (*DPP10-AS1, CDKN2B-AS1*, and *AL928742.12*) genes were validated. In addition, expression of protein-coding gene *DUOX2* was also measured by qPCR. All cDNAs were prepared using 750 ng of DNA-free RNA using iScriptTM cDNA synthesis kit (BioRad) with a mixture of random and oligo(dT) primers following the manufacturer's instructions. Real-time PCR was performed with 7.5 ng of cDNA per well template for the all the protein-coding genes and lncRNAs with Brilliant III Ultra-Fast QPCR Master Mix (Agilent Technologies). For PCR amplification, the following thermal profile was used: 3 min 95°C; 40 x (5 sec 95°C, 10 sec 60°C). Expression of each lncRNA and protein-coding gene tested was represented as a fold change using the $2^{-\Delta\Delta CT}$ method. *GAPDH* was used as the reference gene.

Identification of IBD-loci associated IncRNAs

All IBD loci marker SNPs and associated genes were retrieved from the ImmunoBase [29]. In total, 233 unique marker SNPs for IBD, CD, and UC regions were retrieved and mapped to the 22,007 lncRNAs (Gencode v15) using intersect feature of the BedTools [30]. The susceptibility locus for IBD was defined based on 500 kb long genomic region with the IBD marker SNP in the middle. The differentially expressed lncRNAs from five contrasts (iCD vs control, iUC vs control, iCD vs niCD, iUC vs niUC, and iCD vs iUC), were mapped to the IBD loci to identify the IBD loci-associated lncRNAs. Regulatory evidence for the IBD-associated SNPs was retrieved from Mokry et al. [7] and RegulomeDB [31].

Functional annotation and Gene Ontology (GO) analysis of differentially expressed lncRNAs

For the differentially expressed lncRNAs, the nearest protein-coding neighbors within a span of <10 kb were identified. For the antisense overlapping or intronic overlapping lncRNAs, intersecting protein-coding gene(s) were identified using intersect feature of BedTools [30]. PANTHER (protein annotation through evolutionary relationship) classification system [32] was used to perform functional annotation and GO analysis of genes that overlap with or are neighbors of the differentially expressed lncRNAs. Likewise, for the IBD loci-associated lncRNAs, GO analysis was performed using the above described nearest neighbor approach. The enrichment for over-represented GO functional terms was calculated based on Binomial test in PANTHER.

Sample classification using SVM based on differentially expressed genes identified by LIMMA

Support Vector Machines (SVM) [33] was used for classifying the CD and UC cases from the controls based on differentially expressed genes identified by LIMMA in five contrasts (iCD vs control, iCD vs niCD, iUC vs control, iUC vs niUC and iCD vs iUC). SVM classification was applied to all five contrasts using the Leave-one-out cross-validation (LOOCV) for differentially expressed lncRNAs and protein-coding genes. To explore the effect of various clinical parameters (age, sex, smoking, disease index and biopsy location) on overall disease outcome, we used the following linear regression function:

```
y = err + w1^*age + w2^*sex + w3^*smoking + w4^*disease index + w5^*biopsy location
```

Here, y = 1 for iCD or iUC disease phenotype and for rest of the samples it is 0. For clinical parameters: age and disease index, we used original values, while for sex and smoking, we used the following binary outcomes: male = 1, female = 0 and smoker =1, non-smoker = 0. In case of six biopsy locations, we used values ranging from 0 to 5. To control any input bias, same analysis was performed on a randomized lncRNA gene list of the same number as the total differentially expressed lncRNA genes. The feature values were normalized to values ranging from 0 to 1 using (x-min)/(max-min). Linear regression was applied using Scikit-learn [34] package in python and least square method was used for optimization in our analysis. Furthermore, differentially expressed genes identified by LIMMA were verified by Support Vector Machines- Recursive Feature Elimination (SVM-RFE) method [35]. SVM-RFE recursively prunes genes whose absolute weights are the smallest until desired number of features is reached. For each contrast, we used SVM-RFE to identify the same number of differentially expressed genes as identified by LIMMA.

Co-expression network analysis

To identify CD and UC specific networks clusters (modules) based on highly correlated genes, weighted correlation network analysis (WGCNA) method was used [36]. We used the normalized expression data as input and removed the outlier samples. The clinical parameters were represented as following: numeric for age and disease, binary for sex, ethnicity (3 categories), smoking (4 categories), clinical subgroup (5 categories), and biopsy location (6 categories). The standard procedure of WGCNA was applied for network construction and module identification. The trait-based gene significance measure is defined as the absolute correlation and correlation test p-value between the trait and the gene expression profile. Gene Ontology analysis of modules was performed with GOstats package in R [37] using adjusted p-value < 0.001. We controlled for study bias in the GO analysis by running the same analysis for randomized gene sets of the same module sizes.

Results

The overall summary describing sample information is provided in Table 1. Both, CD and UC samples were divided based on inflammation status confirmed by both macroscopic, microscopic evaluations and pro-inflammatory gene signatures into inflamed (iCD, iUC) and non-inflamed (niCD, niUC) categories. The total number of samples included 21 iCD, 23 niCD, 15 iUC, 9 niUC and 22 healthy controls. In total, 90 intestinal pinch biopsies (45 individuals (13 CD, 20 UC patients and 12 healthy controls)) from multiple colonic regions were harvested from both inflamed and non-inflamed mucosa (Figure 1A). Detailed sample information including the ethnic background, disease index, previous treatment regimens and other clinical parameters are listed in Table 2.

Diagnosis	Number of samples	Number of individuals	
iCD	21	13	
niCD	23		
iUC	21 (15 unique)	20	
niUC	9		
Controls	22	12	
Total	96 (90 unique samples)	45	

Table 1 Overall study design and sample information

90 biopsy samples extracted from different colonic locations from 45 patients (CD = 13, UC = 20, Controls = 12). 6 samples from UC patients were used as technical replicates.

Figure 1 Study design and inflammatory gene signature for iCD and iUC. (A) Study design included 90 pinch biopsies from multiple colonic regions for both inflamed and non-inflamed mucosa (21 iCD, 23 niCD, 15 iUC, 9 niUC and 22 healthy controls samples). (B) Principal component analysis (PCA) separated inflamed-CD (iCD) and inflamed-UC (iUC) samples from non-inflamed and healthy controls. PC1 and PC2 together explained 15% of the total variation. (C) Unsupervised hierarchical clustering of the most dynamic probes (coefficient of variance >0.05) across the samples resulted in clustering of samples according to their clinical subgroups. (D) The log2 ratio and -log10 adj.P-values are plotted in the form of volcano plots for iCD vs control, iUC vs control and iCD vs iUC contrasts. The probes in red, blue and orange colors represents up-regulated (FC >1.5 and adj.P-value <0.05), downregulated (FC < -1.5 and adj.P-value < 0.05) and significant with small fold change (FC >-1.5 and <1.5), respectively. The non-significant probes are represented in black color. The selected protein-coding genes and lncRNAs labeled in black and green, respectively. (E) Venn diagram shows the overlap between differentially expressed genes identified in iCD vs control, iUC vs control and iCD vs iUC contrasts. The up-regulated genes are depicted in italics, down-regulated as underlined and contra-regulated in red.

Number of individuals	CD	UC	Controls
	13	20	12
Age	31 (19–59)	46 (18-68)	54 (18–77)
median years (range)			
Average age at diagnosis (years)	27	33	NA
Average years with disease (Disease duration)	8	9.3	NA
Female / Male	6/7	13/7	8/4
Smoking			
Smoker (S)	8	1	4
Previous (P)	4	8	1
Never (N)	1	11	5
Not disclosed (ND)	-	-	2
Ethnicity			
Danish (DK)	9	19	12
European (EU)	1	-	-
Middle Eastern (ME)	3	1	-
Number of individuals with family history of other autoimmune diseases	1 (7%)	6 (30%)	5 (41%)
Number of patients on medication			NA
5-ASA	2	13	
Solumedrol	2	1	
Azathioprin	2	2	
Budesonide	1	1	
Prednisolon	1	2	
Disease Index	HB index $= 3-36$	SCCAI index = $2-12$	

Table 2 Clinical parameters

Each column summarizes characteristics for all patients contributing with samples to the corresponding sample groups. 5-ASA – 5-aminosalicylic acid. HB index – Harvey Bradshaw index. SCCAI index – Simple Clinical Colitis Activity index.

Microarray analysis of lncRNAs and protein-coding gene expression

In Gencode v15 lncRNA microarray design, each lncRNA transcript is targeted by two probes covering 22,001 lncRNA transcripts corresponding to 12,963 lncRNA genes. In addition, each array contains 17,535 randomly-selected protein-coding targets, of which 15,182 (unique 12,787) correspond to protein-coding genes. Six samples analyzed in duplicates, hybridized on separate chips, and used as technical replicates showed strong positive Pearson correlation ($r^2 \ge 0.98$, p-value < 2.2e-16) (Additional file 2: Figure S1). Based on the principal component analysis (PCA) (see methods section for details), separation of iCD and iUC samples from niCD, niUC and healthy controls were observed (Figure 1B). However, there was no apparent separation between iCD and iUC samples. The scatterplot matrices describing the first four principal components are described in Additional file 2: Figure S2. Unsupervised hierarchical clustering of the most dynamic probes (coefficient of variance >0.05) across the samples ensued in clustering of samples according to their clinical subgroups (Figure 1C). The probes targeting lncRNAs and protein-coding genes separately also clustered samples in a similar manner (Additional file 2: Figure S3A and S3B).

Differential transcriptional signature of lncRNAs and protein-coding genes in CD and UC

To define CD and UC specific transcriptional signatures based on intestinal inflammation, we identified differentially expressed genes using LIMMA [24] (based on a cutoff of log2 fold change (FC) >1.5 (up-regulated), FC < -1.5 (down-regulated) and adj.P-value <0.05 (moderated t-test)) in all contrasts (Table 3). The log2 ratio and -log10 adj.P-values are plotted and represented as volcano plots for iCD vs control, iUC vs control and iCD vs iUC contrasts in Figure 1D. For the non-inflamed tissues contrasts (iCD vs niCD and iUC vs niUC), the volcano plots are shown in Additional file 2: Figure S4A and S4B, respectively.

ed genes			
iCD	niCD	iUC	niUC
1477	73	2429	44
	435	73	
			1814
iCD	niCD	iUC	niUC
1039	61	1684	25
	328	50	
			1215
iCD	niCD	iUC	niUC
438	12	745	19
	107	23	
			599
	iCD 1477 iCD 1039 iCD	iCD niCD 1477 73 435 iCD niCD 1039 61 328 iCD niCD 438 12	iCD niCD iUC 1477 73 2429 435 73 iCD niCD iUC 1039 61 1684 328 50 iCD niCD iUC 438 12 745

	Table 3 Total	number of	differentially	expressed genes
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Total differentially expressed genes identified in seven pairwise contrasts (iCD vs control, iUC vs control, iCD vs niCD, iUC vs niUC, niCD vs control, niUC vs control and iCD vs iUC).

Differential gene expression analysis identified the following up/down-regulated genes: 761/278 protein-coding genes and 254/184 lncRNAs in iCD vs control and 1085/599 proteincoding genes and 370/375 lncRNAs in iUC vs control (Table 3 and Figure 1E). The top upregulated and down-regulated lncRNAs and protein-coding genes (based on FC) for iCD vs control and iUC vs control are listed in Tables 4 and 5. Interestingly, lncRNA *RP11-731 F5.2* (whose 3' end partly spans into the start of the *IGHG2* gene) and antisense lncRNA *MMP12* were found significantly up-regulated, whereas, the antisense *DPP10-AS1*, *ANRIL* (*CDKN2B-AS1*) and *DIO3OS* lncRNAs were significantly down-regulated in both iCD vs control and iUC vs control contrasts (Tables 4 and 5).

Up-regulated IncRNAs	Up-regulated protein-coding genes				
Gene Name	Transcript	FC	Gene Name	Transcript	FC
RP11-731 F5.2	ENST00000460164.1	14.14	REG3A	NM_138938	52.71
MMP12	ENST00000532855.1	6.64	DUOXA2	NM_207581	47.26
MMP12	ENST00000326227.5	6.52	DEFA5	NM_021010	37.73
RP11-465 L10.10	ENST00000419897.1	5.69	DEFA6	NM_001926	28.33
RP11-44 K6.2	ENST00000520185.1	3.83	CHI3L1	NM_001276	26.29
FAM66D	ENST00000526690.1	3.36	CXCL1	NM_001511	14.8
LINC01272	ENST00000445003.1	3.35	DMBT1	NM_007329	13.45
RP11-44 K6.4	ENST00000522970.1	3.24	SAA1	NM_000331	12.67
SAA2-SAA4	ENST00000524555.1	3.16	CXCL9	NM_002416	12.07
KIF9-AS1	ENST00000429315.2	3.14	IGHG3	ENST00000390551	11.52
Down-regulated IncRNAs			Down-regulated protein-coding genes		
DPP10-AS1	ENST00000432658.1	-8.57	PCK1	NM_002591	-5.55
PDZK1P2	ENST00000401008.2	-4.11	SLC26A2	NM_000112	-3.82
DIO3OS	ENST00000553575.1	-3.01	C10orf116	NM_006829	-3.8
DIO3OS	ENST00000554694.1	-3.01	GUCA2B	NM_007102	-3.61
DIO3OS	ENST00000557532.1	-2.99	LCN15	NM_203347	-3.43
DIO3OS	ENST00000557109.1	-2.98	AQP7P1	NR_002817	-3.32
ANRIL (CDKN2B-AS1)	ENST00000422420.1	-2.97	TRPM6	NM_017662	-3.19
ANRIL (CDKN2B-AS1)	ENST00000428597.1	-2.97	TNNC2	NM_003279	-3.1
DIO3OS	ENST00000554441.1	-2.96	UGT2A3	NM_024743	-2.97
DIO3OS	ENST00000554735.1	-2.95	ADH1C	NM_000669	-2.96

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Table 4 Top ten	differentially e	expressed	genes in i(1)
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Top ten up and down-regulated lncRNAs and protein-coding genes in iCD vs control contrast. The log2 fold change is denoted as FC.

Table 5 Top ten differentially expressed genes in iUC

Up-regulated IncRNAs	Up-regulated	protein-coding genes			
Gene Name	Transcript	FC	Gene Name	Transcript	FC
RP11-731 F5.2	ENST00000460164.1	20.64	DUOXA2	NM_207581	109.61
MMP12	ENST00000532855.1	17.05	CHI3L1	NM_001276	39.71
MMP12	ENST00000326227.5	16.54	SAA1	NM_000331	30.67
RP11-465 L10.10	ENST00000419897.1	9.52	CXCL1	NM_001511	25.92
KIF9-AS1	ENST00000429315.2	5.75	MMP7	NM_002423	21.2
FAM66D	ENST00000526690.1	5.73	SLC6A14	NM_007231	20.52
SAA2-SAA4	ENST00000524555.1	5.66	IGHG3	ENST00000390551	20.14
CLRN1-AS1	ENST00000476886.1	4.64	MMP12	NM_002426	15.76
RP11-1149O23.3	ENST00000517774.1	4.29	C4orf7	NM_152997	14.76
RP5-1028 K7.2	ENST00000578280.1	4.21	CXCL2	NM_002089	11.91
Down-regulated lncRNAs			Down-regulate	ed protein-coding genes	
ANRIL (CDKN2B-AS1)	ENST00000422420.1	-8.67	PCK1	NM_002591	-15.24
ANRIL (CDKN2B-AS1)	ENST00000428597.1	-8.31	OSTalpha	NM_152672	-11.33
ANRIL (CDKN2B-AS1)	ENST00000585267.1	-7.06	ANPEP	NM_001150	-11.02
ANRIL (CDKN2B-AS1)	ENST00000580576.1	-6.92	SLC26A2	NM_000112	-10.46
ANRIL (CDKN2B-AS1)	ENST00000577551.1	-6.74	GBA3	NM_020973	-9.28

ANRIL (CDKN2B-AS1)	ENST00000581051.1	-6.72 GUCA2A	NM_033553	-9.22
ANRIL (CDKN2B-AS1)	ENST00000582072.1	-6.68 SLC3A1	NM_000341	-9.21
PDZK1P2	ENST00000401008.2	-6.67 GUCA2B	NM_007102	-8.84
DPP10-AS1	ENST00000432658.1	-5.95 TMIGD1	NM_206832	-8.17
ANRIL (CDKN2B-AS1)	ENST00000421632.1	-5.78 SLC1A7	NM_006671	-6.57
TE (1.1 1	(1.1 D)14 1 ('	1		0.611.1

Top ten up and down-regulated lncRNAs and protein-coding genes in iUC vs control contrast. The log2 fold change is denoted as FC.

In case of the protein-coding genes, the top differentially expressed genes included, DUOXA2, CHI3L1, CXCL1 and SAA1, which were all significantly up-regulated, whereas, PCK1, SLC26A2, GUCA2B, were significantly down-regulated (Tables 4 and 5). In case of iCD vs controls, REG3A was >52 fold up-regulated (adj.P-value = 2.17e-04). The top differentially expressed lncRNAs and protein-coding genes for iCD vs niCD and iUC vs niUC contrasts, displayed similar expression patters as healthy controls (Additional file 1: Table S1 and S2).

On comparing niCD vs control and niUC vs control, only a small number of up/downregulated genes (61/25 and 8/17 protein-coding, 12/19 and 9/10 lncRNAs) were identified for niCD and niUC, respectively. Nearly, all of the differentially expressed genes in niCD vs control were also present in iCD vs control contrast with the exception of protein-coding gene *CRYBB2* (FC = -1.5) (Additional file 1: Table S3). Conversely, in case of niUC vs control, majority (15 out of 17) of the up-regulated genes including 4 small nucleolar RNAs (snoRNAs: *SNORD97, SNORA28, SNORA53,* and *SNORA74A*) and the down-regulated genes, *MAST3, CPT1B, LOC338799, EXOC3L4,* and *MAPK8IP3* were specifically found in niUC only (Additional file 1: Table S4). Importantly, in the case of iCD vs iUC contrast, 18/32 protein-coding genes and 13/10 lncRNAs were significantly found to be up/downregulated. The top up/down-regulated lncRNAs and protein-coding genes for iCD vs iUC are shown in Table 6. The annotations for the Gencode v15 [38] microarray features for lncRNAs are summarized in Figure 2A. Majority of the differentially expressed lncRNAs identified in our analysis belonged to three main classes: antisense, processed transcripts and intergenic lincRNAs (Figure 2B), as described in the following section.

Up-regulated IncRN	As	Up-regulated p	protein-coding genes		
Gene Name	Transcript	FC	Gene Name	Transcript	FC
FLJ42969	ENST00000514926.1	2.6	C8G	NM_000606	2.75
AC007182.6	ENST00000455232.1	2.42	SLC25A34	NM_207348	2.43
RP11-542 M13.2	ENST00000599411.1	2.04	UGT1A6	NM_001072	2.25
RP11-399 F4.4	ENST00000453998.1	1.87	LRRC66	NM_001024611	2.20
FAM95B1	ENST00000455995.1	1.87	EXOC3L4	NM_001077594	1.95
RP3-395 M20.8	ENST00000432521.2	1.69	ANO7	NM_001001891	1.95
RP3-395 M20.8	ENST00000448624.2	1.65	GLYCTK	NM_145262	1.90
OPLAH	ENST00000426825.1	1.61	CLEC10A	NM_182906	1.89
OPLAH	ENST00000534424.1	1.61	FAM95B1	NR_026759	1.89
SPPL2B	ENST00000592738.1	1.59	LPIN3	NM_022896	1.87
Down-regulated Inc	RNAs		Down-regulated protein-coding genes		
AL928742.12	ENST00000412518.1	-2.01	SERPINB3	NM_006919	-3.87
RP11-444D3.1	ENST00000540811.1	-1.84	SLC6A14	NM_007231	-3.51
AL928742.12	ENST00000427543.1	-1.8	GAL	NM_015973	-2.50
FAM25D	ENST00000426412.2	-1.69	GJB4	NM_153212	-2.38
RP11-274 N19.2	ENST00000515643.1	-1.64	IGHV1-58	ENST00000390628	-2.29
RP11-838 N2.4	ENST00000579007.1	-1.59	CRYM	NM_001888	-2.28
RP11-279 F6.3	ENST00000558941.1	-1.57	SLC26A4	NM_000441	-2.22
RP11-279 F6.3	ENST00000559212.1	-1.55	DEFB103B	NM_018661	-2.22

Table 6 Top ten differentially expressed genes in iCD vs iUC

LINC00524	ENST00000555860.1	-1.54	LAMC2	NM_005562	-2.20	
VAV3-AS1	ENST00000438318.1	-1.52	TUSC3	NM_178234	-2.02	
Top ten up and down-regulated lncRNAs and protein-coding genes in iCD vs iUC. The log2 fold change is denoted as FC.						

Figure 2 Gencode v15 annotation of the total differentially expressed lncRNAs in IBD and microarray validation by qPCR. (A) The Gencode v15 array targeted 22007 lncRNA transcripts falling into seven major annotation classes (antisense, processed transcripts, intergenic (lincRNAs), sense overlapping, sense intronic and retained introns). Three classes (ambiguous orf, non-coding RNAs and TEC (to be experimentally confirmed) with small number of lncRNAs were merged into miscellaneous (misc) class for better representation. (B) Three major classes of differentially expressed lncRNAs identified in our study: intergenic (lincRNAs), processed transcripts and antisense lncRNAs. (C) The differences between the expression levels of top 3 most up-and down regulated protein-coding genes (in blue) and lncRNA genes (in red) in different the clinical were tested using Kruskal-Wallis with Dunn's multiple comparison test. The top 3 up-regulated protein-coding genes (DUOXA2, CHI3L1 and CXCL9) and lncRNAs genes (MMP12, FAM66D and SAA2-SAA4), showed increasing signal intensity from control group to inflamed CD and UC groups based on averaged gene expression levels (p-value < 0.001). While, in case of the top 3 downregulated protein-coding genes (PCK1, GUCA2B and TNNC2) and lncRNA genes (DPP10-ASI, PDZK1P2 and ANRIL), a decreasing signal intensity across the clinical subgroups was observed from iCD, iUC to controls (p-value < 0.001). (D) A total of 8 genes were selected for real-time PCR validation of the microarray data in iCD vs control (red) and iUC vs control (blue). The log2 fold change (FC) is plotted on the y axis.

Additionally, we also tested the differences between the clinical subgroups for the top differentially expressed protein-coding and lncRNA genes. The top 3 up-regulated protein-coding genes (*DUOXA2, CHI3L1* and *CXCL9*) and lncRNAs genes (*MMP12, FAM66D* and *SAA2-SAA4*), showed increasing signal intensity based on the averaged gene expression levels across the spectrum of clinical subgroups from control to iCD and iUC (p-value < 0.001) (Figure 2C). In case of top 3 down-regulated protein-coding genes (*PCK1, GUCA2B* and *TNNC2*) and lncRNA genes (*DPP10-AS1, PDZK1P2* and *ANRIL*), we observed decreasing signal intensity across the clinical subgroups from iCD, iUC to controls (p-value < 0.001) (Figure 2C). Importantly, eight major isoforms (out of total 17 annotated isoforms) of *ANRIL* were found to be down-regulated in iCD and iUC when compared to controls and non-inflamed tissues in our data (Tables 4 and 5). *ANRIL* was -2.97 and -2.72 fold down-regulated in iCD vs control and iCC vs niCD contrasts, respectively.

Furthermore, for the validation of microarray results by quantitative real-time PCR (qPCR), we selected 8 top differentially expressed genes (based on FC) common between iCD vs control and iUC vs control (up-regulated: *DUOXA2, CHI3L1, DUOX2, MMP12, RP11-731 F5.2*; down-regulated: *PCK1, DPP10-AS1, ANRIL*). The qPCR analysis confirmed the microarray expression results with respect to the fold change values (Additional file 1: Table S5). We also performed qPCR analysis for *DUOX2* although it was not probed on our microarray, but it has been implicated along with its maturation factor *DUOXA2* in IBD pathogenesis (see discussion). Both *DUOXA2* and *DUOX2* were found to be significantly up-regulated in iCD vs control (FC = 8.83 and 5.85) and iUC vs control (FC = 9.14 and 6.05) (Figure 2D). In case of the remaining four contrasts, we also tested 5 differentially expressed genes by qPCR validation (Additional file 1: Table S6 and S7).

Overlap of differentially expressed genes in iCD and iUC

A Venn diagram illustrating the relationship between lncRNAs and protein-coding genes differentially expressed in iCD and iUC is shown in Figure 3. In total, 337 differentially expressed lncRNAs were identified as common between iCD and iUC with 100 unique lncRNAs for iCD and 400 unique lncRNAs for iUC (when compared to the healthy controls) (Figure 3A). While in case of the protein-coding genes, 901 differentially expressed genes were found to be common for iCD and iUC with 128 unique for iCD and 739 unique for iUC (Figure 3B). Conversely, in iCD vs iUC contrast, 19 out of 23 and 45 out of 50 differentially expressed lncRNAs and protein-coding genes, respectively overlapped with iCD vs control and iUC vs control.

Figure 3 Overlap of differentially expressed lncRNAs and protein-coding genes between iCD and iUC. Venn diagram shows an overlap of 337 lncRNAs (**A**) and 901 protein-coding genes (**B**) that were differentially expressed (fold change >1.5, adj.P-value <0.05) between patients with iCD and iUC when compared with healthy controls. We observed contraregulated genes between iCD/iUC vs control contrasts when compared with iCD vs iUC contrast. The up-regulated genes are depicted in italics, down-regulated as underlined and contra-regulated in red. Heat maps of average normalized gene expression for the overlapping 337 lncRNAs (**C**) and 901 protein-coding genes (**D**) between iCD and iUC in the five clinical subgroups (iCD, niCD, iUC, niUC and controls) are displayed. Selected up-regulated and down-regulated genes are listed.

The unsupervised hierarchical clustering showed that both inflamed groups (iCD and iUC) cluster together in contrast to the non-inflamed (niCD and niUC) which clustered with healthy controls. The normalized gene expression values from the above mentioned 337 lncRNAs and 901 protein-coding genes common to both iCD and iUC conditions were averaged for each of the five clinical subgroups and visualized in a heat map in Figure 3C and D. The expression patterns for the specific up-regulated and down-regulated genes showed increasing or decreasing signal intensity across the clinical subgroups (from iCD, iUC, niCD, niUC and healthy controls). Collectively, these overlapping differentially expressed genes between iCD vs control and iUC vs control define a distinct inflammatory iCD/iUC gene expression signature. Importantly, this inflammatory gene signature included the key drivers of the innate and adaptive immune responses (for example *DUOXA2* and *CXCL1*).

Comparison of expression levels of top differentially expressed genes in patients and healthy controls

To stratify iCD and iUC samples from the healthy controls, we also compared the expression profiles of top 20 up/down-regulated lncRNAs and top 20 up/down-regulated protein-coding genes (based on FC) through unsupervised hierarchical clustering. The expression map of these top 40 differentially expressed genes displayed a clear separation of the patients from the control groups (Figure 4A and B), except for the two iUC samples B11 and 17_3 which were misclassified in the clustering. The magnitude of log2 intensity signal for these top differentially expressed genes displayed in Figure 4 was >6 in both iCD and iUC. Interestingly, in case of the iCD vs iUC contrast, clustering was unable to distinguish between iCD and iUC patients (Additional file 2: Figure S5). In addition to the top candidates, we also compared the expression profiles of all differentially expressed lncRNAs

and protein-coding genes, and observed similar results as described above (Additional file 2: Figure S6A and S6B).

Figure 4 Comparison of expression levels of the top 40 differentially expressed lncRNAs and protein-coding genes. Unsupervised hierarchical clustering of samples (patients in red, controls in blue) based on normalized expression values from the top 40 up and down-regulated lncRNAs and protein-coding genes for (A) iCD vs control and (B) iUC vs control. The log2 normalized expression values are shown in the color key. A clear separation between the diseased from control group is visible in case of iCD vs control contrast.

Inflammatory response and antimicrobial peptide (AMPs) genes are dysregulated in iCD and iUC

Antimicrobial peptides (AMPs) play an important role in protection of the host intestinal mucosa against microorganisms and AMPs dysregulation have been associated with IBD pathogenesis (see discussion for details). Therefore, we investigated whether there were differences in expression of genes involved in inflammatory response and AMPs, between different clinical subgroups. Our analysis identified key genes associated with inflammatory response, including the pro-inflammatory chemokines and cytokines. *CCL11*, *CCL19*, *CCL4* and *CXCL9*, were significantly up-regulated, in both iCD vs control and iUC vs control. In addition, we also found key antimicrobial response genes to be significantly up-regulated in iCD and iUC when compared to healthy controls (Figure 5). *REG3A*, *DEFA5* and *DEFA6* were >30 fold up-regulated only in iCD vs control. Chemokines *CXCL1* and *CXCL2* were >15 and >25 fold up-regulated in both iCD vs control and iUC vs control, respectively. *CXCL5*, *IL15* and *C3AR1* were specifically up-regulated in iCD vs control (Figure 5). Notably, *NOD2* gene was >2 fold up-regulated in iCD vs control, iUC vs control and iCD vs niCD contrasts. *DEFB1* and *NPY* were the only AMP genes that were significantly down-regulated in both iUC and iCD.

Figure 5 Differentially expressed protein-coding genes involved in antimicrobial and autoimmune response. Key genes involved autoimmune and inflammatory immune responses and AMPs were found to be dysregulated in both iCD and iUC when compared to healthy controls as well as to non-inflamed tissues. The log2 fold change values are plotted on the y axis.

Differentially expressed genes in iCD and iUC are enriched within IBD loci

Since majority of disease associated susceptibility SNPs map to the non-coding regions in the genome, we looked for the presence of known IBD associated SNPs (total 233 SNPs) within the Gencode v15 annotated lncRNAs. Interestingly, 29 IBD risk variants intersected 37 lncRNAs, of which only *IFNG-AS1* antisense lincRNA (*ENST00000536914.1*) harboring UC susceptibility SNP rs7134599 was found to be differentially expressed in our study. *IFNG-AS1* was up-regulated in iUC vs control (FC = 1.54) and iUC vs niUC (FC = 1.52). Furthermore, we identified IBD loci associated lncRNAs and protein-coding genes by intersecting the IBD susceptibility loci, which was defined as 500 kb long genomic region with the IBD risk variant in the middle. In total, 1040 IBD loci-associated lncRNAs were identified, out of which 96 lncRNAs were found to be differentially expressed (Additional file 1: Table S8). These differentially expressed lncRNAs co-localized with 57 IBD risk variants (within 500 kb locus), and were found to be enriched within IBD loci (p-value <

0.0001, Pearson's Chi-squared test). In case of protein-coding genes, 681 genes were found to be associated with IBD loci, out of which 154 were differentially expressed and enriched within IBD loci (p-value < 0.0001, Pearson's Chi-squared test). Based on unsupervised hierarchical clustering of averaged normalized gene expression values of 96 and 154 differentially expressed IBD loci-enriched lncRNAs and protein-coding genes, respectively, enabled independent stratification of disease from the controls and further distinguished inflamed from the non-inflamed conditions in both CD and UC (Figure 6).

Figure 6 Averaged gene expression for differentially expressed IBD loci-associated lncRNAs and protein-coding genes. Unsupervised hierarchical clustering of averaged normalized expression values for (A) 96 differentially expressed IBD loci-associated lncRNAs and (B) 154 protein-coding genes in different clinical subgroups. The range for the expression values is shown in the color scale.

Regulatory IBD-associated SNPs co-localize with differentially expressed IBD loci-associated lncRNAs

Next, we asked whether active regulatory regions within the IBD loci overlap with the differentially expressed IBD loci-associated lncRNAs. IBD associated SNPs overlapping active regulatory elements in intestinal epithelium were retrieved from Mokry et al. study [7]. In their study, the active regions overlapping IBD associated SNPs were identified based on H3K27ac chromatin immunoprecipitation and sequencing (ChIP-Seq). Out of 96 differentially expressed IBD loci-associated lncRNAs, 68 lncRNAs were found to be associated with 24 IBD loci SNPs co-localizing with the active regulatory elements in intestinal epithelium and immune cells (Additional file 1: Table S8). These overlapping IBD loci-associated active regulatory elements have been reported to frequently co-localize with the known transcription factor binding motifs [7]. A number of IBD-associated SNPs potentially affect the binding affinity of transcriptional factors, and thus perturb the gene expression. Additionally, IBD-associated risk variants also act as expression quantitative trait loci (eOTLs) signals for number of genes (Additional file 1: Table S8). For example, IBDassociated risk variant rs10797432 located within IBD loci-associated lncRNA RP3-395 M20.8 (ENSG00000238164) alters the binding motifs for TFAP2A and CTCF. Furthermore, it is also known to acts as cis-eQTL for MMEL1 in monocytes. Regulatory IBD-associated SNP rs1569723 located within IBD loci-associated lncRNA RP11-465 L10.10 (ENSG00000204044) acts as cis-eQTL for CD40 in monocytes. Also, SNP rs12946510 associated with lncRNAs RP11-387H17.4 (ENSG00000264968) and RP11-94 L15.2 (ENSG00000264198) is known to perturb the binding sites for transcription factors FOXO1, ELF3, and SRF. In addition, this SNP also acts as a cis-eQTL for pseudogene KRT222P, transcriptional co-activator complex component MED24, transcription factor NR1D1, and ORMDL3 in lymphoblastoid cell lines. In case of the antisense lncRNA CTD-2196E14.5 (ENSG00000261266), the associated SNP rs7404095 acts as a cis-eQTL for PRKCB in lymphoblastoid and *PRKCB1* in monocytes. Moreover, SNP rs734999 associated with IncRNA RP3-395 M20.8 (ENSG00000238164) acts as a cis-eQTL for TNFRSF14 in lymphoblastoid cell line.

Cis-acting correlation of expression between differentially expressed IBD lociassociated lncRNAs and protein-coding genes

We computed pairwise Pearson correlations in order to explore the possible co-expression patterns between IBD loci-associated differentially expressed lncRNAs and protein-coding genes. Pairwise correlations of expression involving neighboring lncRNAs and proteincoding genes associated with each IBD-associated SNP (500 kb loci with SNP in the middle) were computed. We found positive ($r^2 \ge 0.5$) and extreme positive ($r^2 \ge 0.9$) correlations between the overlapping as well as *cis*-neighboring differentially expressed IBD lociassociated lncRNA-protein-coding gene pairs (p-value < 0.05). The pairwise correlations for six intersecting IBD loci associated lncRNA-protein-coding gene pairs LSP1 and ENST00000509204.1 (rs907611), HLA-DQB1 and ENST00000443574.1 (rs9268853, rs6927022), MST1 and ENST00000563780.1 (rs9822268 and rs3197999), TSPAN33 and ENST00000498745.1 (rs4728142), SLC22A5 and ENST00000417795.1 (rs2188962, rs12521868), DGRD and ENST00000442524.1 (rs12994997, rs3792109) are plotted in Figure 7. Interestingly, lncRNA ENST00000563780.1 and MST1 protein-coding gene exhibited extreme positive correlation ($r^2 \ge 0.99$) (Figure 7). Enrichment for positive correlations has been reported for the lncRNAs intersecting protein-coding genes in antisense orientation [35]. Indeed, we also observed strong positive correlation ($r^2 \ge 0.7$) for the intersecting antisense IncRNA ENST00000417795 and SLC22A5 protein-coding gene.

Figure 7 Correlations of expression for cis-neighboring pairs of IBD loci-associated differentially expressed lncRNAs and protein-coding genes. Expression correlations for *cis*neighboring pairs of IBD loci-associated lncRNAs and protein-coding genes. Overall positive correlations between overlapping protein-coding and lncRNA gene-pairs (a) LSP1 and and ENST00000443574.1 ENST00000509204.1 **(b)** *HLA-DOB1* (c) *MST1* and *ENST00000563780.1* (d) *TSPAN33* and *ENST00000498745.1* (e) *SLC22A5* and ENST00000417795.1 (f) DGRD and ENST00000442524.1 were observed. An extreme positive ($r^2 > 0.99$, p-value < 2.2e-16) correlation was observed in case of MST1 and its intersecting lncRNA ENST00000563780.1 associated with IBD risk variants rs9822268 and rs3197999. Protein-coding expression is plotted on the x-axis, and lncRNA expression is shown on the y-axis. Each point represents a biopsy sample from different clinical subgroups.

Functional annotation of differentially expressed lncRNAs

The functional annotations of lncRNAs have mostly been based on nearest-neighbor approach i.e. "guilt-by-association" analyses, for example Cabili et al. [39]. We therefore analyzed the GO terms of genes that overlap with or are neighbors of the differentially expressed lncRNAs. We identified 516 nearest protein-coding neighbors within a span of <10 kb covering 610 differentially expressed lncRNAs. In addition, we also included 712 neighboring protein-coding genes for the 57 IBD risk variants (associated with 96 differentially expressed IBD loci-associated lncRNAs) based on 1 Mb locus size for each variant. The most significant over-represented GO terms in the biological process category included, antigen processing and presentation (p-value 7.39e-08), immune system process (p-value 2.5e-05) and natural killer cell activation (p-value 9.6e-05) (Additional file 1: Table S9). In the cellular component category, we found enrichment for MHC protein complex (p-value 5.95e-09). Furthermore, we also observed enrichment for over-represented GO terms in molecular function category which included protein binding, receptor binding and cytokine activity.

Cross validation of differentially expressed genes by SVM

Support Vector Machines (SVM) [33] was used for classifying IBD cases from controls and for cross-validating differentially expressed genes identified by LIMMA. The best SVM classifier performance was obtained from differentially expressed lncRNAs identified in iCD vs control followed by iUC vs control contrast (see methods for details). The classifier distinguished iCD and iUC from controls with 100% and 94.6% accuracy, 100% and 100% specificity and 100% and 86.7% sensitivity, respectively. In addition, the classifier was also able to distinguish iCD and iUC from niCD and niUC samples with an accuracy of 86.4% and 91.7%, with 78.3% and 88.9% specificity and 95.2% and 83.3% sensitivity, respectively. While for iCD vs iUC contrast, the accuracy of classifier was 77.8%, with 60.0% specificity and 90.4% sensitivity (Figure 8A). For the differentially expressed protein-coding genes, the classifier achieved accuracy of 100% and 94.6%, with 100% and 100% specificity, with %100 and 86.7% sensitivity, in discriminating iCD and iUC from controls, respectively (Figure 8B). Similar to the above described observations, the classifier also allowed distinction between iCD and iUC from niCD and niUC samples with an accuracy of 81.8% and 83.3%, with 78.2% and 77.8% specificity and 85.7% and 86.7% sensitivity, respectively. While for iCD vs iUC contrast, the accuracy of classifier was 88.9%, with 80.0% specificity and 95.2% sensitivity (Figure 8B). Furthermore, our classifier achieved the similar performance when using combined differentially expressed protein-coding and lncRNA genes or only protein-coding genes (data not shown). The effect of clinical parameters (Table 2 and Additional file 2: Figure S7) on disease (iCD or iUC) was described by the following linear function:

Figure 8 ROC curve analysis of differentially expressed lncRNAs and protein-coding genes. Receiver operating characteristic (ROC) curve analysis of differentially expressed (A) lncRNAs and (B) protein-coding genes for five contrasts.

y = 0.511 + (-0.212*age) + (-0.114*sex) + (-0.339*smoking) + (1.185*disease score) + (-0.058*biopsy location)

Using t-statistics, p-values for linear regression coefficients for age, sex, smoking, disease index and biopsy location were 1.40e-01, 2.29e-01, 2.98e-03, 5.02e-07 and 6.43e-01, respectively. Our analysis indicated that disease index had strongest effect on defining iCD and iUC phenotype, followed by smoking, sex and age (Additional file 2: Figure S7, Figure S8). Differentially expressed genes identified by LIMMA were verified by Support Vector Machines- Recursive Feature Elimination (SVM-RFE) [35], which revealed a robust concordance rate in terms of total number of overlapping differentially expressed genes identified by the two methods (Additional file 2: Figure S9). In both iCD vs control, and iUC vs control contrasts, the overlap of about 66% was observed. To control for any input bias, a randomized lncRNA gene list of same size as differentially expressed lncRNAs was also used in this analysis (Additional file 2: Figure S8).

Impact of clinical parameters on disease diagnosis

We next investigated impact of the clinical parameters for disease diagnosis, and in the question if expression profiles of differentially expressed genes are also correlated to other clinical parameters. The applied strategies were linear regression model and weighted correlation network analysis (WGCNA) [36]. The regression analysis showed a strong impact of the disease index (Harvey-Bradshaw Index (HB) for CD and Simple Clinical Colitis

Activity Index (SCCAI) for UC) (p-value < 10e-6, t-test) which by definition is positively correlated to the severity of the disease, and a significant impact of smoking (p-value < 0.05, t-test) which was however, 3.5 times lower than the disease index and lower than the error rate. However, biopsy location did not show any significant effect on the severity of the disease. In agreement, the network analysis identified 10,435 genes significantly correlated to the disease index (p-value < 0.05, t-test), which is a clinical parameter with most related gene expression profiles. However, only 509 of these genes were differentially expressed between disease and control. Conversely, the expression profile of 1006 differentially expressed genes significantly associated with age (p-value < 0.05, t-test; Additional file 1: Table S10). These results suggest that even though the sample diagnosis for disease is only partly related to other clinical parameters, especially disease index and smoking, many differentially expressed gene significance measures for all genes in a given module are summarized in Additional file 1: Table S10.

Overall, the network analysis identified three large co-expression modules enriched for differentially expressed genes between iCD / iUC and control (p-value < 10e-100, Pearson's Chi-squared test; Additional file 2: Figure S10 and S11). The three modules comprised of 2054 out of 2737 differentially expressed genes. The gene network of the "brown" module was found to be enriched for immune and pro-inflammatory response (Additional file 1: Table S11), the "green" and "red" module were driven by genes involved in small molecule trans-membrane transport, anionic and cationic transport (Additional file 1: Table S12 and S13). GO analysis was also performed for the randomized gene sets of the same module sizes. None of the randomized modules had significant GO terms.

Discussion

The present study was intended to explore the transcriptomic landscape of the lncRNAs in IBD, with particular focus on CD and UC. To explore the transcriptomic profiles of CD and UC patients, colonic pinch biopsies were analyzed using gene expression microarrays. Our results revealed widespread dysregulation of lncRNAs and protein-coding gene expression in both CD and UC. It is noteworthy that although our main focus was transcriptome analysis of lncRNAs, we also profiled a significant number of protein-coding genes (~12,000, see methods). The Gencode v15 lncRNA microarray has been extensively used and the levels of both mRNAs and lncRNAs are comparable and show strong correlations (ranging from 0.62) to 0.75) with results obtained from RNA sequencing (RNAseq) [38]. These correlations are also comparable with the previous lncRNA microarray versions [35]. The Gencode v15 lncRNA microarray has been designed to capture both poly(A) and non-poly(A) transcripts (out of total 22,007 lncRNA transcripts targeted by the microarray, 9273 lncRNA transcripts are polyadenylated). In recent years, many studies have been conducted to profile lncRNAs using RNAseq, however, it is expensive and time consuming because of the requirement of doing deep sequencing particularly lncRNAs which are relatively expressed at lower levels than the protein-coding genes [38]. It has also been reported that microarrays are more sensitive to detect whether a lncRNA is expressed or not as compared to RNAseq [40].

SVM based classifiers have been previously used to cross-validate the circulating microRNA based biomarker panels in UC [13]. We also verified robustness of the differentially expressed genes by SVM and the predictive capability of these genes to discriminate CD and UC was tested using SVM-RFE based classifiers. HB Index and SCCAI are symptom based indices used to assess the disease activity in CD and UC, respectively. Among various

clinical parameters tested, we found strong influence of disease index followed by smoking, age and sex, on iCD and iUC phenotypes. Smoking is known to have deleterious effects in CD while it has been found to be protective against UC [41]. Furthermore, smoking has also been known to influence colonic gene expression profile in CD [42]. Additionally, based on linear regression and WGCNA analysis, we did not find any significant effect of biopsy location on overall gene expression. However, regional variation in gene expression along the colonic mucosa has been reported to have influence on the expression profiling studies in IBD [43,44]. These modest regional variations are more pronounced in healthy controls and un-inflamed biopsies and largely remain masked when comparing inflamed biopsies [44]. On the contrary, other studies suggest no such gene expression differences due to regional variation [17,45]. These reports highlight the importance and impact of various confounding factors like smoking, sex, biopsy locations, among many other clinically relevant parameters in gene expression analysis in IBD.

Our analysis identified common expression patterns between the lncRNAs and proteincoding genes in iCD and iUC as confirmed by unsupervised hierarchical clustering (Figure 3). A distinctive inflammatory (iCD/iUC) gene expression signature included the key drivers of the innate and adaptive immune responses (chemokines, cytokines and defensins) for example DUOXA2 [dual oxidase maturation factor 2], CXCL1 [chemokine (C-X-C motif) ligand 1], CXCL9 [chemokine (C-X-C motif) ligand 9] and also included a significant number of lncRNAs. Expression levels of both DUOX2 and DUOXA2 have been reported to be up-regulated in association with iUC, and in UC-CRC (UC-associated colorectal dysplasia and colorectal cancer) and are involved specifically in inflammation and regulated on a cryptby-crypt basis in UC [46]. We also observed a global up-regulation of DUOXA2 in iCD and iUC when compared to both non-inflamed and healthy controls. Both DUOX2 and its maturation factor DUOXA2 are part of the NADPH oxidase family of enzymes involved in release of hydrogen peroxide (H₂O₂) [47]. These enzymes are essential components of evolutionarily conserved mechanisms through which organisms are known to defend themselves against bacterial, viral, or parasitic infections, yet allowing tolerance of commensals [48,49] Suppression of DUOX2-generated H₂O₂ production by Mesalazine (5aminosalicylic acid; 5-ASA) has been demonstrated to reduce reactive oxygen species (ROS)-induced genetic lesions and thereby lowering the risk of UC-CRC [46].

Our results revealed significant down-regulation of lncRNA ANRIL [antisense non-coding RNA in the INK4 locus] in both iCD (FC < -2.7, p-value < 0.05) and iUC (FC < -7.9, pvalue < 0.05) when compared to non-inflamed and healthy controls. ANRIL, encoded on the chromosome 9p2.3 region is a known hotspot for the disease-associated SNPs [50]. ANRIL has emerged as an important regulatory molecule mediating human disease at various levels and cellular settings. Nevertheless, the role of ANRIL has not yet been described specifically in context of the IBD pathology. ANRIL has been found to be up-regulated in leukemia, prostate cancer, basal cell carcinoma and glioma, whereas, depletion of ANRIL has been implicated with reduced proliferation, indicating its role in cancerogenesis [51-53]. Remarkably, in our study, eight major ANRIL isoforms, including the isoforms known to form circular variants (*cANRIL*), were found to be universally down-regulated in both iCD and iUC. Importantly, endogenous expression of *cANRIL* has been associated with risk for atherosclerosis [54]. In this context, dysregulation of ANRIL in IBD is highly intriguing, particularly the down-regulation of *cANRIL* isoform. Indeed, recently, circular RNAs have been shown to be involved in stabilizing the sense transcripts and also act as sponges for microRNAs [55] however; biological functions of circRNAs have recently been debated [56].

It is therefore imperative to investigate comprehensively the potential roles of *cANRIL* in IBD pathogenesis.

Unsurprisingly, our results also enabled us to distinguish between iCD and iUC, although the number of differentially expressed genes were small, which emphasizes the close pathogenic nature of CD and UC. An interesting distinction between iCD and iUC expression was observed in *SERPINB3* [serpin peptidase inhibitor, clade B (ovalbumin), member 3] expression, where *SERPINB3* was significantly down-regulated (FC < -3.8) in iCD vs iUC contrast (Table 6 and Figure 1G). *SERPINB3* has been found to be over-expressed in certain squamous epithelial cancers, for instance uterine cervix carcinoma, head and neck carcinomas, and esophagus carcinoma [57] Although, the precise physiological functions of *SERPINB3* are elusive, it has been hypothesized that *SERPINB3* might be involved in the development of autoimmunity [58].

In our study, we found significant enrichment for the 96 differentially expressed lncRNAs within IBD loci. Collectively, we found differentially expressed IBD loci-associated IncRNAs overlapping active regulatory elements in intestinal epithelium and immune cells located within known binding motifs [7]. LncRNA RP3-395 M20.8 was found to be associated with the regulatory IBD risk variant rs10797432 which affects the binding motifs for AP-2 [transcription factor AP-2 alpha (activating enhancer binding protein 2 alpha)] and *CTCF* [CCCTC-binding factor (zinc finger protein)] (Additional file 1: Table S5). Moreover, IBD risk variant rs1569723 is known to act as a *cis*-eQTL for *CD40* [CD40 molecule, TNF receptor superfamily member 5] which was significantly up-regulated in iUC and associated with lncRNA RP11-465 L10.10. Additionally, lncRNA IFNG-ASI harboring UC susceptibility SNP rs7134599 was found to be up-regulated in iUC. SNP rs7134599 is associated with IBD26 (12q15) genetic locus and with regulatory pro-inflammatory cytokines IFNG [interferon, gamma] and IL-2 [interleukin 2] and anti-inflammatory cytokine IL-26 [interleukin 26]. *IFNG* gene encodes interferon gamma (*IFN*- γ), a soluble cytokine that is pivotal for the host's innate and adaptive immunity against viral, certain bacterial and protozoal infections. Aberrant expression of $IFN-\gamma$ has been linked with a number of autoimmune and inflammatory diseases, and mucosal expression of $IFN-\gamma$ is known to play a vital role in pathogenesis IBD [59]. IL-2 is encoded by IL2 gene and is involved in immune responses to microbial infections and intestinal inflammation activation in IBD. Antiinflammatory IL-26 has been shown to be overexpressed in CD [60]. These findings suggest potential involvement of differentially expressed lncRNAs overlapping the active regulatory elements in IBD pathogenesis.

Interestingly, we also found positive $(r^2 \ge 0.5)$ and extreme positive $(r^2 \ge 0.9)$ correlations between the overlapping as well as *cis*-neighboring differentially expressed IBD lociassociated lncRNA-protein-coding gene pairs. A strong positive correlation was observed between lncRNA *AC051649.12* and protein-coding gene *LSP1* [lymphocyte-specific protein 1] associated with IBD risk variant rs907611. SNP rs907611 affects the binding affinity of transcriptional factors *YY1* and *NF-muE1* and thus alters the gene expression. It is plausible that the differentially expressed IBD loci-associated lncRNAs intersecting protein-coding genes somehow contribute to the regulation of the latter [61]. Taken together, these data suggests role of lncRNAs in regulating the expression of IBD loci-associated genes.

Additionally, we also noticed dysregulation of AMPs and inflammatory response genes such as, pro-inflammatory chemokines and cytokines in various clinical subgroups. For example, key antimicrobial response genes, *REG3A* (Regenerating islet-derived 3 alpha), *DEFA5*

(Defensin, alpha 5, Paneth cell-specific) and *DEFA6* (Defensin, alpha 6, Paneth cell-specific), were >30 fold up-regulated specially in iCD vs control. Consistent with our results, *REG3A*, *DEFA5* and *DEFA6*, have been shown previously to be significantly up-regulated and linked to Paneth cell metaplasia in IBD [62,63]. Mutations in the cytoplasmic pathogen recognition receptor *NOD2* (nucleotide-binding oligomerization domain containing 2) gene have been associated with ileal CD and Paneth cell dysfunction [64] and importantly, *NOD2* was found to be up-regulated in both iCD and iUC. Concordant with the findings by Arjis et al., we also found two AMPs *DEFB1 and NPY* significantly down-regulated in both iCD and iUC [59]. *IL15* (interleukin 15) was found specifically up-regulated in iCD and not in iUC which supports the notion that it contributes to acute intestinal inflammation in CD [65].

For all the differentially expressed lncRNAs and protein-coding genes, we evaluated biological functional processes through analysis of GO terms based on "guilt-by-association" and WGCNA approach. Unsurprisingly, we found enrichment for immune response, proinflammatory cytokine activity, extracellular matrix organization, and ion membrane transport genes (Additional file 1: Table S9, S11, S12 and S13). Given the idiopathic nature of IBD, the overall up-regulation of pro-inflammatory immune response-related gene expression manifestation could be largely due to the infiltrating immune cells, rather than due to the underlying disease phenotype. Indeed, persistent inflammation in CD and UC is known to be elicited by the activation of innate and adaptive immune cells by foreign antigens, which in turn produce and release pro-inflammatory cytokines that give rise to the acrimonious circle of inflammation thereby leading to chronic tissue injury and epithelial damage [66]. Nevertheless, differentially expressed genes identified in non-inflamed (niCD and niUC) vs control (Additional file 1: Table S3 and S4), might be disease specific. In summary, our findings suggest that dysregulated lncRNAs could be involved in the IBD pathogenesis. However, these findings warrant a systematic experimental follow-up in cellular and murine based models with additional validation in a larger cohort in order to elucidate the role and biomarker potential of these dysregulated lncRNAs in IBD.

Conclusions

In conclusion, we show that lncRNA expression profiling can be effectively used to stratify iCD and iUC from healthy controls. Additionally, our data indicates the underlining potential of lncRNA transcriptional signatures associated with clinical parameters as biomarkers for IBD.

Competing interests

The authors declare that they have no competing interests.

Authors' contributions

Conceived and designed the experiments: FP, JG, CHBB, AHM, MV, KSF. Performed the experiments: AHM, CHBB, SES, XP. Analyzed the data and wrote the manuscript: AHM. Intellectual guidance: FP, JG. All authors read and approved the final manuscript.

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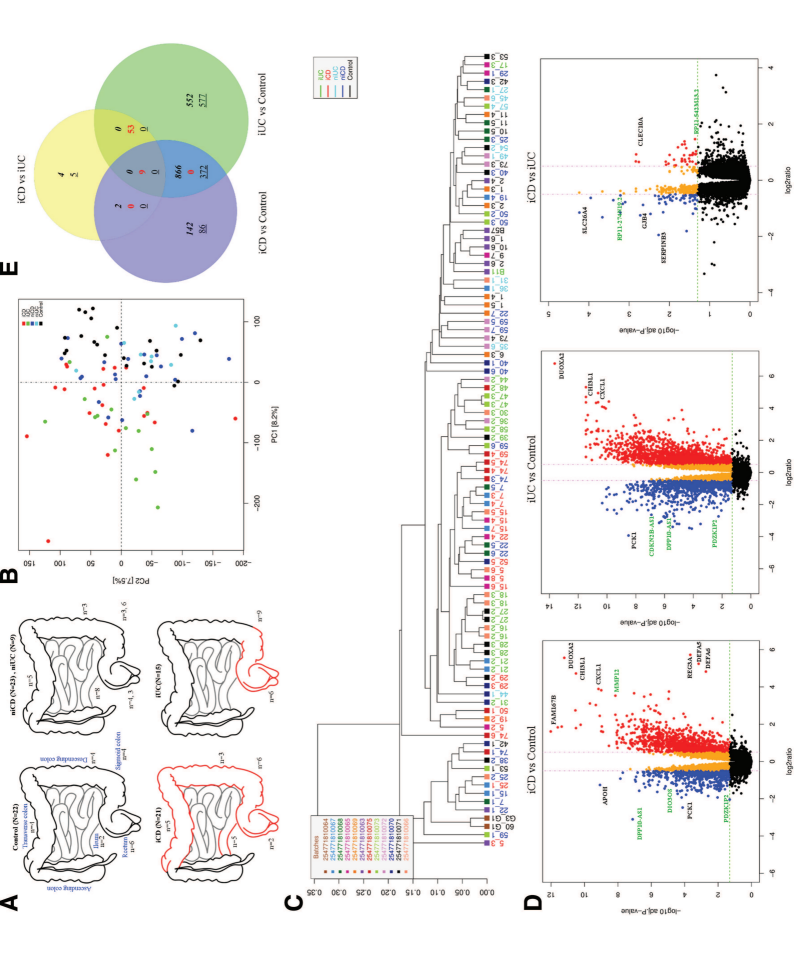
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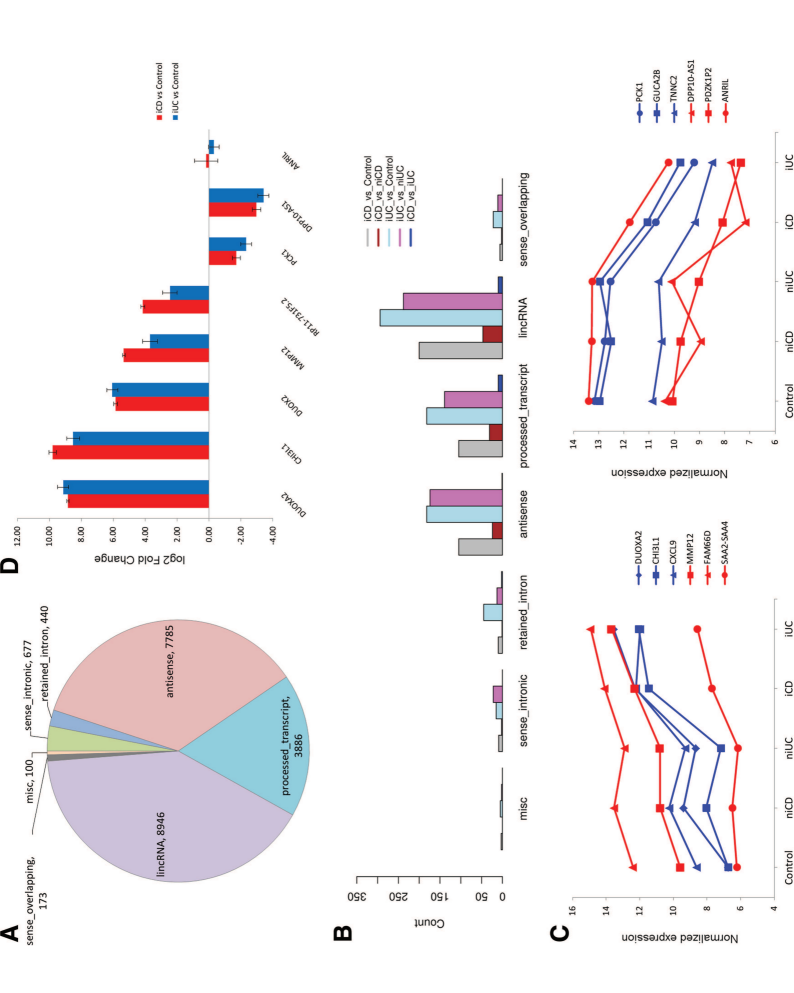
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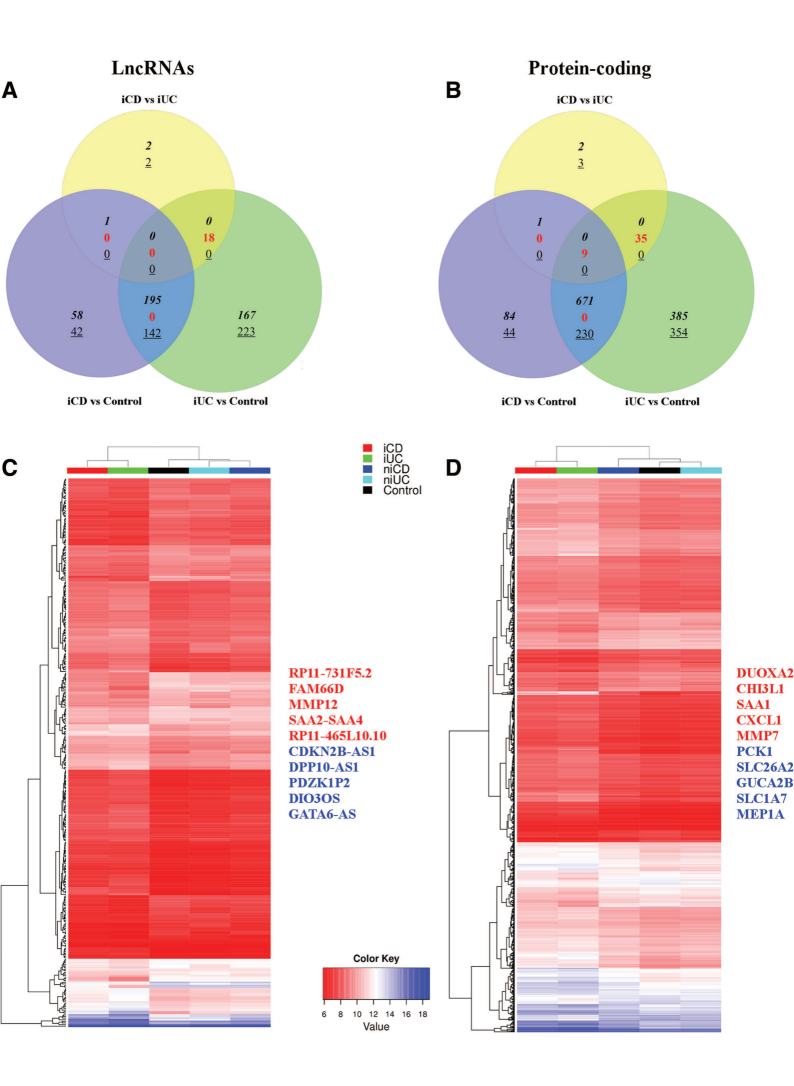
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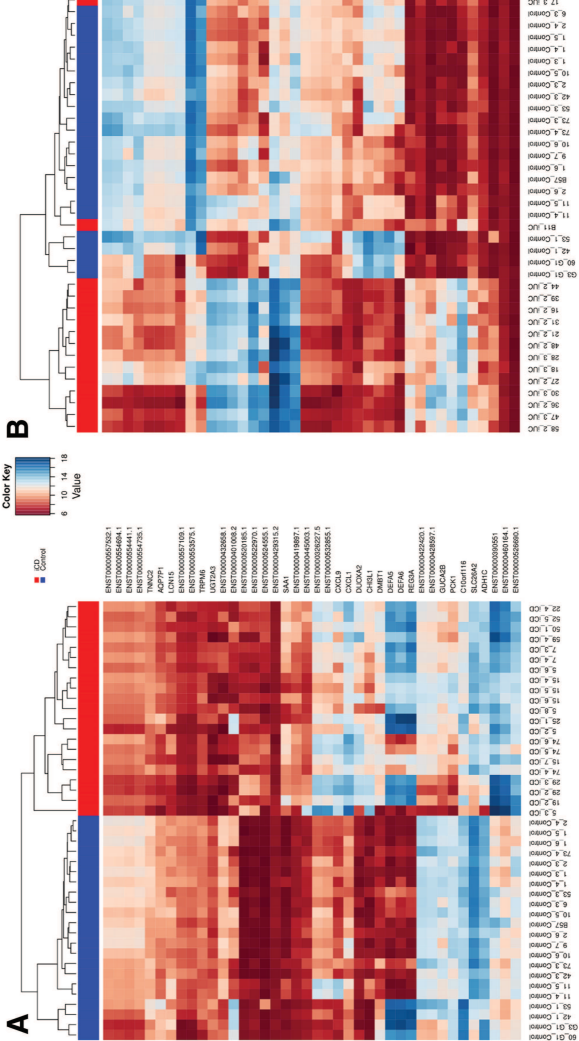
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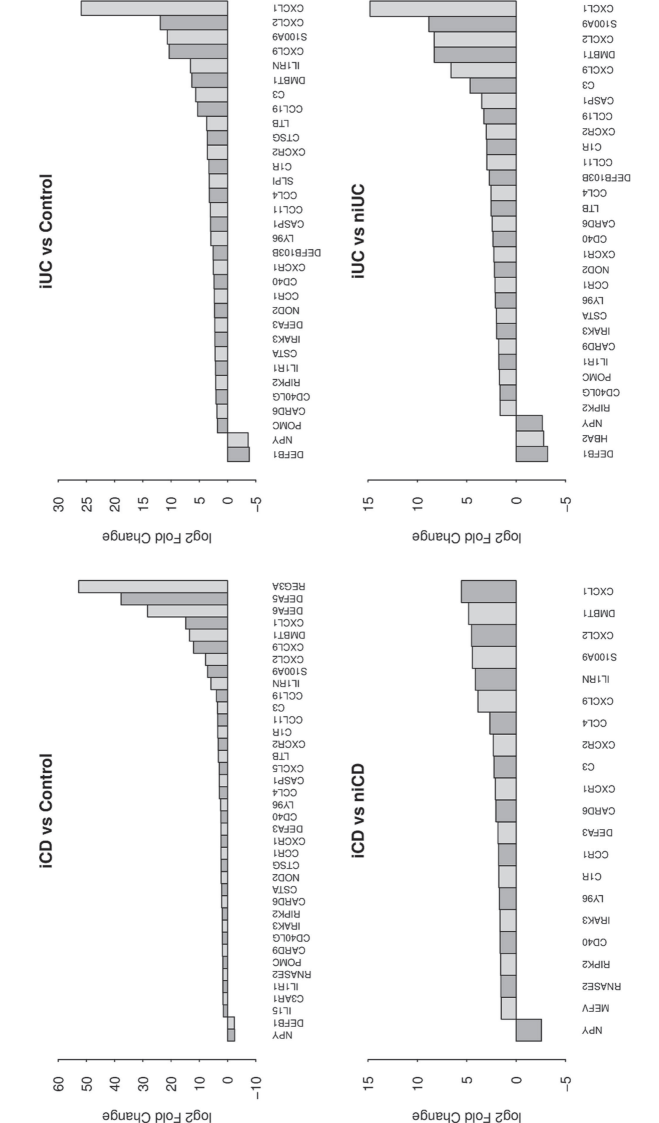


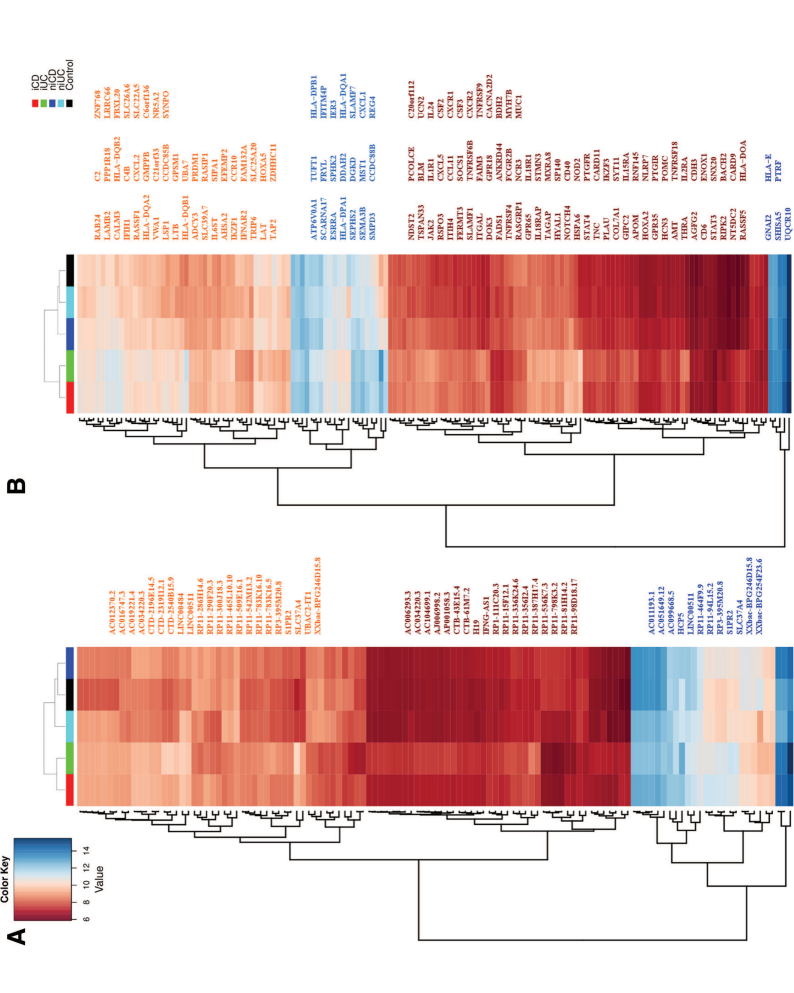


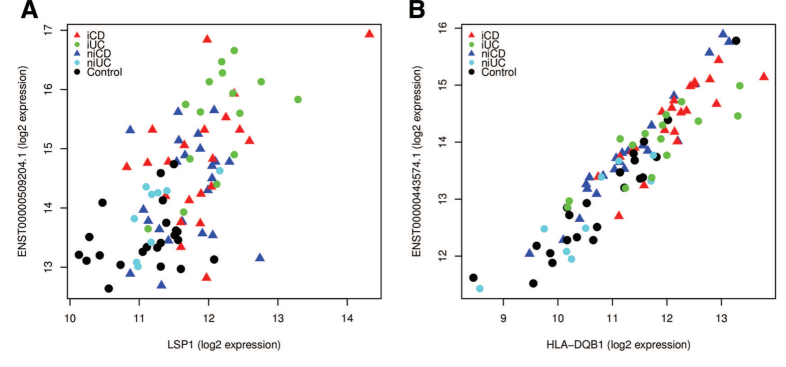
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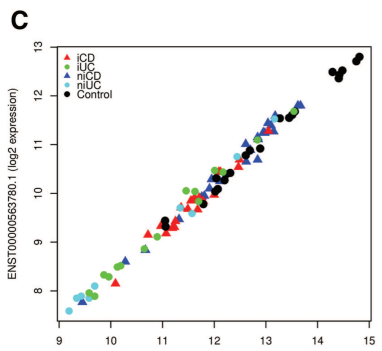
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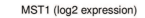
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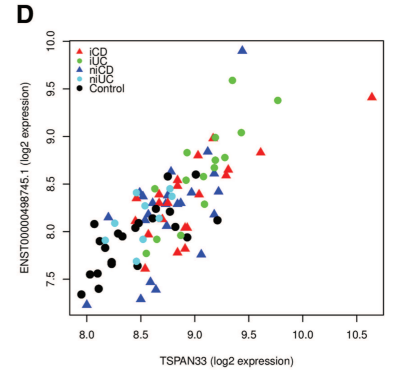


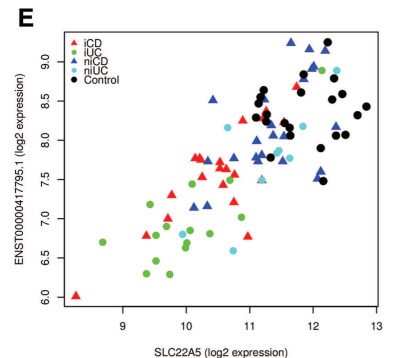


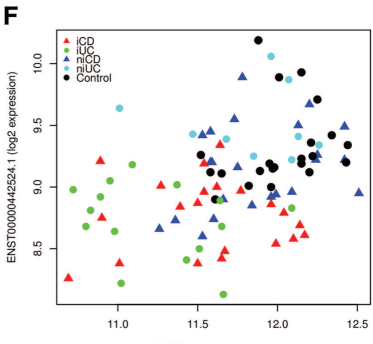




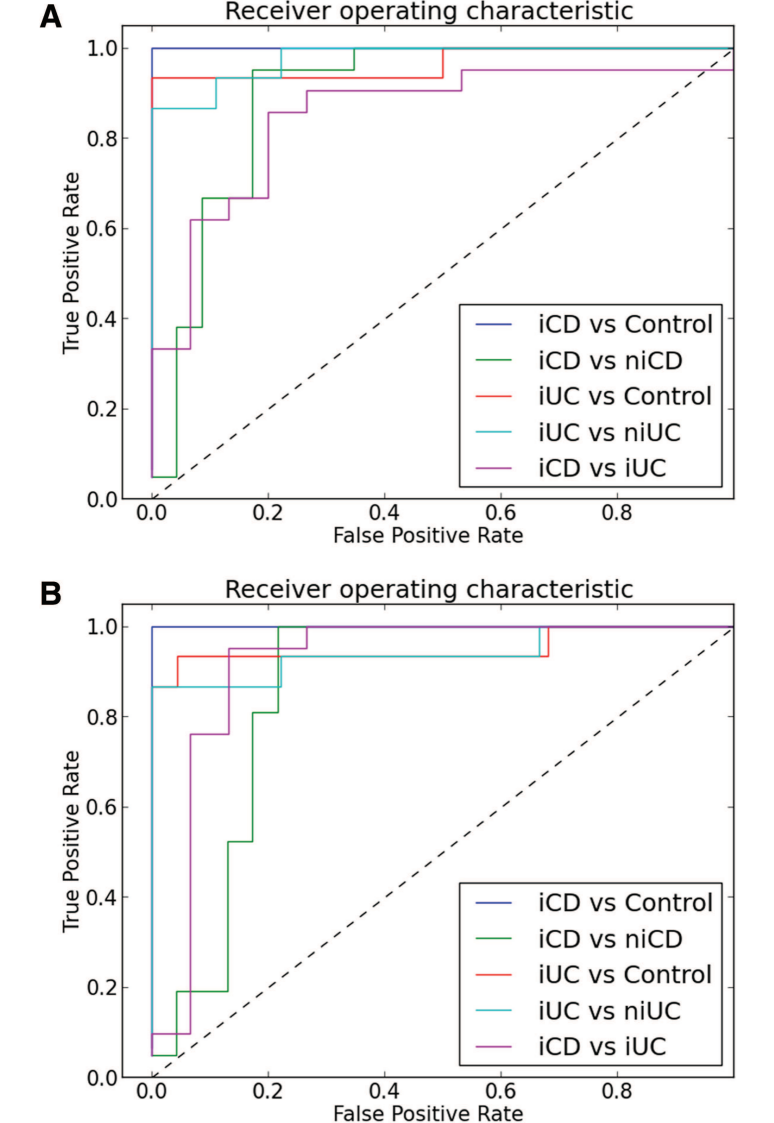








DGKD (log2 expression)



Additional files provided with this submission:

Additional file 1. Table S1, S2, S3 and S4: The top differentially expressed lncRNAs and protein-coding genes for inflamed vs. non-inflamed tissues in CD and UC. Table S1: Top 10 up-down regulated lncRNAs and protein-coding genes in iCD vs niCD. Table S2: Top 10 up-down regulated lncRNAs and protein-coding genes in iUC vs niUC. Table S3: Common differentially expressed genes found between iCD vs control and niCD vs control. Table S4: Common differentially expressed genes between iUC vs control and niUC vs control. Table S5, S6 and S7: Table S5: List of oligonucleotides used for validating microarray data using real-time PCR analysis for selected 15 genes. Table S6: Validation of microarray results by Real-time PCR analysis for 8 differentially expressed genes in iCD vs control and iUC vs control. Table S7: Validation of microarray results by Real-time PCR analysis for 5 differentially expressed genes in iCD vs iUC. Table S8: 96 differentially expressed lncRNAs found significantly enriched within IBD-loci. Table S9: Functional Annotation of differentially expressed lncRNAs based on nearest neighbor approach. Table S10, S11, S12 and S13: Weighted correlation network analysis (WGCNA) analysis. Table S10: Gene significance. The co-expression network identified the gene significance for the different clinical parameters. Table S11: Brown module- Immune and inflammatory response: 4216 genes in the module of which 1748 have an Entrez ID (considered in GO analysis).. Table S12 -Green module - Small molecule trans-membrane transport: 2934 genes in the module of which 1210 have an Entrez ID (considered in GO analysis).. Table S13: Red module - Anionic and cationic transport: 2486 genes in the module of which 1173 have an Entrez ID (considered in GO analysis) (625kb)

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Additional file 2. Figure S1: The Pearson correlations calculated for the technical replicated (six samples analyzed in duplicates on separate chips: 16 2, 18 3, 27 2, 28 3, 47 3 and 21 2). Figure S2: The scatterplot matrices describing the variation explained by the first four principal components for 90 biopsy samples. Figure S3: Unsupervised hierarchical clustering of the most dynamic probes (coefficient of variance >0.05) targeting lncRNAs (S3A) and protein-coding genes (S3B) across the samples in different clinical subgroups. Figure S4: The log2 ratio and -log10 adj.P-values plotted and represented as volcano plots for the non-inflamed tissues contrasts iCD vs niCD (S4A) and iUC vs niUC (S4B).. Figure S5: The expression map of the top 40 differentially expressed lncRNAs and protein-coding genes in iCD vs iUC based on unsupervised hierarchical clustering. Figure S6: The expression map of the total differentially expressed lncRNAs and protein-coding genes in iCD vs controls (S6A) and iUC vs controls (S6B) (patients in red, controls in blue) based on unsupervised hierarchical clustering. Figure S7, S8 and S9: Figure S7: Dendrogram of samples and heatmap of clinical parameters. Linear regression model and weighted correlation network analysis (WGCNA) were used to investigate the impact of clinical parameters on disease diagnosis. Figure S8: Receiver operating characteristic (ROC) curve analysis for age (a), sex (b), disease index (c), smoking (d) classification using differentially expressed lncRNAs in all five contrasts. Figure S9: Overlap of differentially expressed genes identified by LIMMA and SVM. Figure S10 and S11: Figure S10: Co-expression network is built by hierarchical clustering and Dynamic Tree Cut. Modules are clusters of highly interconnected genes. The "brown", "green" and "red" modules are enriched for differentially expressed genes between iUC / iCD and Control. Figure S11: Module significance is determined as the average absolute gene significance measure for all genes in a given module (2562kb)

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