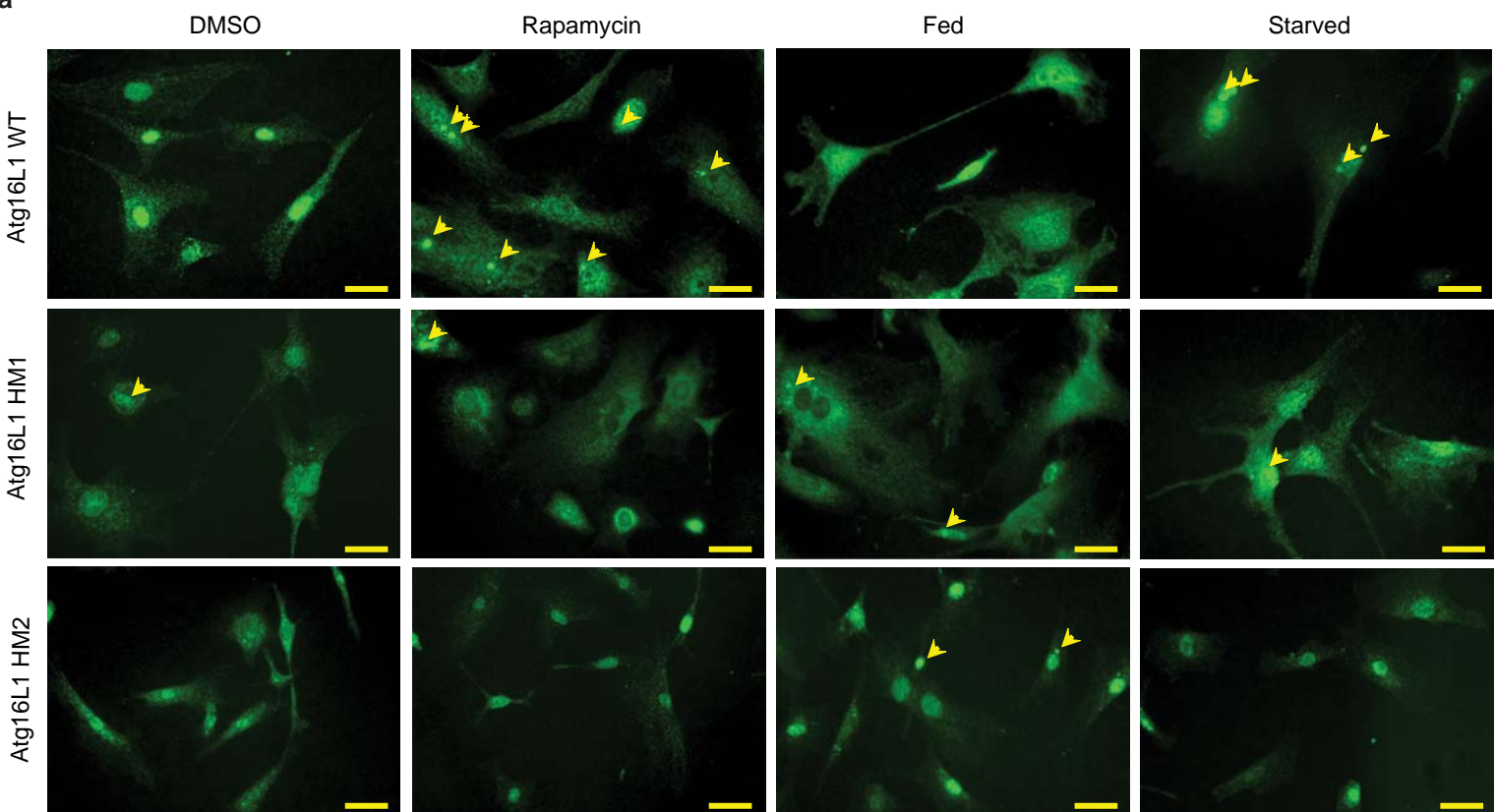
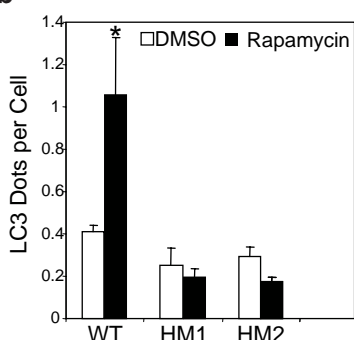
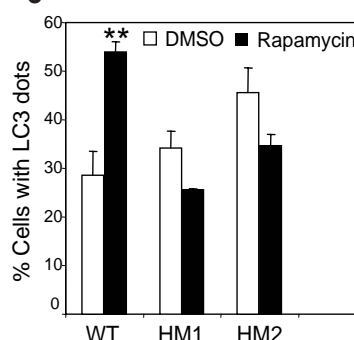
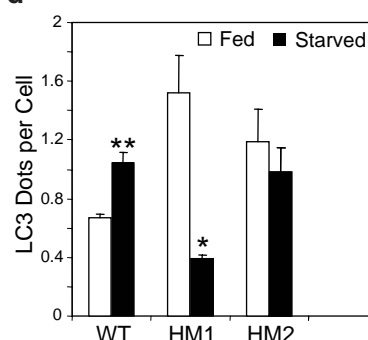
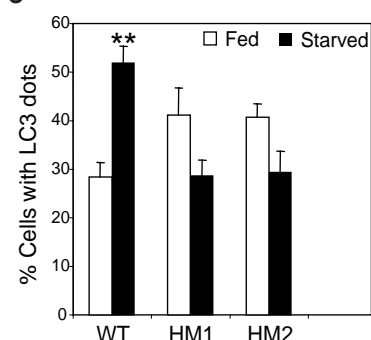
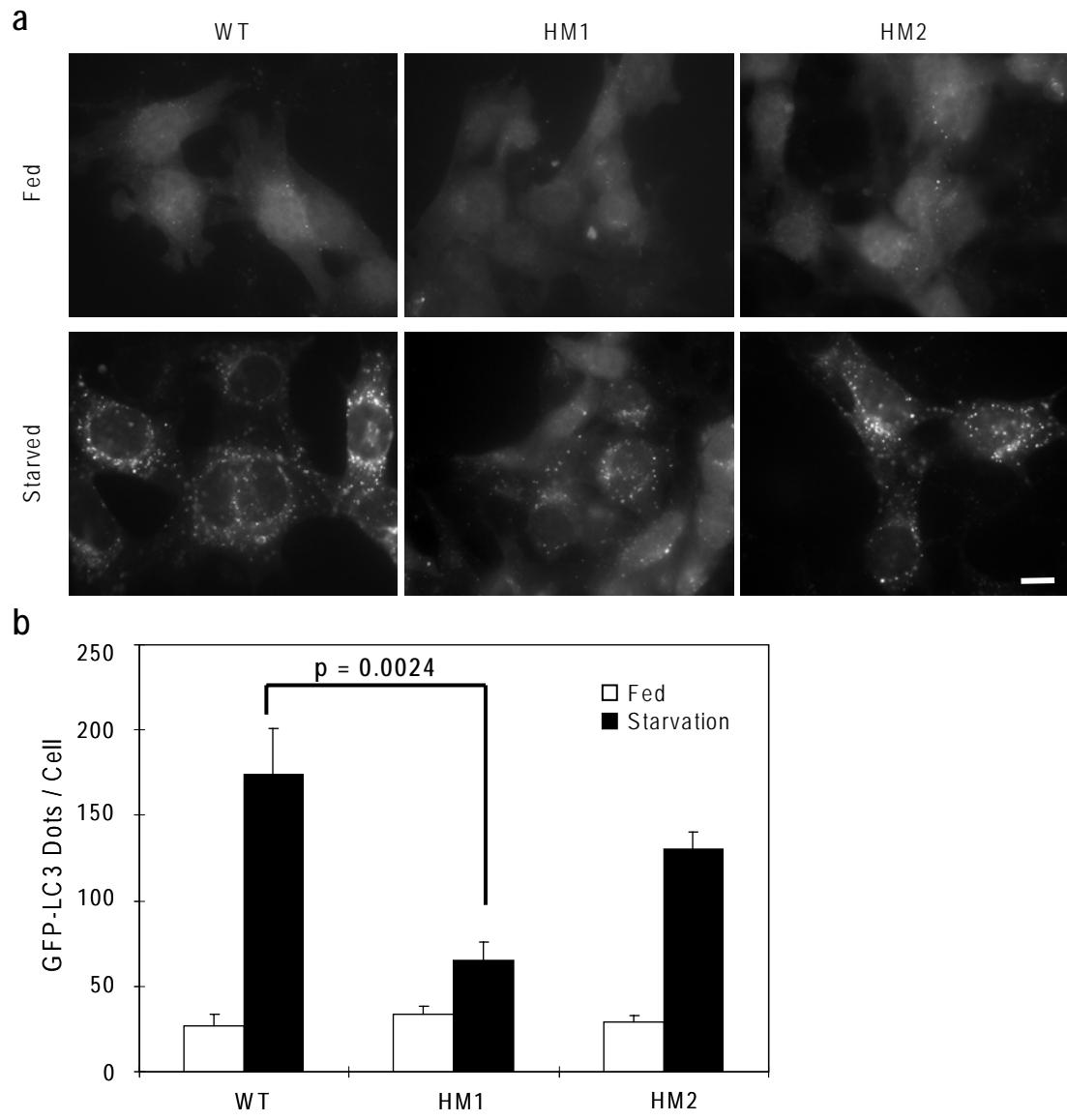


Supplementary Figure 1. Allelic inheritance in $\text{Atg16L1}^{\text{HM}}$ mice and rescue of autophagy deficiency in Atg16L1 deficient fibroblasts. a, Mice heterozygous for the gene trap mutation were bred to each other and produced wild-type (wt), heterozygous (hm/+) and homozygous (hm/hm) progeny which were separated by genotype ~4 weeks after birth. Both gene trap alleles displayed Mendelian inheritance (no significant deviation by Fisher's chi-square test, $p>0.2$ for $\text{ATG16L1}^{\text{HM}1}$ and $p>0.3$ for $\text{ATG16L1}^{\text{HM}2}$) b-c, Atg16L1 cDNA or vector control were reintroduced into $\text{Atg16L1}^{\text{HM}1}$ MEFs via retroviral transduction. MEFs were selected for successful transduction with 5 $\mu\text{g}/\text{ml}$ puromycin, then grown in the presence of the chemical inducer of autophagy rapamycin (50 $\mu\text{g}/\text{ml}$) and cyclohexamide (5 $\mu\text{g}/\text{ml}$) for 0, 4, and 8 h. Western blot analysis of cell lysates show a restoration of p62 degradation in MEFs selected for Atg16L1 expression but not vector control (b) which was quantified by densitometry and normalized to actin ($n = 3$) (c).

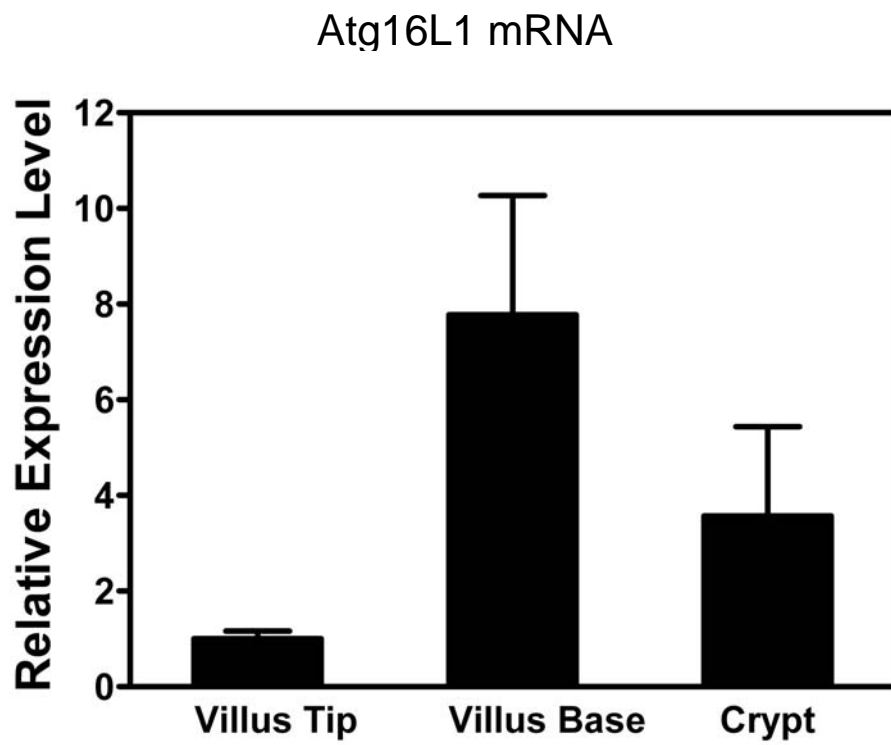
a**b****c****d****e**

*p<0.04, ** p<0.005

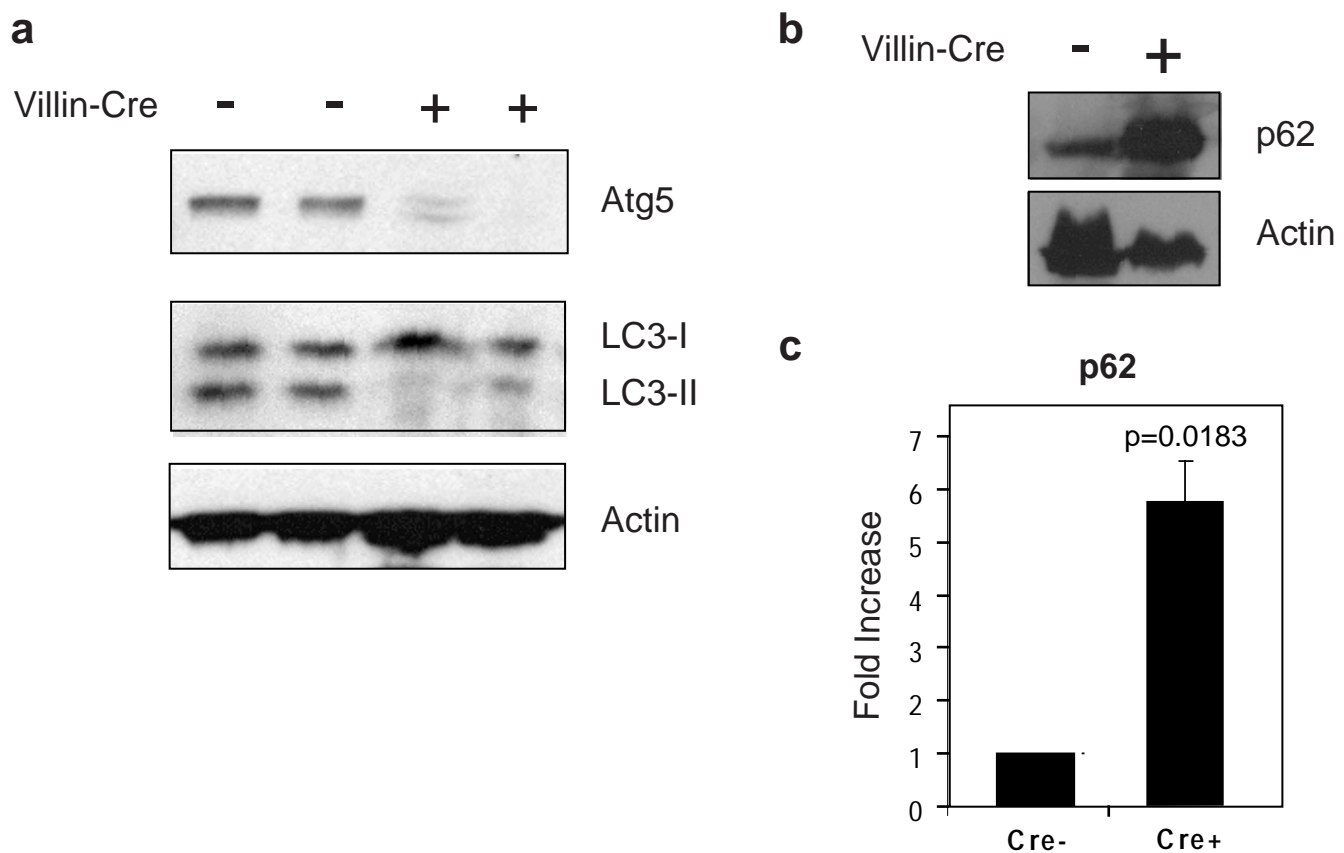
Supplementary Figure 2. MEFs derived from $\text{Atg16L1}^{\text{HM1}}$ and $\text{Atg16L1}^{\text{HM2}}$ lines show reduced endogenous LC3 dot formation. **a**, Representative immunofluorescence images of primary low-passage MEFs grown in 50 $\mu\text{g}/\text{ml}$ rapamycin for 4 h or in amino acid and serum replete EBSS media for 2 h to induce autophagy and stained with LC3. LC3-positive dots $>0.5\mu\text{m}$ in diameter are indicated by arrow heads. Scale bar, 20 μm . **b-e**, Quantification of LC3 dots revealed that both the accumulation of LC3 dots per cell (**b** and **d**) or % cells with LC3 dots (**c** and **e**) after autophagy induction were compromised in $\text{Atg16L1}^{\text{HM}}$ cells, indicating that autophagosome formation was aberrant under these conditions ($n = 3$, at least 70 cells were analyzed per sample). The increase in LC3 dots was statistically significant in all WT samples. There was no statistically significant increase in $\text{Atg16L1}^{\text{HM}}$ cells (note: $\text{Atg16L1}^{\text{HM1}}$ cells display a statistically significant decrease in dots per cell under starvation conditions). P values were calculated using two-tailed student's t test. Error bars represent SEM.



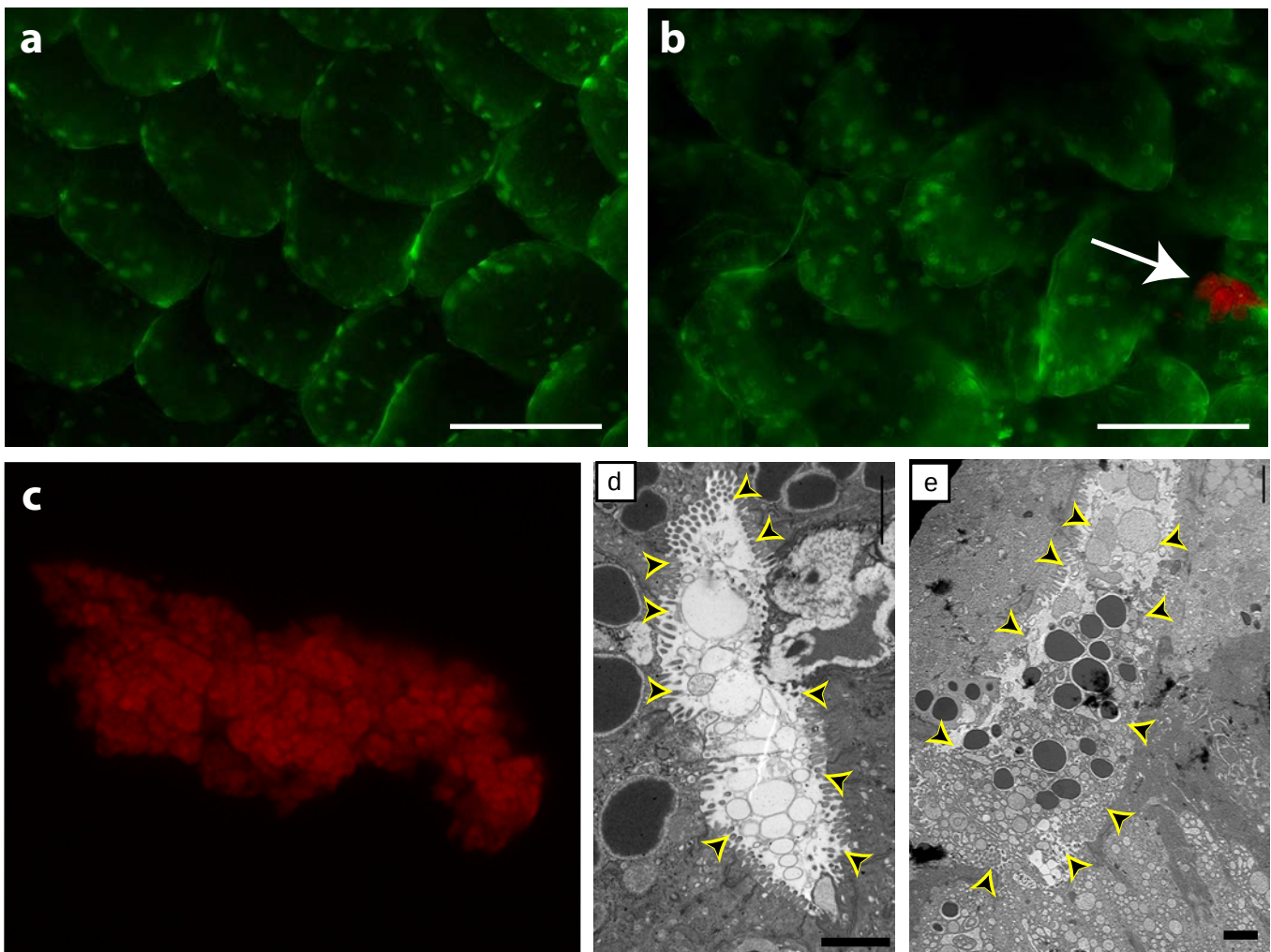
Supplementary Figure 3. GFP-LC3 dot formation is reduced in Atg16L1^{HM1} MEFs. **a**, Representative fluorescence images of immortalized Atg16L1^{HM} MEFs stably expressing GFP-LC3. Cells were cultured in DMEM with 10% FBS or DMEM without amino acids and serum for 2 h. Scale bar, 10 μ m. **b**, Quantification of the number of GFP-LC3 dots per cell (counted in at least 5 different images) show a significant reduction in dot formation in Atg16L1^{HM1} cells. Atg16L1^{HM2} cells did not show a statistically significant reduction in dot formation, consistent with the higher expression of Atg16L1 in these cells. P values were calculated using two-tailed student's t test. Error bars represent SEM.



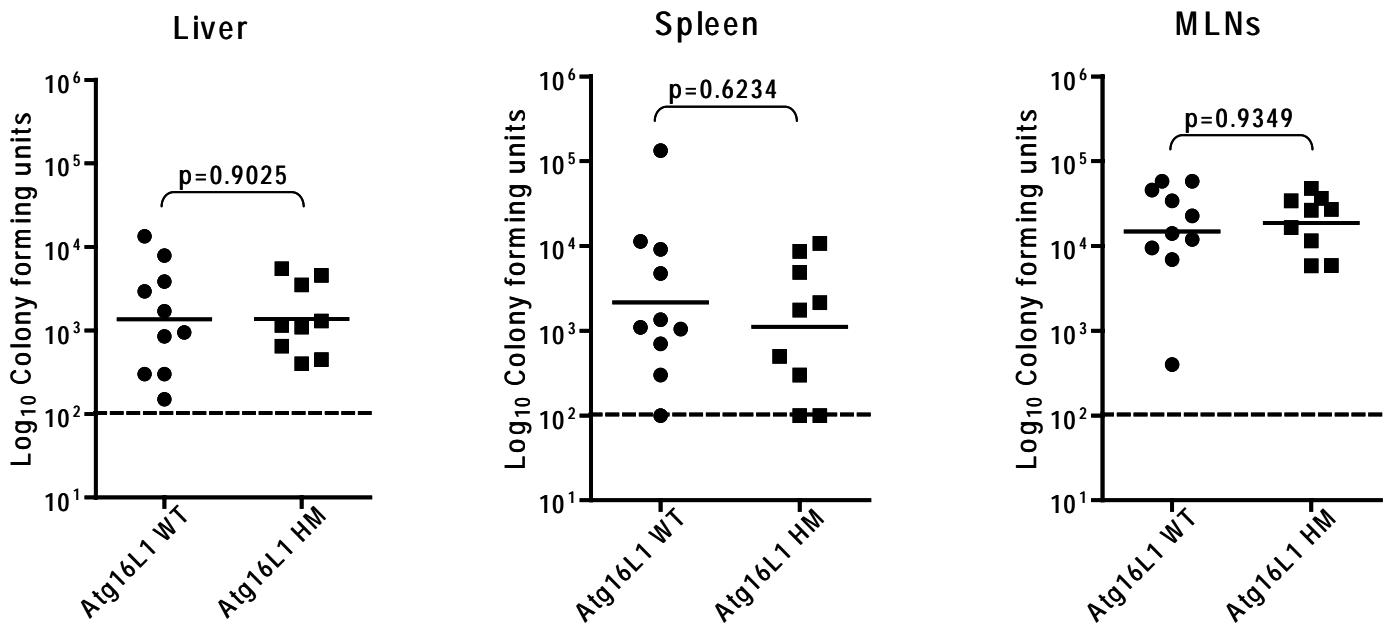
Supplementary Figure 4. Atg16L1 is expressed throughout the ileal crypt-villus axis. RNA was procured by LCM from the villus tip, villus base, and crypt base of the distal ileum from Atg16L1^{HM} mice. qRT-PCR analysis shows detectable Atg16L1 transcripts in all three compartments ($n = 3$). There was a statistically significant difference between the villus tip and the villus base ($p < 0.1$) or crypt ($p < 0.05$) indicating that Atg16L1 transcripts are enriched in the villus base and crypt. P values were calculated using two-tailed student's t test. Error bars represent SEM.



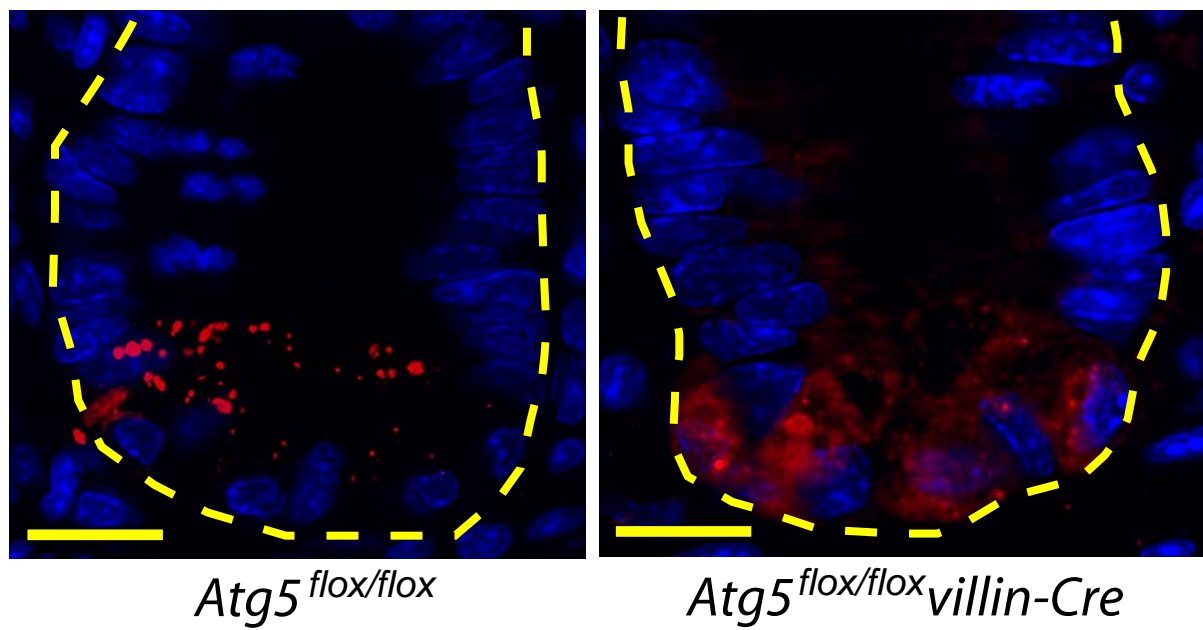
Supplementary Figure 5. Conditional deletion of *Atg5* in the intestinal epithelium leads to reduced LC3 conversion and accumulation of p62. **a**, Western blot analysis of ileal lysates from *Atg5*^{flox/flox}*villin-Cre* mice reveal decreased Atg5 expression and an increase in LC3-I to LC3-II ratio similar to Atg16L1^{HM} mice suggesting a critical role for these proteins in intestinal autophagy (n = 3 of each genotype, 2 of each shown). **b-c**, *Atg5*^{flox/flox}*villin-Cre* mice also display an increase in p62 protein expression in the ileal epithelium (b) similar to Atg16L1^{HM} mice. Quantification of p62 levels by densitometry normalized to actin revealed ~7-fold increase in *Atg5*^{flox/flox}*villin-Cre* mice (n = 3) (c). P values were calculated using two-tailed student's t test. Error bars represent SEM.



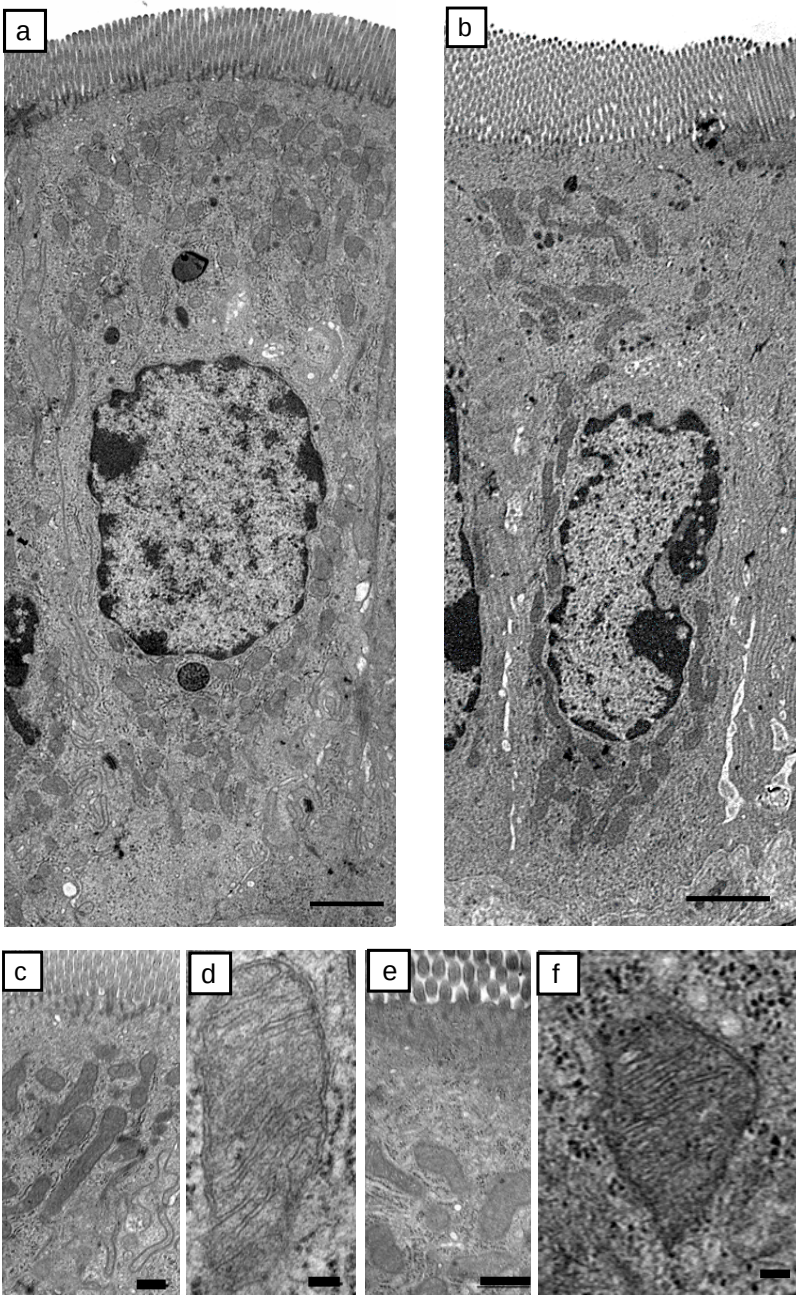
Supplementary Figure 6. Abnormal Paneth granule exocytosis in Atg16L1 deficient mice. a-c, Whole mounts of the small intestines from control (a) and Atg16L1^{HM} (b, c) mice stained with FITC-conjugated *Helix pomatia* lectin that labels goblet cell mucus (green) and antisera directed against lysozyme (red). Lectin positive goblet cells stud the surface of the villi. No accumulated mucin is shown in these fields. Strikingly, the lysozyme staining in the Atg16L1^{HM} mice is concentrated in small clusters of spherical aggregates (white arrow in b) that are present in the crypt lumen. High power view of the aggregate in (c) is 40 μm in its greatest dimension. d-e, EM analysis of the Atg16L1^{HM} ileum reveals diminished microvilli on Paneth cells (d) and the adjacent crypt lumen (indicated by arrow heads) contains intact Paneth granules and cytoplasm (e). Scale bars: a, b, 200 μm ; d, e, 2 μm .



Supplementary Figure 7. Atg16L1 mutant mice do not display increased susceptibility to oral *L. monocytogenes* infection. Littermate WT ($n = 10$) and Atg16L1^{HM} ($n = 5$ for HM1 and $n = 4$ for HM2) mice were infected orally with 10^9 *L. monocytogenes* re-suspended in 200 μ l of 5% sodium bicarbonate by gastric gavage. The number of bacteria in liver, spleen, and mesenteric lymph nodes were determined 72 h after infection and displayed no significant differences with WT. P values were calculated using two-tailed Mann-Whitney test. Horizontal bars represent the geometric mean.



Supplementary Figure 8. Mice with conditional deletion of *Atg5* in the intestinal epithelium have abnormal lysozyme distribution in Paneth cells similar to observations made in *Atg16L1* deficient mice. Representative images of indirect immunofluorescence of sections stained for lysozyme (red) in $Atg5^{flox/flox}$ control and $Atg5^{flox/flox}villin-Cre$ mouse ileal crypts. Dotted lines denote crypt unit. Scale bars: 10 μm .



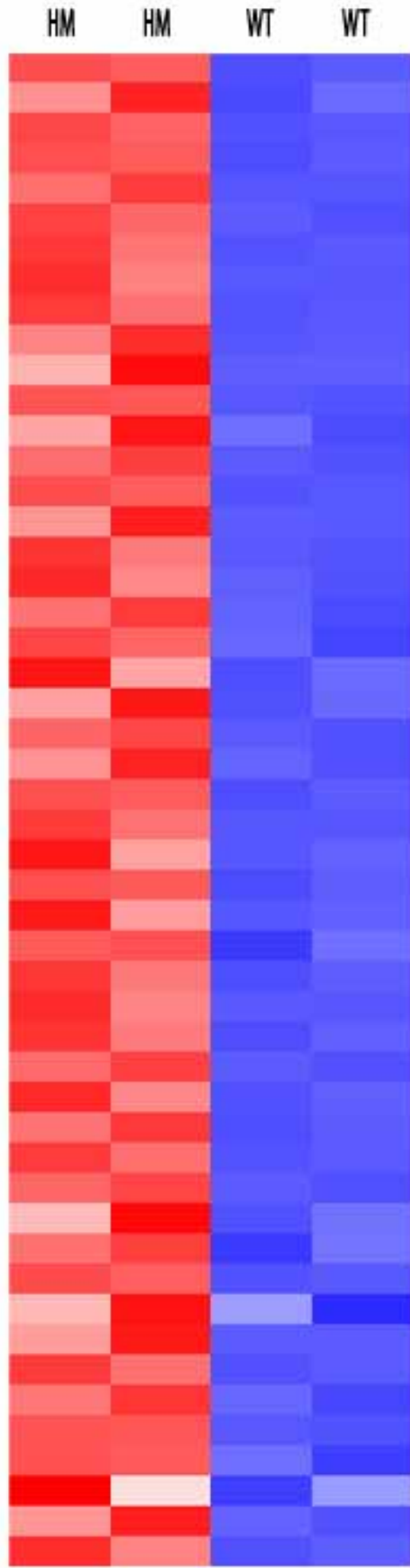
Supplementary Figure 9. Enterocytes in the ileal epithelium are similar between Atg16L1 mutant and wild-type controls by ultrastructural analysis. a-b, EM of ileal villus enterocytes from WT (a, c, d) and Atg16L1^{HM} (b, e, f) mice show similar morphology with intact mitochondria. Scale bars: a, b, 2 μm ; c, e, 500 nm; d, f, 100 nm.

Gene	qRT-PCR Validation	Fold Change	Low Bound of FC
Selenoprotein P, plasma, 1	114.0 ± 10.8	1.10	1.03
Peripheral myelin protein 22	169.3 ± 10.2	1.40	1.27
Serine/cysteine peptidase inhibitor, clade B, member 1a	14.0 ± 1.1	1.38	1.20
Serum/glucocorticoid regulated kinase 1	2.8 ± 0.1	-2.05	-1.82
BTB and CNC homology 1	2.5 ± 0.4	-1.09	-0.99
Protein tyrosine phosphatase, non-receptor type 9	3.3 ± 0.7	-1.27	-1.04
Villin 1	Enterocyte marker*	1.12	1.04
Mucin 2	Goblet cell marker*	-1.13	-1.05
Chloride channel calcium activated 3	Goblet cell marker*	1.21	1.01
Chromogranin B	Enteroendocrine cell marker*	-1.25	-1.16

Supplementary Figure 10. Validation of lack of contamination of LCM procured Paneth cells with villus epithelial cells. a, We quantified 6 genes by qRT-PCR known to be enriched in villus epithelial cells¹ and not involved in fatty acid or cholesterol metabolism. We compared gene expression in LCM-procured Paneth cells versus villus epithelial cells. The fold-enrichment for villus epithelial cells is indicated for each gene (experiments were performed in triplicate). For each of these 6 genes and 4 well-characterized markers of villus epithelial cells (indicated by an asterisk), we compared the fold-change for Atg16L1^{HM} Paneth cells versus WT. Transcripts highly enriched in villus epithelial cells were not enriched in LCM procured Paneth cells.

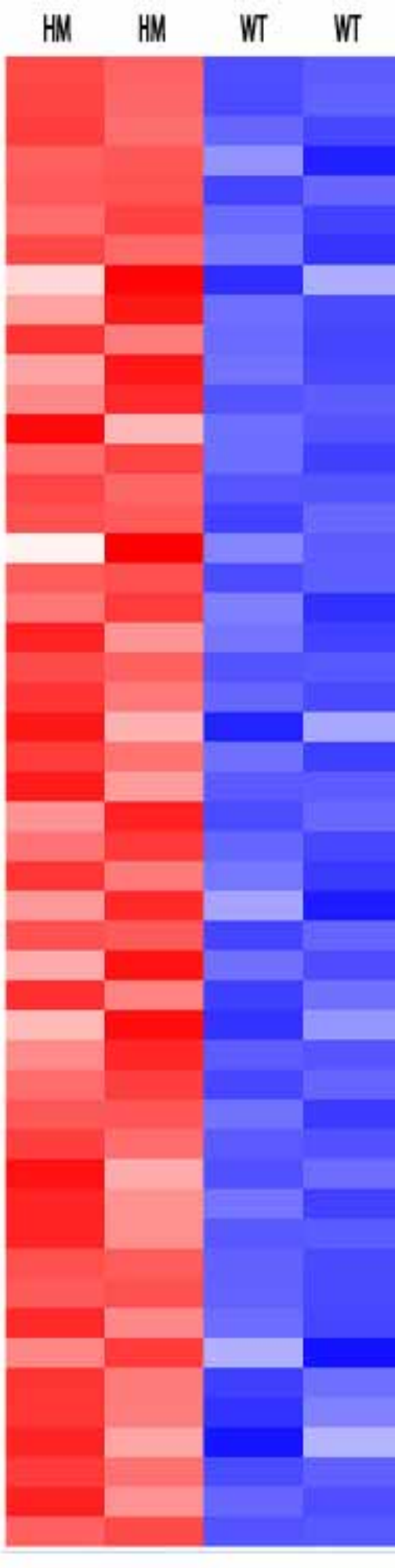
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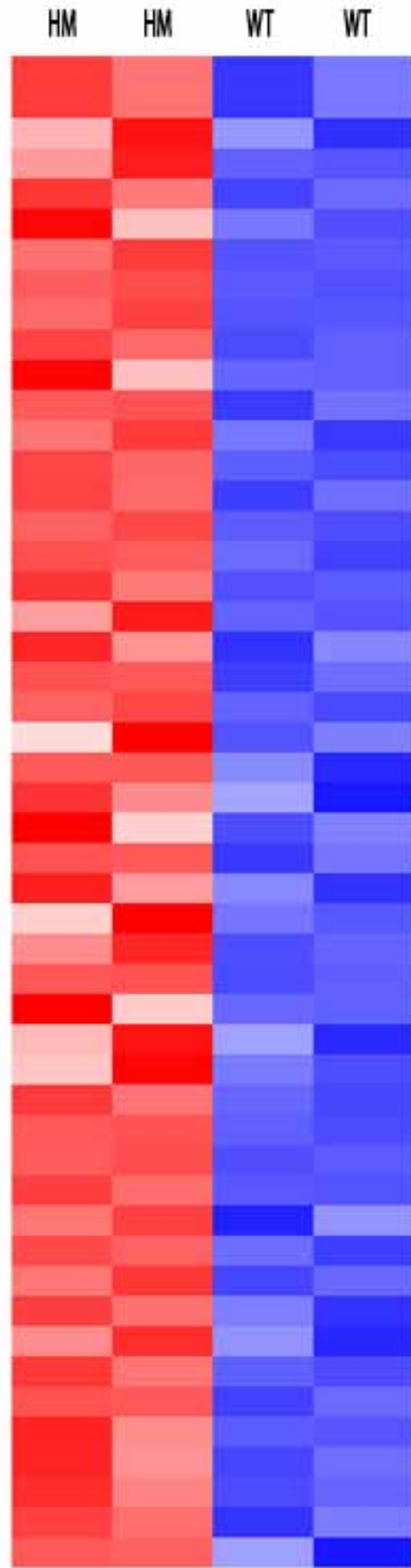


1.3 0.9 0.6 0.2 -0.2 -0.6 -0.9 -1.3

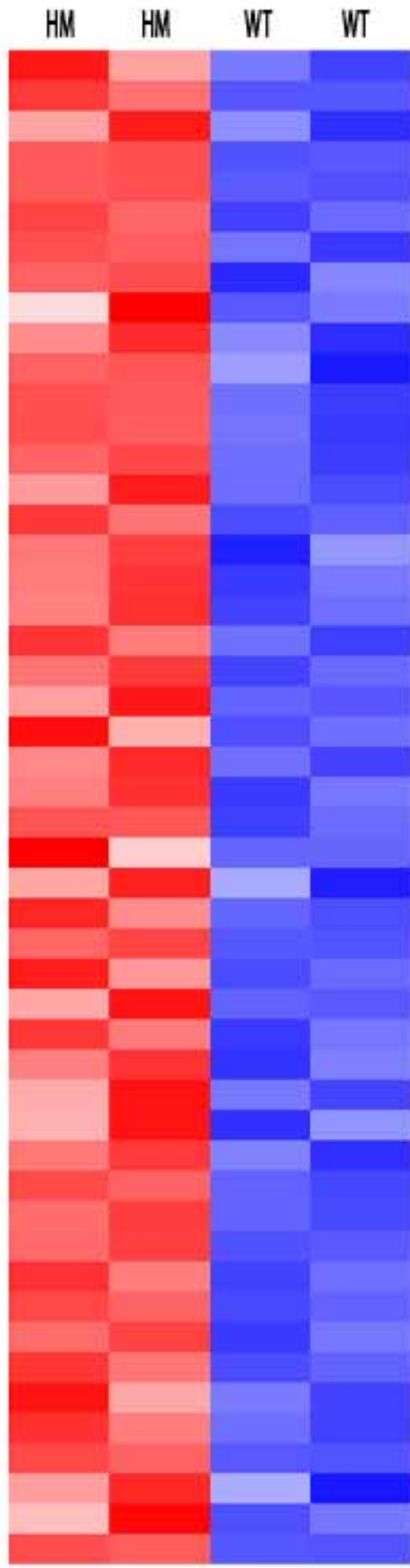
- Sod1: stearoyl-Coenzyme A desaturase 1
- Odc: cell death-inducing DFFA-like effector c
- Sod1: stearoyl-Coenzyme A desaturase 1
- LOC699888 // Lpk(lipo protein lipase) // similar to Lipoprotein lipase precursor (LPL)
- Car3: carbonic anhydrase 3
- BB144871: expressed sequence BB144871
- Thrsp: thyroid hormone responsive SPOT14 homolog (Rattus)
- Pldm: pallidin
- Cyp2e1: cytochrome P450, family 2, subfamily e, polypeptide 1
- Fabp4: fatty acid binding protein 4, adipocyte
- Fabp1: fatty acid binding protein 1, liver
- Adipoq: adiponectin, C1Q and collagen domain containing
- Slc5a12: solute carrier family 5 (sodium/glucose cotransporter), member 12
- Fabp1: fatty acid binding protein 1, liver
- Hp: haptoglobin
- Saa1: serum amyloid A1
- Saa1: serum amyloid A1
- Lep: leptin
- Cfb: complement factor B (adipin)
- Slc36a2: solute carrier family 36 (proton/amino acid symporter), member 2
- Pox: pyruvate carboxylase
- Retnla: resistin like alpha
- Serpina10: serine (or cysteine) peptidase inhibitor, clade A [alpha-1 antiproteinase, antitrypsin], member 10
- Dhh: desert hedgehog
- Apoa4: apolipoprotein A-IV
- Cf // LOC100047470: complement component factor I // similar to complement component factor I
- Mir.2156861
- S3-12: plasma membrane associated protein, S3-12
- Krt12: keratin 12
- Apoa4: apolipoprotein A-IV
- AA481121: expressed sequence AA481121
- 201002M12Rik: Riken cDNA 201002M12 gene
- Acaa1b: acetyl-Coenzyme A acyltransferase 1B
- Cd36: CD36 antigen
- Fmn2: farnin 2
- Socs3: suppressor of cytokine signaling 3
- Bst1: bone marrow stromal cell antigen 1
- Hbb-b1 // Hbb-b2: hemoglobin, beta adult major chain // hemoglobin, beta adult minor chain
- Neurod2: neurogenic differentiation 2
- Socs3: suppressor of cytokine signaling 3
- Ct: ctan
- Kif3b: kinesin family member 3B
- Rhov: rach homolog gene family, member V
- LOC100047573 // Rbp4: retinol binding protein 4, plasma // similar to retinol binding protein
- LOC636537 // Ssr1: signal sequence receptor, alpha // similar to signal sequence receptor, alpha
- Lpk(lipo protein lipase)
- Tim15: tripartite motif protein 15
- Mir.173525.1
- Mir.161838.1
- Chac1: Chac, cation transport regulator-like 1 (E. coli)



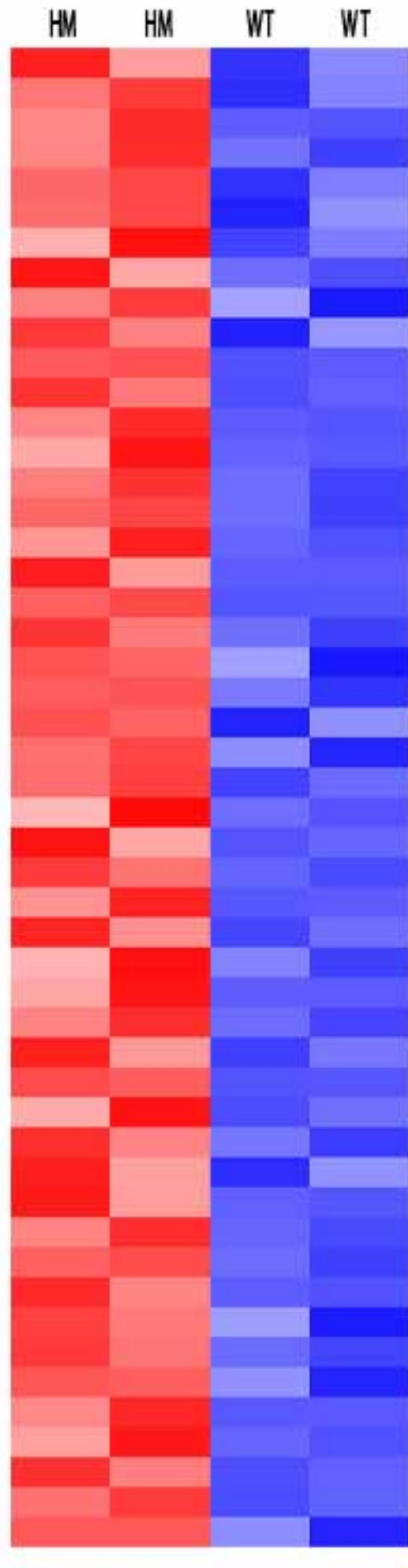
Amyl: amylase 1, salivary
Npr3: natriuretic peptide receptor 3
Hba-a1 // *Hba-a2*: hemoglobin alpha, adult chain 1 // hemoglobin alpha, adult chain 2
Sucnr1: succinate receptor 1
Kctd14: potassium channel tetramerisation domain containing 14
Ecd1: exonuclease 3'-5' domain-like 1
Gant: granzyme B
Zscan1: zinc finger with KPAB and SCAN domains 1
Dbh: dopamine beta hydroxylase
Scp2: sterol carrier protein 2, liver
Lhfp2: lipoma HMGIC fusion partner-like 2
Foxo4l: F-box protein 41
Mm.214071.1
LOC552908: hypothetical LOC552908
Mtp: microsomal triglyceride transfer protein
LOC100046877: similar to developmentally regulated RNA-binding protein 1
Phf21a: PHD finger protein 21A
Tmem116: transmembrane protein 116
Chst1: carbohydrate (keratan sulfate Gal- β) sulfotransferase 1
Mm.214384.1
Car3: carbonic anhydrase 3
Arsc: arylsulfatase K
Hspa12a: heat shock protein 12A
Upk1b: uroplakin 1B
Wif1on2: WIF, follistatin/kazal, immunoglobulin, kunitz and netrin domain containing 2
Rdak1: PDZ domain containing 1
Alpt: alkaline phosphatase, intestinal
G53001100681c: RIKEN cDNA G530011006 gene
Tmle: transmembrane inner ear
Hba-a1 // *Hba-a2*: hemoglobin alpha, adult chain 1 // hemoglobin alpha, adult chain 2
Ap1ml1: adaptor-related protein complex AP-1, mu subunit 1
Inmt: indolethylamine N-methyltransferase
Mm.157620.1
Ggt1: gamma-glutamyltransferase 1
Thpo: thrombopoietin
Gsdmc2 // *LOC100045250*: galectin C2 // hypothetical protein LOC100045250
Dgat2: diacylglycerol O-acyltransferase 2
Wdfy1: WD repeat and FYVE domain containing 1
Gsdmc2 // *Gsdmc4*: galectin C4 // galectin C2
Cdk8t: coiled-coil domain containing 80
Tif: transferrin
Cd36: CD36 antigen
Rad2: radical S-adenosyl methionine domain containing 2
Agtr2l1: alanine-glyoxylate aminotransferase 2-like 1
Apol1: apolipoprotein A-I
Mm.207711.1
493140801481c: RIKEN cDNA 4931408014 gene
Lfc: leukocyte tyrosine kinase
120001380881c: RIKEN cDNA 1200013808 gene
Clec2h: C-type lectin domain family 2, member h



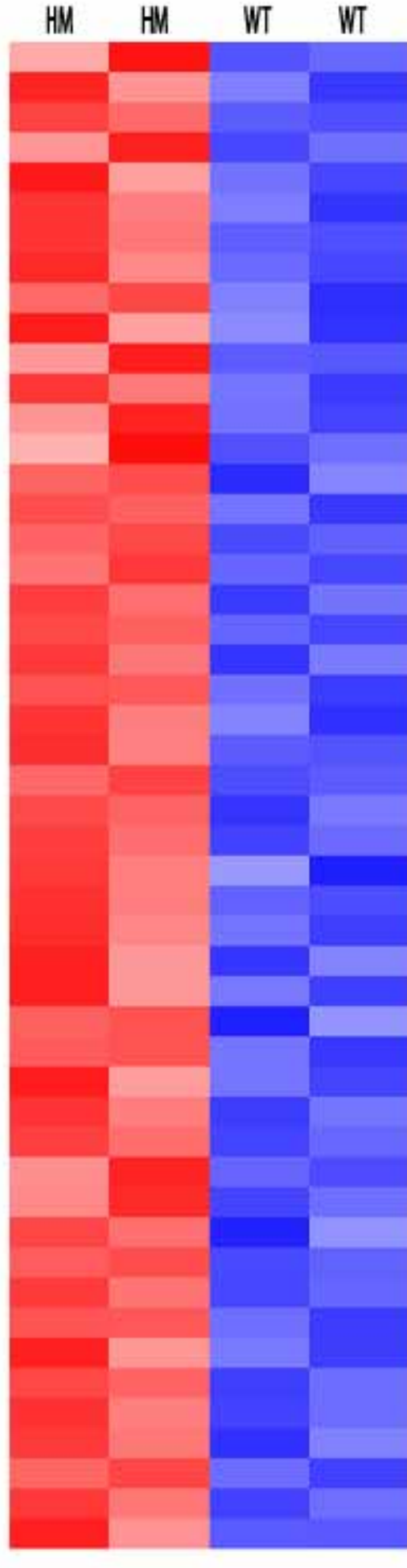
Apoa1: apolipoprotein A-I
Apoa1: apolipoprotein A-I
Mir_4687_2
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Npc1l1: NPC1-like 1
Ifit1: interferon-induced protein with tetratricopeptide repeats 1
Fgrt: Fc receptor, IgG, alpha chain transporter
Hm47: histocompatibility 47
Isg15 // LOC100038882 // LOC100044225 // LOC677168; ISG15 ubiquitin-like modifier // similar to ISG15 ubiquitin-like modifier
ORF9: open reading frame 9
 0 day neonate cerebellum cDNA, RIKEN full-length enriched library, clone C230047L12 product undetectable, full insert sequence
 LOC100041156 // LOC100041932; hypothetical protein LOC100041156 // hypothetical protein LOC100041932
Vnn1: vanin 1
Akr1b7: aldo-keto reductase family 1, member 17
Hac: hydroxyacid oxidase (glycolate oxidase) 3
 OTTMUSG0000016644; predicted gene, OTTMUSG0000016644
Apoc2: apolipoprotein C-II
 5730508809Rik: RIKEN cDNA 5730508809 gene
Fabp4: fatty acid binding protein 4, adipocyte
Tha1: threonine aldolase 1
Ptgfr3: prostaglandin E receptor 3 [subtype EP3]
 2900026402Rik: RIKEN cDNA 2900026402 gene
Cst6: cystatin E/M
 Transcribed locus
Pou4f1: POU domain, class 4, transcription factor 1
Podh17: protocadherin 17
Cdh63: CD163 antigen
Smyd1: SET and MYND domain containing 1
Eosc10: eosinone component 10
Fuz: fuzzy homolog (Drosophila)
Mir_2146_9_1
Plekhm1: pleckstrin homology domain containing, family M (with RIN domain) member 1
Cxcl1: chemokine (C-X-C motif) ligand 1
Cherp: calcium homeostasis endoplasmic reticulum protein
Myo1a: myosin Va
Tmem140: transmembrane protein 140
Ace2: angiotensin I converting enzyme (peptidyl-dipeptidase A) 2
 LOC100046740 // *Sparc*: secreted acidic cysteine rich glycoprotein // similar to Secreted acidic cysteine rich glycoprotein
Gpt1: glutamic pyruvate transaminase 1, soluble
Trib3: tribbles homolog 3 (Drosophila)
Steap4: STEAP family member 4
Vnn1: vanin 1
Bola2: bolaA-Hte 2 (E. coli)
Mip5: microtubular associated protein 5
Acacb // LOC100047358; acetyl-Coenzyme A carboxylase beta // similar to acetyl-Coenzyme A carboxylase beta
P13ap1: phosphoinositide-3-kinase adaptor protein 1
Cdc109b: coiled-coil domain containing 109B
Cibr: cubulin (intrinsic factor-cobalamin receptor)
Apob: apolipoprotein B
 6430597G12Rik: RIKEN cDNA 6430597G12 gene



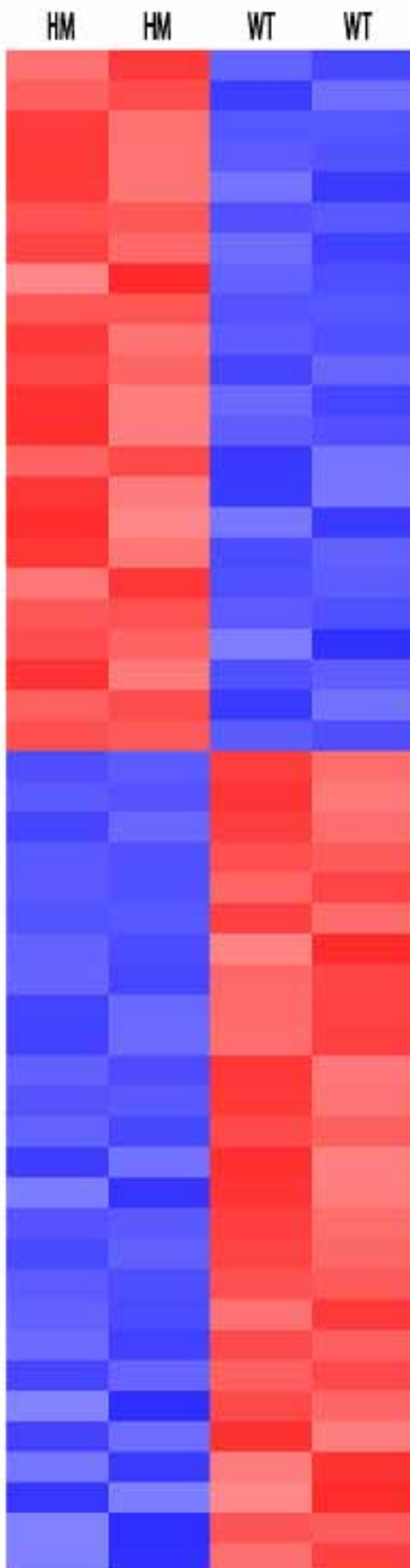
Syt1: synaptotagmin 1
 Ostα: organic solute transporter alpha
 Mm.173854.1
 Col1a1: collagen, type I, alpha 1
 4930539E08Rik: RIKEN cDNA 4930539E08 gene
 Elof6: ELOVL family member 6, elongation of long chain fatty acids [yeast]
 Slc1: secretory leukocyte peptidase inhibitor
 Indo: indoleamine-pyruvate 2,3 dioxygenase
 Elcelastin
 Homer2: homer homolog 2 (Drosophila)
 Genrb1: centaurin, beta 1
 Pygl: liver glycogen phosphorylase
 Sdc4: syndecan 4
 LOC100046081 // Coub1: OTU domain, ubiquitin aldehyde binding 1 // similar to OTU domain, ubiquitin aldehyde binding 1
 Mm.132077.1
 1700011H14Rik: RIKEN cDNA 1700011H14 gene
 Col5a2: collagen, type V, alpha 3
 Cnfs: camello-like 5
 Ifih3: interferon-induced protein with tetratricopeptide repeats 3
 Adams2: a disintegrin-like and metalloproteinase [reprolysin type] with thrombospondin type 1 motif, 2
 Slc5a9: solute carrier family 5 [sodium/glucone co-transporter], member 9
 Tcez2: transcription elongation factor A (SII) 2
 9430081123Rik: RIKEN cDNA 9430081123 gene
 Lce1i: late cornified envelope 1
 Rab8b: RAB8B, member RAS oncogene family
 Slc3a1: solute carrier family 3, member 1
 Pcolce2: procollagen C-endopeptidase enhancer 2
 Cxcl1: chemokine (C-X-C motif) ligand 1
 Trim40: tripartite motif-containing 40
 Stc2: stanniocalcin 2
 Tbc3: tetratricopeptide repeat domain 3
 1810030J14Rik // LOC100044832: RIKEN cDNA 1810030J14 gene // similar to RIKEN cDNA 1810030J14 gene
 Apoal: apolipoprotein A-I
 Mod1: MAX dimerization protein 1
 Transcribed locus
 Arid4a: AT-rich interactive domain 4A (Pbp1 like)
 2610019F08Rik: RIKEN cDNA 2610019F08 gene
 Mogat2: monoacylglycerol O-acyltransferase 2
 16 days neonate cerebellum: cDNA, RIKEN full-length enriched library
 Lbh // LOC100048380: limb-bud and heart // similar to limb-bud and heart
 Aldh1l2: aldehyde dehydrogenase 1 family, member 12
 Ace2: angiotensin I converting enzyme (peptidyl-dipeptidase A) 2
 Zymn5: zinc finger, MYM-type 5
 Ada: adenosine deaminase
 Cyp2d13: cytochrome P450, family 2D, subfamily 13, polypeptide 13
 2010305C02Rik: RIKEN cDNA 2010305C02 gene
 App1: amyloid beta (A4) precursor-like protein 1
 Ubcl1: ubiquitin-activating enzyme E1-like
 Trim12: tripartite motif protein 12
 Sh3bp4: SH3-domain binding protein 4



Klt1: kallikrein 1
Dpepl: dipeptidase 1 (renal)
Bc215: Bc2-like 15
Ghr: growth hormone receptor
Slc26a6: solute carrier family 26, member 6
Pex19: peroxisome biogenesis factor 19
Lgals1: lectin, galactose binding, soluble 1
Vip: vasoactive intestinal polypeptide
Hk3: hexokinase 3
Fbl: fibrin light chain 1
Aldh1a1: aldehyde dehydrogenase family 1, subfamily A7
1810007E14Rik: RIKEN cDNA 1810007E14 gene
Gm16: grameyne A
Zeb2: zinc finger E-box binding homeobox 2
Ntrk2: neurotrophic tyrosine kinase, receptor, type 2
AA409316: expressed sequence AA409316
1810010M01Rik: RIKEN cDNA 1810010M01 gene
Mpp1: membrane protein, palmitoylated
Lgals1: lectin, galactose binding, soluble 1
Tmem120a: transmembrane protein 120A
Gabarap: gamma-aminobutyric acid receptor associated protein
Cntf4: ciliary-like 4
Dmc: desmusulin
2310008M10Rik: RIKEN cDNA 2310008M10 gene
2610204M08Rik // LOC10047142: RIKEN cDNA 2610204M08 gene // similar to INF2
Mn.151642.1
Trim30: tripartite motif protein 30
Gadmc1: gademin C1
Fbox8: F-box and WD-40 domain protein 8
Mn.44320.1
Mn.121603.1
Adam1a // LOC10047083: a disintegrin and metalloproteinase domain 1a // similar to MK-5 type 2
I627: interferon, alpha-inducible protein 27
Slc9a3rl: solute carrier family 9 (sodium/hydrogen exchanger), isoform 3 regulator 1
B2m: beta-2 microglobulin
Cgn: cingulin
Ttc22: tetraakis peptide repeat domain 22
A1956758: expressed sequence A1956758
Klt1: kallikrein 1
Dnaj2: DnaJ (Hsp40) homolog, subfamily C, member 2
Tmem66: transmembrane protein 66
2010003K15Rik: RIKEN cDNA 2010003K15 gene
3110021A11Rik: RIKEN cDNA 3110021A11 gene
Acot1: acyl-Coenzyme A oxidase 1, palmitoyl
Wh3: wuchia related kinase 3
Cd5: chemokine (C-C motif) ligand 5
Srp14: signal recognition particle 14
Oas1a: 2'-5' oligoadenylate synthetase 1A
2310034G01Rik: RIKEN cDNA 2310034G01 gene
Ddaht1: dimethylarginine dimethylaminohydrolase 1

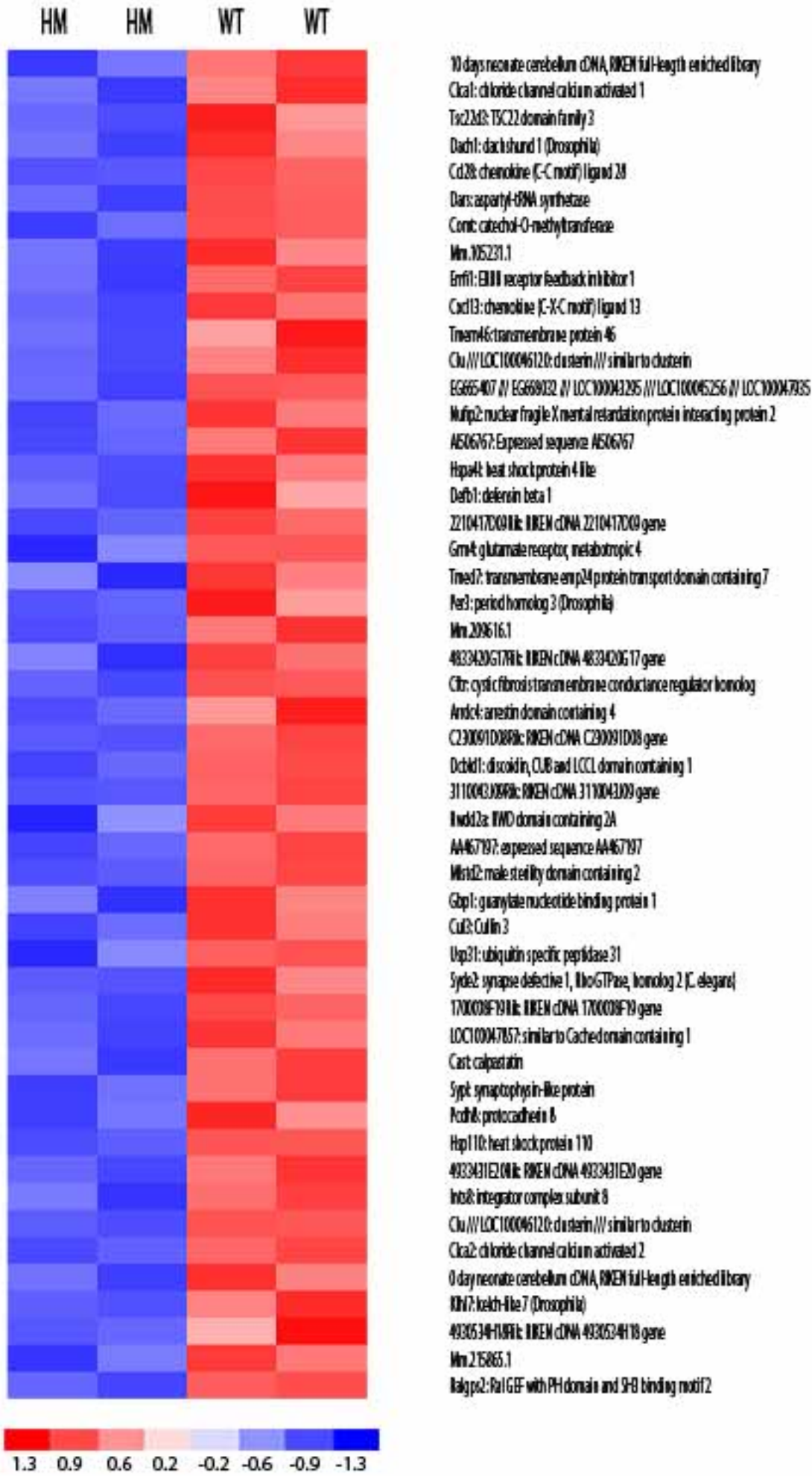


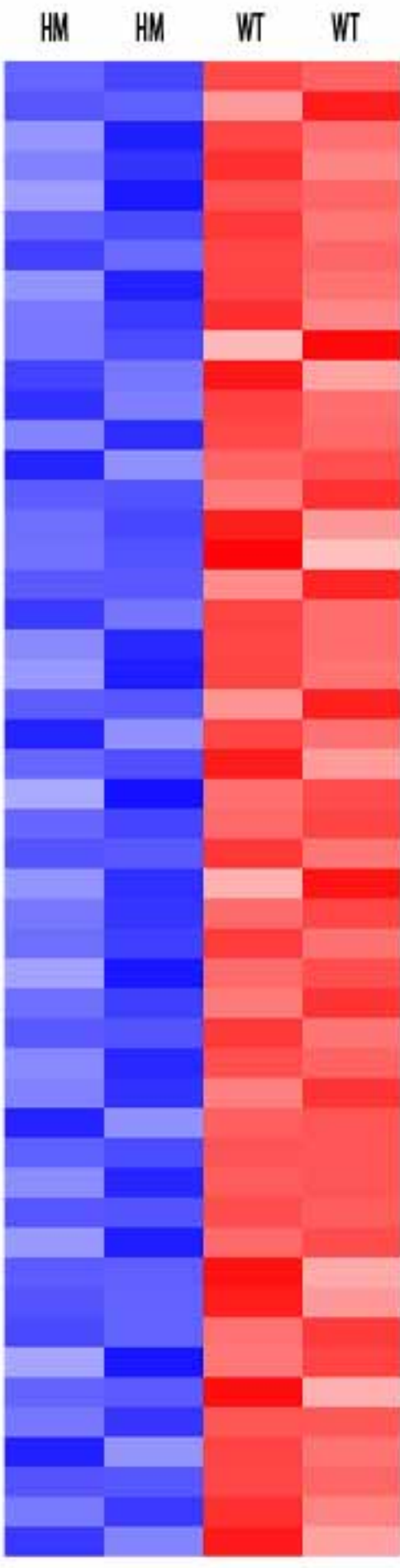
Nup62: nucleoporin 62
Prap1: proline-rich acidic protein 1
Dera: 2-deoxyribose-5-phosphate aldolase homolog [C. elegans]
Bckdhb ||| LOC10048676: branched chain ketoacid dehydrogenase E1, beta polypeptide
Col3a1: collagen, type III, alpha 1
G0s2: G0/G1 switch gene 2
Ap13: adaptor-related protein complex AP-1, sigma 3
Fgf13: fibroblast growth factor 13
Ostb: organic solute transporter beta
Gbp2: guanylate nucleotide binding protein 2
Lip1: lipin 1
Gbp2: guanylate nucleotide binding protein 2
Clqtn2: Clq and tumor necrosis factor related protein 2
Apol7a: apolipoprotein L 7a
Bcl3: B-cell leukemia/lymphoma 3
20106E10Rik: RIKEN cDNA 20106E10 gene
Wolly1: WD repeat and PWE domain containing 1
Figf: C-fos induced growth factor
Ldhb: lactate dehydrogenase B
Ctra6: chloride channel calcium activated 6
Transcribed locus
Rtp4: receptor transporter protein 4
Mgp: matrin Gla protein
Gtpbp2: GTP binding protein 2
Slc6a8t: solute carrier family 6 (neurotransmitter transporter, creatine), member 8
Spn2u: small proline-rich protein 2A
Cyp4f14: cytochrome P450, family 4, subfamily f, polypeptide 14
Rppi1p1: retinitis pigmentosa GTPase regulator interacting protein 1
Slc34a2: solute carrier family 34 (potassium phosphate), member 2
Fasn: fatty acid synthase
Trim34: tripartite motif protein 34
Pla2g12b: phospholipase A2, group XIb
Phd1b: pleckstrin homology-like domain, family 8, member 1
Figf: C-fos induced growth factor
Mttx: microsomal triglyceride transfer protein
Hsd3b3: hydroxysteroid 3 beta- and steroid delta-isomerase 3
4833409A17Rik: RIKEN cDNA 4833409A17 gene
Krt4: keratin 4
LOC100047419 ||| Maf: avian musculoaponeurotic fibrosarcoma (v-maf) A542 oncogene homolog ||| similar to c-Maf long form
Clec7a: C-type lectin domain family 7, member a
Rab15: RAB15, member of RAS oncogene family-like 5
Acot2: angiotensin I converting enzyme (peptidyl-dipeptidase A) 2
Art3: ADP-ribosyltransferase 3
1500011B03Rik: RIKEN cDNA 1500011B03 gene
Gpd1: glycerol-3-phosphate dehydrogenase 1 [soluble]
Transcribed locus; weakly similar to XP_001175112.1, similar to pol protein, partial [Pan troglodytes]
S100a10: S100 calcium binding protein A10 (calactin)
D12Ertd647e: DNA segment, Chr 12, ERATO Doi 647, expressed
Slc6a8t: solute carrier family 6 (neurotransmitter transporter, creatine), member 8
Mep1b: mepin 1 beta



1.3 0.9 0.6 0.2 -0.2 -0.6 -0.9 -1.3

Scl7a4: solute carrier family 7 (fattyacid transporter), member 4
Cldn5: chloride intracellular channel 5
Cdkn2b: cyclin-dependent kinase inhibitor 2B (p15, inhibits CDK4)
Rad23a: RAD23a homolog [S. cerevisiae]
Ardydig: Rho GDP dissociation inhibitor (GDI) gamma
Fbn1: fibrillin 1
Gmfg: glia maturation factor, gamma
Fbxl17: F-box and leucine-rich repeat protein 17
Alg8: asparagine-linked glycosylation 8 homolog (yeast, alpha-1,3-glucosyltransferase)
Canik1d: calcium/calmodulin-dependent protein kinase 1D
Sorbin1: Sorbin and SH3 domain containing 1
Nrp1: neuropilin 1
Trim31: tripartite motif-containing 31
Rab11t: RAB, member of RAS oncogene family-like 4
Tm4sf1: transmembrane 4 superfamily member 1
9130409J20Rik: RIKEN cDNA 9130409J20 gene
Cd59a: CD59a antigen
Fbxo17: F-box and WD-40 domain protein 17
Sifbd1: Si RNA binding domain 1
Visual cortex cDNA, RIKEN full-length enriched library
Inpkca: inositol 1,4,5-trisphosphate 3-kinase A
Atp1a2: ATPase, Na⁺/K⁺ transporting, alpha 2 polypeptide
Skl44a4: solute carrier family 44, member 4
Chordc1: cysteine and histidine-rich domain (CHORD)-containing, zinc-binding protein 1
Timed7: transmembrane emp24 protein transport domain containing 7
Nedd1t: neural precursor cell expressed, developmentally down-regulated gene 1
Dock10: dedicator of cytokinesis 10
Zfp326: zinc finger protein 326
Galactc2: chondroitin sulfate GalNAcT-2
Bace2: beta-site APP-cleaving enzyme 2
Dmd2: Dmd-like 2
Arf14: ADP-ribosylation factor-like 14
Hoxd3: homeobox D3
Fos: FBJ osteosarcoma oncogene
Zfp106: zinc finger protein 106
Aebp2: AE binding protein 2
Cd44: CD44 antigen
Gcg: glucagon
Tulp4: tulip-like protein 4
Mffr1: mitochondrial fission regulator 1
Bace2: beta-site APP-cleaving enzyme 2
Tubgcp5: tubulin, gamma complex associated protein 5
Ralgap2: Ral GEF with PH domain and SH3 binding motif 2
6430706D20Rik / *A73000H12Rik*: RIKEN cDNA A73000H12 gene / RIKEN cDNA 6430706D20 gene
Dnaj4: DnaJ (Hsp40) homolog, subfamily B, member 4
Znf41: zinc finger and BTB domain containing 41 homolog
Hir1: hippocampus abundant gene transcript 1
Tcfcp2l1: transcription factor CP2-like 1
Kif11: kinesin family member 11
D630004K10Rik: RIKEN cDNA D630004K10 gene

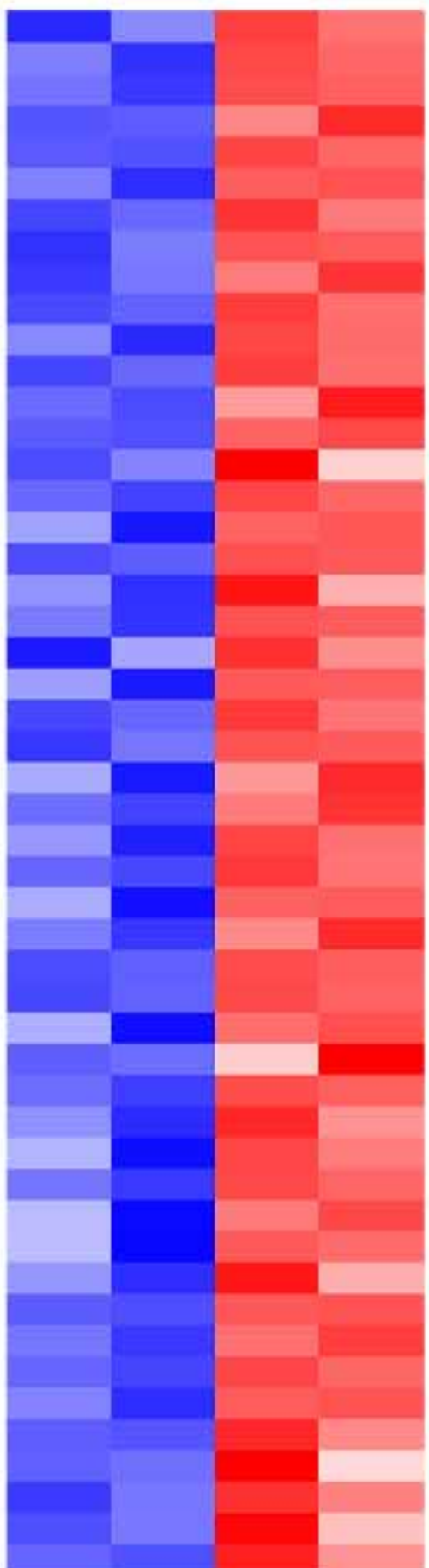




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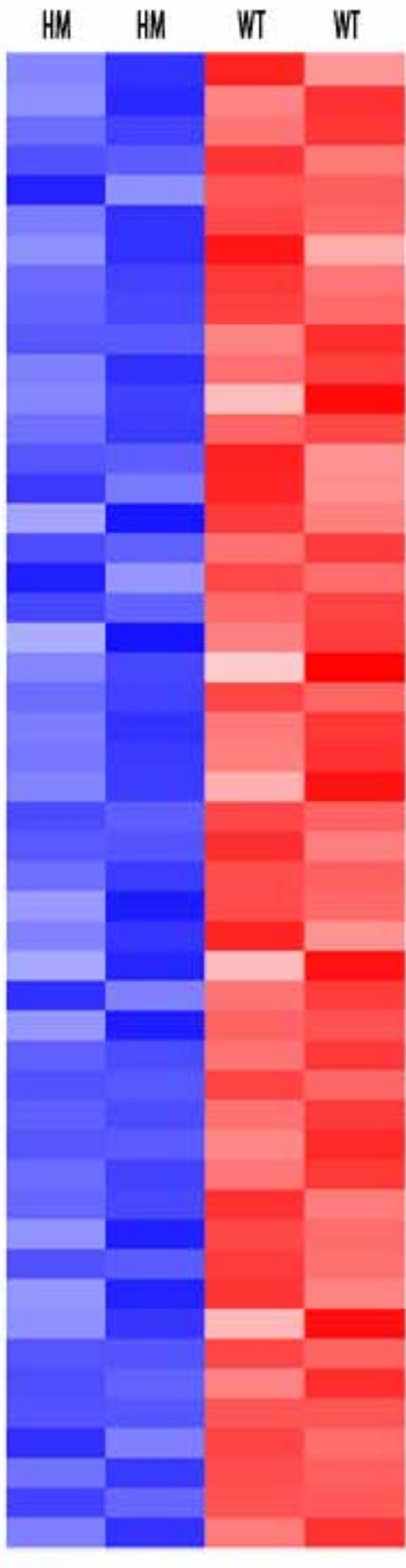
- Fgl1: FSHD region gene 1
 Alr1c14: aldo-keto reductase family 1, member C14
 Sctd4: SET domain containing 4
 6720463N24Rik: RIKEN cDNA 6720463N24 gene
 Sis12ip1: SF1512-interacting protein 1
 Slc5STE20-like kinase (yeast)
 Htr4: 5-hydroxytryptamine (serotonin) receptor 4
 46324110K06Rik: RIKEN cDNA 46324110K06 gene
 Defb1: defensin beta 1
 Cox15: COX15 homolog, cytochrome c oxidase assembly protein (yeast)
 Sh11: splicing factor, arginine/serine-rich 11
 Pbx1: pre-B-cell leukaemia transcription factor 1
 Ywhaz: tyrosine 3-monooxygenase tryptophan 5-monooxygenase activation protein, zeta polypeptide
 Ig-VI: immunoglobulin lambda chain, variable 1
 Jai1d1: jumonji AT-rich interactive domain 1D (Jbp2-like)
 LOC100430621//LOC100446651//Ywhaz:tyrosine 3-monooxygenase
 C33006P01Rik: RIKEN cDNA C33006P01Rik gene
 Aqp4: aquaporin 4
 Slc25a16: solute carrier family 25 (mitochondrial carrier, Grana disease autoantigen), member 16
 G15: Glyceraldehyde 3-phosphate dehydrogenase
 Tox3: TOX high mobility group box family member 3
 Rgn13: regulator of G-protein signalling 13
 Mm39929.1
 Mm84118.1
 Eno3: enolase 3, beta muscle
 AA467657: expressed sequence AA467657
 Sloc3a1: solute carrier organic anion transporter family, member 3a1
 Zfp518: zinc finger protein 518
 Gpr177: G protein-coupled receptor 177
 Sepintob5: serine (or cysteine) peptidase inhibitor, clade II, member 5
 1700110KT7Rik: RIKEN cDNA 1700110KT7 gene
 Zbtb16: zinc finger and BTB domain containing 16
 251004919Rik: RIKEN cDNA 251004919 gene
 Otu1//LOC100046120: clusterin// similar to clusterin
 Tubgcp4: tubulin, gamma complex associated protein 4
 Vav3: vav 3 oncogene
 Lin7c: lin-7 homolog C (*C.elegans*)
 2610021A01Rik: RIKEN cDNA 2610021A01 gene
 Pphn1: periphilin 1
 Bcl2: B-cell lymphoma/lymphoma 2
 4432414E09Rik: RIKEN cDNA 4432414E09 gene
 Transcribed locus
 Myb: myeloblastosis oncogene
 Csp1: centrosome and spindle pole associated protein 1
 2210019G11Rik: RIKEN cDNA 2210019G11 gene
 Oca2: chloride channel calcium activated 2
 Mstnd2: male sterility domain containing 2
 Dck1: doublecortin-like kinase 1
 D330012F22Rik: RIKEN cDNA D330012F22 gene
 Usp49: ubiquitin specific peptidase 49

HM HM WT WT



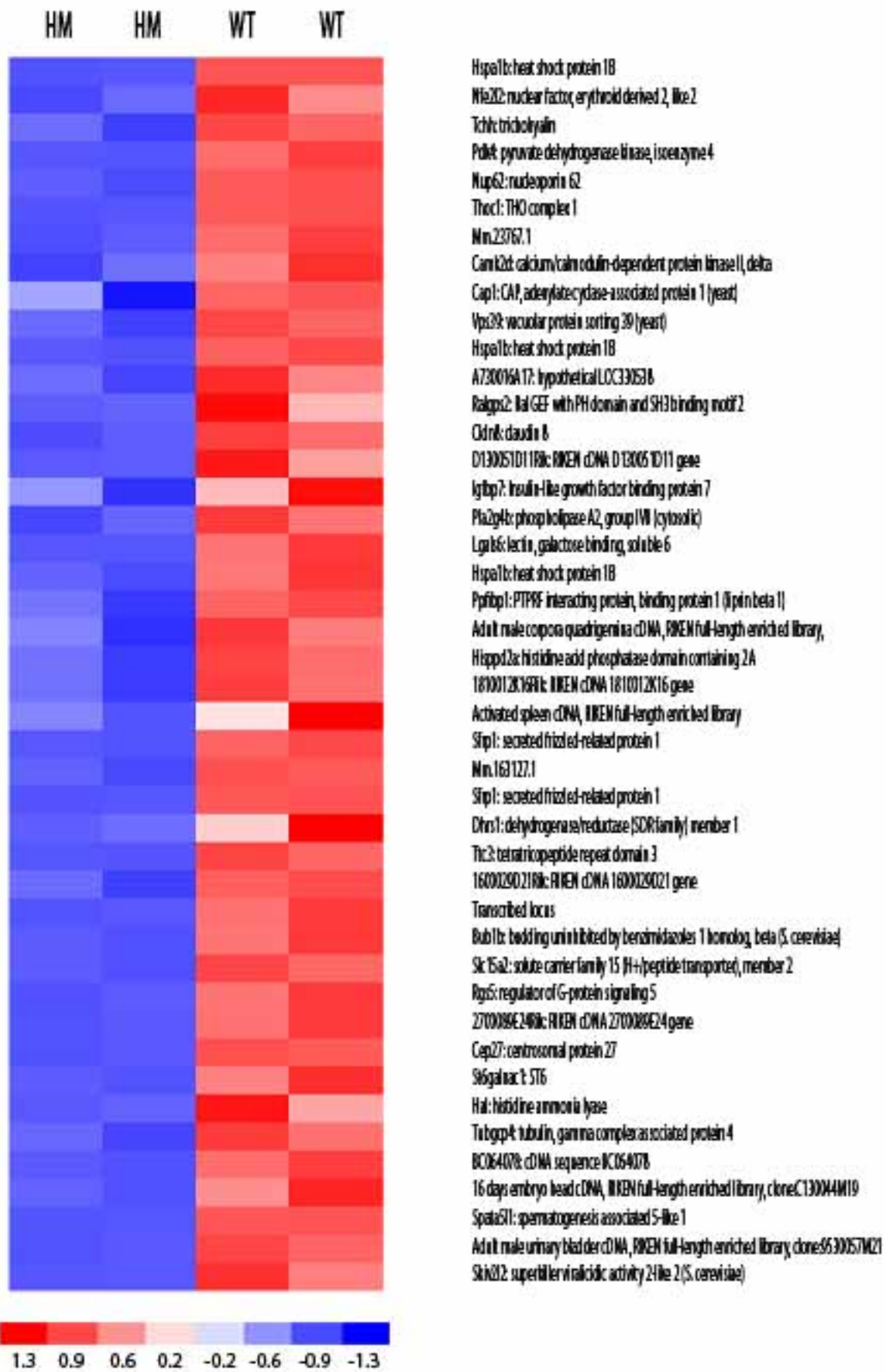
Sync: syncillin
 Nrn_1232.1
 B0X0267: cDNA sequence B0X0267
 Sult1d1: sulfotransferase family 1D, member 1
 Mt2: metallothionein 2
 Rnf180: ring finger protein 180
 Zdhhc14: zinc finger, DHHC domain containing 14
 Greb1 // LOC10045413: gene regulated by estrogen in breast cancer protein
 Miph: melanophilin
 Trt: transthyretin
 Hspa4l: heat shock protein 4 like
 Trt: transthyretin
 Pip1: PC4 and SRS1 interacting protein 1
 Cascan_10: CBA-related cell adhesion molecule 10
 Ky: kynurenic acid peptidase
 Fnkld3: FAS7 kinase domains 3
 Rhdk: P-R5 domain containing 1
 Cascan_10: CBA-related cell adhesion molecule 10
 5530402H23Rik: RIKEN cDNA 5530402H23 gene
 15 days embryo lead cDNA, RIKEN full-length enriched library
 Gtac2: glutathione S-transferase, C-terminal domain containing
 Lonrd: LON peptidase N-terminal domain and ring finger 3
 Zdhhc14: zinc finger, DHHC domain containing 14
 Sto2: storkhead box 2
 Slc6a14: solute carrier family 6 (neurotransmitter transporter), member 14
 Nrn_215886.1
 Acadc: arylacetamide deacetylase (esterase)
 Cox7c // LOC10041521 // LOC10047065 // LOC10048613: cytochrome c oxidase, subunit Vc
 493635809Rik: RIKEN cDNA 493635809 gene
 Cd177: CD177 antigen
 Nrip: nitric oxide synthase interacting protein
 LOC677143 // Shtn2: sidefelin 2 / similar to Sidefelin-2
 4839416E15Rik: ENPP1: ectonucleotide pyrophosphatase/phosphodiesterase 1
 Ptov1: prostate tumor over expressed gene 1
 Comt: catechol-O-methyltransferase
 Nrn_45081.1
 Snap23: synaptosomal-associated protein 23
 Ppargc1a: peroxisome proliferative activated receptor, gamma, coactivator 1 alpha
 4921509J17Rik: RIKEN cDNA 4921509J17 gene
 Atp13a2: ATPase type 13A3
 Erbg: estrogen-related receptor gamma
 A930005H10N: RIKEN cDNA A930005H10 gene
 Spre1: sprouty protein with BH-1 domain 1, related sequence
 Reg4: regenerating islet-derived family member 4
 A1024213: expressed sequence A1024213
 2810487A22Rik: RIKEN cDNA 2810487A22 gene
 A330021E22Rik: RIKEN cDNA A330021E22 gene
 Nsl1: Nasalith homolog 1 (Drosophila)
 Gdap2: ganglioside-induced differentiation-associated-protein 2
 Sult1c2: sulfotransferase family cytosolic 1C, member 2

1.3 0.9 0.6 0.2 -0.2 -0.6 -0.9 -1.3



Lig3: leucine-rich repeats and immunoglobulin-like domains 3
 Ythdc2: YTH domain containing 2
 Slc22a15: solute carrier family 22 (organic cation/cation transporter), member 15
 Syt1: serum/glucocorticoid regulated kinase 1
 Rnf187: ring finger protein 187
 LOC675567 //| Rcc1: RCC1 domain containing 1 //| similar to MYC binding protein 2
 Transcribed locus
 Mm.1232.1
 LOC633417: similar to histocompatibility 2, T-region locus 3
 Ppargc1a: Peroxisome proliferative activated receptor, gamma, coactivator 1 alpha
 Cdc32: coiled-coil domain containing 32
 Vcam1: vascular cell adhesion molecule 1
 LOC10044194 //| Ncc: mutated in colorectal cancers //| hypothetical protein LOC10044194
 Transcribed locus, moderately similar to XP_001117064.1 BTB (POZ) domain containing 9 [Macaca mulatta]
 2610204H10Rik: RIKEN cDNA 2610204H10 gene
 Hsp40: heat shock protein 40 kDa
 Dnajc4 //| LOC10048331: DnaJ (Hsp40) homolog, subfamily A, member 4
 Stox2: Storkhead box 2
 483420G17Rik: RIKEN cDNA 483420G17 gene
 P4ha1: procollagen-proline 2-oxoglutarate 4-oxygenase (proline 4-hydroxylase), alpha 1 polypeptide
 Hvd: isovaleryl coenzyme A dehydrogenase
 603042H21Rik: RIKEN cDNA 603042H21 gene
 Ppargc1a: Peroxisome proliferative activated receptor, gamma, coactivator 1 alpha
 Per1: period homolog 1 [Drosophila]
 D7End75e: cDNA segment, Chr7, BTA7. D7e, expressed
 EG633640: predicted gene EG633640
 5730437M04H1c: RIKEN cDNA 5730437M04 gene
 Trh: trithorax
 Oip5: Opa interacting protein 5
 Ankyrin repeat K
 LOC10046254: similar to Adl1 protein
 LOC10047788: similar to gamma-2a immunoglobulin heavy chain
 Mm.182879.1
 Bmp1: BMP1 homolog
 Igf6: immunoglobulin heavy chain 6 (heavy chain of IgM)
 D2hgdh: D-2-hydroxyglutarate dehydrogenase
 Neto2: neuropilin 1 (NRP) and toll-like (TL)-like 2
 Hspa1a: heat shock protein 1A
 16 days neonatal thymus cDNA, RIKEN full-length enriched library
 Satb2: special AT-rich sequence binding protein 2
 Cdc32c: coiled-coil domain containing 32C
 H3f3b: H3 histone, family 3B
 Usp53: ubiquitin specific peptidase 53
 051004010098c: RIKEN cDNA 051004009 gene
 5730601R069c: RIKEN cDNA 5730601R06 gene
 3110040N111c: RIKEN cDNA 3110040N11 gene
 Gip: gastric inhibitory polypeptide
 Trp53: Transformation related protein 53
 Hist2h2bb: histone cluster 2, H2bb
 Cd163l1: CD163 molecule-like 1

1.3 0.9 0.6 0.2 -0.2 -0.6 -0.9 -1.3



Supplement Figure 11. Heat map for mRNAs that are significantly enriched or diminished in Atg16L1^{HM} thymocytes relative to control.

A 1.3-fold difference and a 90% lower bound cut-off were used.

a

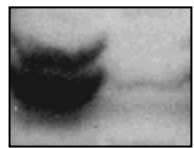
Term	RT	Genes	Count	%	P-Value	Fold Enrichment	Bonferroni	Benjamini	FDR
PPAR signaling pathway	RT		8	6.0	2.0E-6	12.4	3.9E-4	3.9E-4	0.0
lipid metabolic process	RT		16	12.0	6.7E-6	4.1	3.4E-2	3.4E-2	0.0
lipid transporter activity	RT		6	4.5	1.4E-5	19.1	3.8E-2	3.8E-2	0.0
extracellular space	RT		29	21.8	1.9E-5	2.3	1.5E-2	1.5E-2	0.0
cellular lipid metabolic process	RT		14	10.5	3.0E-5	4.1	1.4E-1	7.4E-2	0.1
extracellular region part	RT		29	21.8	5.8E-5	2.2	4.4E-2	2.2E-2	0.1
inflammatory response	RT		9	6.8	6.1E-5	6.6	2.7E-1	9.9E-2	0.1
signal	RT		32	24.1	9.4E-5	2.0	7.8E-2	7.8E-2	0.1
response to wounding	RT		10	7.5	1.1E-4	5.2	4.4E-1	1.4E-1	0.2
acute inflammatory response	RT		6	4.5	1.3E-4	12.2	4.8E-1	1.2E-1	0.2
lipid binding	RT		11	8.3	1.5E-4	4.5	3.3E-1	1.8E-1	0.3
carboxylic acid metabolic process	RT		12	9.0	2.0E-4	3.9	6.4E-1	1.6E-1	0.4
organic acid metabolic process	RT		12	9.0	2.0E-4	3.9	6.5E-1	1.4E-1	0.4
regulation of cholesterol absorption	RT		3	2.3	2.1E-4	124.8	6.6E-1	1.3E-1	0.4
lipid transport	RT		6	4.5	2.2E-4	10.9	6.8E-1	1.2E-1	0.4
extracellular region	RT		29	21.8	2.3E-4	2.0	1.7E-1	5.9E-2	0.4
alcohol metabolic process	RT		9	6.8	2.7E-4	5.3	7.5E-1	1.3E-1	0.5
cholesterol absorption	RT		3	2.3	3.5E-4	99.8	8.3E-1	1.5E-1	0.7
Secreted	RT		19	14.3	3.5E-4	2.5	2.6E-1	1.4E-1	0.5
monocarboxylic acid metabolic process	RT		8	6.0	4.1E-4	5.8	8.8E-1	1.6E-1	0.8
signal peptide	RT		29	21.8	4.3E-4	1.9	1.0E0	1.0E0	0.9
lipid digestion	RT		3	2.3	5.2E-4	83.2	9.3E-1	1.9E-1	1.0
acute-phase response	RT		4	3.0	5.4E-4	24.7	9.4E-1	1.8E-1	1.0
lipoprotein	RT		12	9.0	6.1E-4	3.5	4.1E-1	1.6E-1	0.9
response to external stimulus	RT		11	8.3	7.1E-4	3.7	9.7E-1	2.2E-1	1.3

b

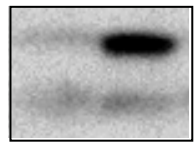
Term	RT	Genes	Count	%	P-Value	Fold Enrichment	Bonferroni	Benjamini	FDR
ubiquitin conjugation	RT		7	7.5	1.1E-3	5.9	6.1E-1	6.1E-1	1.7
amino acid and derivative metabolic process	RT		6	6.5	5.1E-3	5.3	1.0E0	1.0E0	9.3
cytoplasm	RT		30	32.3	5.3E-3	1.5	9.8E-1	9.8E-1	7.8
regulation of catalytic activity	RT		6	6.5	7.4E-3	4.8	1.0E0	1.0E0	13.2
apoptosis	RT		8	8.6	8.3E-3	3.4	1.0E0	1.0E0	14.7
programmed cell death	RT		8	8.6	9.1E-3	3.3	1.0E0	1.0E0	16.0
alternative splicing	RT		21	22.6	9.8E-3	1.7	1.0E0	9.9E-1	14.2
amine metabolic process	RT		6	6.5	1.0E-2	4.5	1.0E0	1.0E0	17.6
chaperone	RT		4	4.3	1.0E-2	8.8	1.0E0	9.5E-1	14.6
metabolic process	RT		37	39.8	1.0E-2	1.3	1.0E0	1.0E0	17.7
cell death	RT		8	8.6	1.1E-2	3.2	1.0E0	1.0E0	19.0
death	RT		8	8.6	1.1E-2	3.2	1.0E0	1.0E0	19.2
regulation of a molecular function	RT		6	6.5	1.3E-2	4.2	1.0E0	1.0E0	22.4
nitrogen compound metabolic process	RT		6	6.5	1.3E-2	4.2	1.0E0	1.0E0	22.4

Supplementary Figure 12. Functional analysis of enriched and diminished transcripts of Atg16L1-deficient Paneth cells. a-b, The Functional Classification Tool from the DAVID Bioinformatics Resource² (NIAID, NIH) was used to determine the enriched functional categories of the two gene lists in Supplementary Table 1 comparing the transcriptome of Atg16L1^{HM} versus WT Paneth cells (enriched (**a**) or diminished (**b**) in Atg16L1^{HM} mice). In both cases, the highest statistically significant gene ontology (GO) terms that were represented within the gene list are indicated. We used medium stringency to cluster the genes. Count = number of genes in the list with a given GO term; % = fractional representation of a GO term within the list. Various statistical methods are available to determine significance (p-value, fold-enrichment, Bonferroni, Benjamini, and false-discovery rate, FDR). Functional groups were sorted by p-value in these tables. The statistical significance of the clustering for the top rated GO terms is much higher in the gene list for the enriched transcripts.

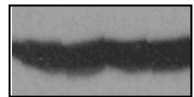
WT HM1



Atg16L1



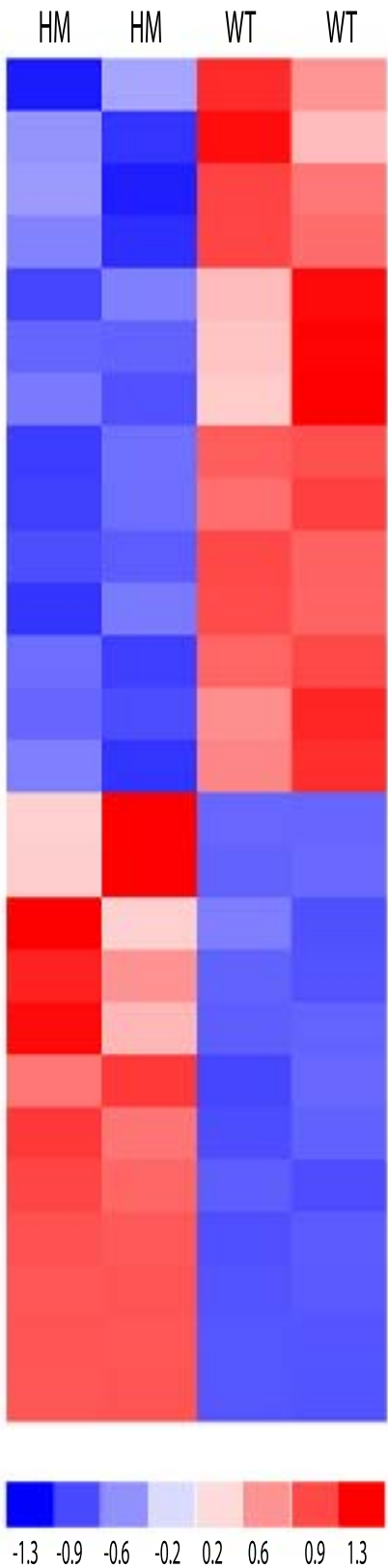
LC3



Actin

Actin

Supplementary Figure 13. Thymocytes from Atg16L1^{HM} mice have reduced Atg16L1 expression and conversion of LC3-I to LC3-II. Western blot analysis of whole cell thymocyte lysates from Atg16L1^{HM} mice reveal decreased Atg16L1 expression and an increase in LC3-I to LC3-II ratio compared to WT control.



Supplement Figure 14. Heat map for mRNAs that are significantly enriched or diminished in Atg16L1^{HM} Paneth cells relative to control.

A 1.3-fold difference and a 90% lower bound cut-off were used.

Type of file: table

Label: 3

Filename: Virgin_Cadwell et al - Supp Info 3_Table.xls

Transcripts decreased in Atg16L1 deficient Paneth cells

probe set	gene	Accession
1457666_s_at	interferon activated gene 202B	AV229143
1421551_s_at	interferon activated gene 202B	NM_011940
1447517_at	superkiller viralicidic activity 2-like 2 (<i>S. cerevisiae</i>)	BM208991
1442632_at	Centaурин, gamma 2	BB116896
1455863_at	spermatogenesis associated 5-like 1	AW215442
1442424_at	CTD (carboxy-terminal domain, RNA polymerase II, polypeptide A) sma	BB368771
1435069_at	cDNA sequence BC064078	AW493518
1429345_at	DNA segment, Chr 2, ERATO Doi 435, expressed	AK016563
1418645_at	histidine ammonia lyase /// similar to Histidine ammonia-lyase (Histidas)	L07645
1421517_at	ST6 (alpha-N-acetyl-neuraminyl-2,3-beta-galactosyl-1,3)-N-acetylgalact	NM_011371
1430577_at	centrosomal protein 27	BE686390
1453208_at	RIKEN cDNA 2700089E24 gene	AV030118
1420941_at	regulator of G-protein signaling 5	BF585144
1417600_at	solute carrier family 15 (H ⁺ /peptide transporter), member 2	NM_021301
1441460_at	FGFR1 oncogene partner 2	BB435465
1447362_at	budding uninhibited by benzimidazoles 1 homolog, beta (<i>S. cerevisiae</i>)	AU045529
1423933_a_at	RIKEN cDNA 1600029D21 gene	BC022950
1441684_at	tetratricopeptide repeat domain 3	BM116591
1415677_at	dehydrogenase/reductase (SDR family) member 1	NM_026819
1448395_at	secreted frizzled-related sequence protein 1	BI658627
1457417_at	gb:BB307346 /DB_XREF=gi:16402288 /DB_XREF=BB307346 /CLONE	BB307346
1438620_x_at	secreted frizzled-related sequence protein 1	BB497685
1442109_at	gb:BE989344 /DB_XREF=gi:10666631 /DB_XREF=UI-M-BZ1-bfr-o-05-	BE989344
1447595_x_at	RIKEN cDNA 1810012K16 gene	AV050083
1442466_a_at	histidine acid phosphatase domain containing 2A	BG071019
1457443_at	gb:BB311369 /DB_XREF=gi:16402696 /DB_XREF=BB311369 /CLONE	BB311369
1443029_at	PTPRF interacting protein, binding protein 1 (liprin beta 1)	AW550306
1422071_at	lectin, galactose binding, soluble 6	AF026794
1427126_at	heat shock protein 1B	M12573
1425045_at	phospholipase A2, group IVB (cytosolic)	BC016255
1437804_at	Insulin-like growth factor binding protein 7	BE446893
1439839_at	RIKEN cDNA D130051D11 gene	BB815394
1449091_at	claudin 8	BC003868
1431704_a_at	Ral GEF with PH domain and SH3 binding motif 2	AK018622
1446731_at	hypothetical protein A730016A17	BB246786
1427127_x_at	heat shock protein 1B	M12573
1451720_at	vacuolar protein sorting 39 (yeast)	BC007479
1417461_at	CAP, adenylate cyclase-associated protein 1 (yeast)	NM_007598
1439168_at	calcium/calmodulin-dependent protein kinase II, delta	BB373572
1448150_at	nucleoporin 62	NM_053074
1431044_at	THO complex 1	BB647938
1444056_at	alanine-glyoxylate aminotransferase 2-like 2	BB553796
1417273_at	pyruvate dehydrogenase kinase, isoenzyme 4	NM_013743
1434425_at	trichohyalin	AI597080
1457117_at	nuclear factor, erythroid derived 2, like 2	AV248273
1452318_a_at	heat shock protein 1B	M12573
1440808_x_at	RIKEN cDNA E430002D04 gene	BB209710
1425868_at	Histone cluster 2, H2bb	BC019122
1438542_at	gb:BB199080 /DB_XREF=gi:16271258 /DB_XREF=BB199080 /CLONE	BB199080

1449908_at	gastric inhibitory polypeptide	NM_008119
1450972_at	RIKEN cDNA 3110040N11 gene	AK019261
1447886_at	RIKEN cDNA 0610040B09 gene	AV302770
1457913_at	RIKEN cDNA 5730601F06 gene	BB374879
1452385_at	ubiquitin specific peptidase 53	BG076275
1430357_at	H3 histone, family 3B	BM241237
1453330_at	RIKEN cDNA 0610010D24 gene	AK002458
1427017_at	special AT-rich sequence binding protein 2	BB104560
1452388_at	heat shock protein 1A	AW763765
1458089_at	FK506 binding protein 5	BB161337
1436309_at	neuropilin (NRP) and tolloid (TLL)-like 2	BB125651
1427351_s_at	immunoglobulin heavy chain 6 (heavy chain of IgM)	BB226392
1437840_s_at	similar to CG3835-PA, isoform A	BB222646
1457728_at	BCNP1 homolog	BB763642
1425385_a_at	Immunoglobulin heavy chain 6 (heavy chain of IgM)	BC018365
1445963_at	Phosphodiesterase 5A, cGMP-specific	BG076280
1449076_x_at	acireductone dioxygenase 1	NM_134052
1453109_at	arylsulfatase K	AK013194
1430617_at	Opa interacting protein 5	BB238604
1426607_at	predicted gene, EG633640	BG068672
1453591_at	RIKEN cDNA 5730437N04 gene	AK017626
1459737_s_at	transthyretin	AA408768
1455087_at	DNA segment, Chr 7, ERATO Doi 715, expressed	AV328498
1449851_at	period homolog 1 (Drosophila)	AF022992
1434099_at	Peroxisome proliferative activated receptor, gamma, coactivator 1 alpha	BB752393
1441799_at	RIKEN cDNA 6030422H21 gene	AI098139
1449001_at	isovaleryl coenzyme A dehydrogenase	AV221988
1419636_at	RIKEN cDNA 4833420G17 gene	BG067119
1452094_at	procollagen-proline, 2-oxoglutarate 4-dioxygenase (proline 4-hydroxylase)	AI314028
1441953_at	Storkhead box 2	BB003572
1418591_at	DnaJ (Hsp40) homolog, subfamily A, member 4	NM_021422
1428966_at	RIKEN cDNA 2610204K14 gene	AK018614
1458385_at	heat shock protein 4 like	AI461691
1448162_at	vascular cell adhesion molecule 1	BB250384
1438081_at	mutated in colorectal cancers	BB794635
1458719_at	Transcribed locus	BM233846
1425470_at	gb:BC003855.1 /DB_XREF=gi:13277980 /FEA=FLmRNA /CNT=238 /T	BC003855
1427726_at	similar to histocompatibility 2, T region locus 3	X03051
1434100_x_at	Peroxisome proliferative activated receptor, gamma, coactivator 1 alpha	BB752393
1460558_at	coiled-coil domain containing 32	AV295080
1416041_at	serum/glucocorticoid regulated kinase	NM_011361
1451157_at	ring finger protein 187	AW488376
1456411_at	RCC1 domain containing 1	BM218704
1445671_at	Transcribed locus	BG228218
1444312_at	solute carrier family 22 (organic anion/cation transporter), member 15	BB017021
1430554_at	leucine-rich repeats and immunoglobulin-like domains 3	BB524113
1437357_at	YTH domain containing 2	AI481820
1424057_at	ganglioside-induced differentiation-associated-protein 2	BC025070
1449409_at	sulfotransferase family, cytosolic, 1C, member 2	NM_026935

Transcripts increased in Atg16L1 deficient Paneth cells

probe set	gene	Accession
1436936_s_at	inactive X specific transcripts	BG806300
1427262_at	inactive X specific transcripts	L04961
1424454_at	transmembrane protein 87A	BC027354
1424729_at	cDNA sequence BC054059	AY092026
1415965_at	stearoyl-Coenzyme A desaturase 1	NM_009127
1452260_at	cell death-inducing DFFA-like effector c	BB221402
1415964_at	stearoyl-Coenzyme A desaturase 1	NM_009127
1431056_a_at	lipoprotein lipase /// similar to Lipoprotein lipase precursor (LPL)	AK017272
1460256_at	carbonic anhydrase 3	NM_007606
1436453_at	expressed sequence BB144871	BB144871
1424737_at	thyroid hormone responsive SPOT14 homolog (Rattus)	BC009165
1457088_at	pallidin	BB780781
1415994_at	cytochrome P450, family 2, subfamily e, polypeptide 1	NM_021282
1417023_a_at	fatty acid binding protein 4, adipocyte	NM_024406
1417556_at	fatty acid binding protein 1, liver	NM_017399
1422651_at	adiponectin, C1Q and collagen domain containing	NM_009605
1437755_at	solute carrier family 5 (sodium/glucose cotransporter), member 12	BB503566
1448764_a_at	fatty acid binding protein 1, liver	NM_017399
1448881_at	haptoglobin	NM_017370
1419075_s_at	serum amyloid A 1	NM_011314
1450788_at	serum amyloid A 1	NM_009117
1422582_at	leptin	U18812
1417867_at	complement factor D (adipsin)	NM_013459
1436521_at	solute carrier family 36 (proton/amino acid symporter), member 2	AI596194
1416383_a_at	pyruvate carboxylase	NM_008797
1449015_at	resistin like alpha	NM_020509
1424758_s_at	serine (or cysteine) peptidase inhibitor, clade A (alpha-1 antiproteinase, BC018416	
1434959_at	desert hedgehog	AV367068
1436504_x_at	apolipoprotein A-IV	AV027367
1418724_at	complement component factor i	NM_007686
1446585_at	gb:BB263909 /DB_XREF=gi:16400167 /DB_XREF=BB263909 /CLONE BB263909	
1418595_at	plasma membrane associated protein, S3-12	NM_020568
1419231_s_at	RIKEN cDNA A830036E02 gene	NM_010661
1417761_at	apolipoprotein A-IV	BC010769
1456123_at	expressed sequence AI481121	BB498095
1457976_at	RIKEN cDNA 2010002M12 gene	AI447935
1424451_at	acetyl-Coenzyme A acyltransferase 1B	BC019882
1450883_a_at	CD36 antigen	BB534670
1431725_at	formin 2	AK013585
1455899_x_at	suppressor of cytokine signaling 3	BB241535
1449453_at	bone marrow stromal cell antigen 1	AI647987
1417184_s_at	hemoglobin, beta adult major chain /// hemoglobin, beta adult minor ch: BC027434	
1456212_x_at	suppressor of cytokine signaling 3	BB831725
1418995_at	gb:NM_010895.1 /DB_XREF=gi:6754823 /GEN=Neurod2 /FEA=FLmRINM_010895	
1458276_x_at	citron	BB730767
1420987_at	kinesin family member 3B	BB448784
1424976_at	ras homolog gene family, member V	BC021307
1426225_at	retinol binding protein 4, plasma	U63146
1417764_at	signal sequence receptor, alpha /// similar to signal sequence receptor,	BG077348
1415904_at	lipoprotein lipase	BC003305
1451916_s_at	tripartite motif protein 15	BC027186

1446042_at	gb:BG068241 /DB_XREF=gi:12550810 /DB_XREF=H3063C07-3 /CLO BG068241
1459802_at	PDZ domain containing RING finger 4
1451382_at	ChaC, cation transport regulator-like 1 (E. coli)
1417765_a_at	amylase 1, salivary
1435184_at	natriuretic peptide receptor 3
1428361_x_at	hemoglobin alpha, adult chain 1
1418804_at	succinate receptor 1
1426633_s_at	potassium channel tetramerisation domain containing 14
1445191_at	Vexonuclease 3'-5' domain-like 1
1419060_at	granzyme B
1447944_at	zinc finger with KRAB and SCAN domains 1
1459848_x_at	dopamine beta hydroxylase
1419975_at	gb:C76618 /DB_XREF=gi:2516948 /DB_XREF=C76618 /CLONE=J001 C76618
1434129_s_at	lipoma HMGIC fusion partner-like 2
1456444_at	F-box protein 41
1460084_at	gb:BB473929 /DB_XREF=gi:16439785 /DB_XREF=BB473929 /CLONE BB473929
1442213_at	hypothetical LOC552908
1419400_at	microsomal triglyceride transfer protein
1437904_at	developmentally regulated RNA binding protein 1
1444679_at	PHD finger protein 21A
1425318_a_at	transmembrane protein 116
1447478_at	RAN, member RAS oncogene family
1449147_at	carbohydrate (keratan sulfate Gal-6) sulfotransferase 1
1449434_at	carbonic anhydrase 3
1453108_at	arylsulfatase K
1434051_s_at	heat shock protein 12A
1455464_x_at	gb:BB427704 /DB_XREF=gi:16423841 /DB_XREF=BB427704 /CLONE BB427704
1428811_at	WAP, follistatin/kazal, immunoglobulin, kunitz and netrin domain containing 1
1431077_at	alkaline phosphatase, intestinal
1431701_a_at	PDZ domain containing 1
1440342_at	RIKEN cDNA G530011O06 gene
1439838_a_at	transmembrane inner ear
1452757_s_at	hemoglobin alpha, adult chain 1
1416307_at	adaptor-related protein complex AP-1, mu subunit 1
1418697_at	indolethylamine N-methyltransferase
1445578_at	ELOVL family member 6, elongation of long chain fatty acids (yeast)
1449569_at	thrombopoietin
1448485_at	gamma-glutamyltransferase 1
1454883_at	expressed sequence AI987692
1422678_at	diacylglycerol O-acyltransferase 2
1430641_at	RIKEN cDNA 9030605I04 gene /// expressed sequence AI987692
1424749_at	WD repeat and FYVE domain containing 1
1425546_a_at	transferrin
1424186_at	coiled-coil domain containing 80
1423166_at	CD36 antigen
1436058_at	radical S-adenosyl methionine domain containing 2
1452975_at	alanine-glyoxylate aminotransferase 2-like 1
1419232_a_at	apolipoprotein A-I
1440682_at	Odd Oz/ten-m homolog 4 (Drosophila)
1431806_at	RIKEN cDNA 4931408D14 gene
1460300_a_at	leukocyte tyrosine kinase
1431240_at	C-type lectin domain family 2, member h

1455201_x_at	apolipoprotein A-I	AI194999
1427007_at	RIKEN cDNA 1200013B08 gene	AK004734
1438840_x_at	apolipoprotein A-I	AI527359
1455568_at	RIKEN cDNA 2310015A05 gene	BI689456
1459144_at	Fibronectin type III domain containing 3a	BB518323
1438514_at	NPC1-like 1	AV377091
1450783_at	interferon-induced protein with tetratricopeptide repeats 1	NM_008331
1452095_a_at	histocompatibility 47	AK005204
1416978_at	Fc receptor, IgG, alpha chain transporter	NM_010189
1431591_s_at	hypothetical protein LOC677168	AK019325
1425122_at	open reading frame 9	AF360358
1447043_at	V-erb-a erythroblastic leukemia viral oncogene homolog 4 (avian)	BB386653
1431213_a_at	RIKEN cDNA 1300007C21 gene /// predicted gene, EG668462 /// predicted gene	BG297038
1447845_s_at	vanin 1	AV360029
1423556_at	aldo-keto reductase family 1, member B7	AV021656
1435529_at	similar to interferon-induced protein with tetratricopeptide repeats 1 /// s	BM245961
1418654_at	hydroxyacid oxidase (glycolate oxidase) 3	NM_019545
1418069_at	apolipoprotein C-II	NM_009695
1447100_s_at	RIKEN cDNA 5730508B09 gene	C80506
1451263_a_at	fatty acid binding protein 4, adipocyte	BC002148
1429089_s_at	RIKEN cDNA 2900026A02 gene	BG063749
1450344_a_at	prostaglandin E receptor 3 (subtype EP3)	NM_011196
1428780_at	threonine aldolase 1	AK005036
1455582_at	Transcribed locus	BI156044
1429667_at	POU domain, class 4, transcription factor 1	BB465134
1453070_at	protocadherin 17	BB305930
1427910_at	cystatin E/M	AK003744
1419144_at	CD163 antigen	NM_053094
1443726_at	SET and MYND domain containing 1	AV010327
1441442_at	gb:BM232203 /DB_XREF=gi:17867473 /DB_XREF=K0319H04-3 /CLO	BM232203
1431025_at	fuzzy homolog (Drosophila)	BB627268
1454746_at	pleckstrin homology domain containing, family M (with RUN domain) member 1	BB528702
1454021_a_at	exosome component 10	AK017621
1457644_s_at	chemokine (C-X-C motif) ligand 1	BB554288

EntrezGene	baseline mean	experiment mean	fold change	lower bound of FC
26388	505.02	1.39	-362.44	-127.07
26388	171.07	2.46	-69.58	-26.11
72198	265.81	5.91	-45	-24.01
347722	771.05	17.45	-44.2	-26.07
214616	720.85	34.22	-21.06	-17.45
329506	146.47	8.55	-17.12	-7.85
408064	207.27	13.39	-15.48	-7.01
51885	181.34	12.34	-14.69	-6.32
15109 /// 638196	127.12	8.92	-14.25	-6.47
20445	213.82	16.67	-12.83	-8.9
66296	399.32	35.5	-11.25	-8.9
381820	866.69	87.48	-9.91	-4.33
19737	1234.73	135.9	-9.09	-7.05
57738	1673.19	222.33	-7.53	-6.13
67529	154.52	21.07	-7.33	-5.05
12236	170.15	23.21	-7.33	-4.38
76509	196.78	30.02	-6.56	-3.89
22129	720.82	112.67	-6.4	-5.24
52585	294.89	46.51	-6.34	-2.51
20377	277.81	48.12	-5.77	-4.36
	183.2	31.93	-5.74	-4.37
20377	1303.18	263.6	-4.94	-4.34
	285.75	59.96	-4.77	-2.04
69108	242.78	51.42	-4.72	-3.04
327655	230.16	58.5	-3.93	-2.63
	149.67	38.42	-3.9	-2.3
67533	177.54	47.42	-3.74	-2.57
16857	3045.05	819.7	-3.71	-3.27
15511	1181.66	318.36	-3.71	-3.01
211429	555.06	152.85	-3.63	-2.89
29817	1147	319.54	-3.59	-1.44
319737	261.47	73.71	-3.55	-2.57
54420	218.54	63.77	-3.43	-2.85
78255	258.52	78.47	-3.29	-2.22
330538	219.87	67.41	-3.26	-2.41
15511	2603.52	821.2	-3.17	-2.95
269338	1627.86	515.29	-3.16	-2.42
12331	179.13	57.05	-3.14	-1.73
108058	222.03	71.32	-3.11	-2.24
18226	324.93	105.13	-3.09	-2.77
225160	557.03	180.04	-3.09	-2.86
72947	248.51	80.49	-3.09	-2.31
27273	504.03	165.97	-3.04	-2.66
99681	153.35	50.82	-3.02	-2.37
18024	153.83	51.45	-2.99	-2.32
15511	4231.36	1440.56	-2.94	-2.8
244233	153.9	52.56	-2.93	-2.06
319189	355.16	121.86	-2.91	-2.45
	345.74	124.13	-2.79	-2.19

14607	164.73	60.17	-2.74	-1.96
67290	1616.37	597.8	-2.7	-2.47
75395	470.44	176.29	-2.67	-2.43
77519	343.09	128.72	-2.67	-2.19
99526	262.34	98.95	-2.65	-1.56
15081	284.07	108.52	-2.62	-1.71
68339	172.35	67.25	-2.56	-2.21
212712	411.77	162.04	-2.54	-1.73
193740	743.89	297.47	-2.5	-2.08
14229	206.77	82.68	-2.5	-2.06
74513	191.15	76.85	-2.49	-2.08
16019	1796.37	723.88	-2.48	-2.29
673662	1139.83	459.77	-2.48	-2.08
66171	324.93	135.1	-2.41	-2.08
16019	417.04	175.7	-2.37	-1.75
242202	274.34	115.77	-2.37	-1.64
104923	317.61	136.89	-2.32	-1.43
77041	378.09	164.31	-2.3	-1.67
70645	180.06	78.52	-2.29	-1.6
633640	433.55	190.06	-2.28	-2.09
70544	216.4	94.89	-2.28	-1.86
22139	724.25	317.08	-2.28	-1.92
52480	214.57	96.94	-2.21	-1.57
18626	202.12	91.85	-2.2	-1.77
19017	348.35	160.78	-2.17	-1.71
402765	740.48	342.73	-2.16	-1.87
56357	684.63	320.52	-2.14	-1.43
67392	300.75	141.96	-2.12	-1.89
18451	252.13	118.95	-2.12	-1.43
71069	321.06	152.49	-2.11	-1.57
58233	244.56	116.22	-2.1	-1.78
67148	1599.78	767.33	-2.08	-1.67
18415	198.41	95.27	-2.08	-1.43
22329	316.41	153.12	-2.07	-1.47
328949	259.9	125.47	-2.07	-1.77
	200.02	96.76	-2.07	-1.62
	439.18	213.46	-2.06	-1.75
633417	331.62	160.89	-2.06	-1.82
19017	336.99	163.67	-2.06	-1.77
269336	390.9	189.59	-2.06	-1.52
20393	3246.21	1580.41	-2.05	-1.82
108660	1866.18	912.3	-2.05	-1.59
269955	435.18	212.44	-2.05	-1.68
	365.95	178.43	-2.05	-1.45
242126	227.88	111.64	-2.04	-1.61
320398	228.2	113.48	-2.01	-1.52
240255	281.39	139.93	-2.01	-1.52
14547	1047.9	524.38	-2	-1.44
69083	259.91	129.93	-2	-1.65

EntrezGene	baseline mean	experiment mean	fold change	lower bound of FC
213742	21.8	4461.91	204.7	97.46
213742	3.23	256.69	79.47	34.42
211499	7.03	325.08	46.22	20.64
246747	12.8	156.54	12.23	5.31
20249	74.3	739.14	9.95	7.57
14311	27.04	245.88	9.09	5.07
20249	681.8	5922.21	8.69	7.39
16956 /// 669888	31.95	262.57	8.22	6.27
12350	68.94	556.61	8.07	6.5
101918	97.18	708.65	7.29	5.9
21835	420.49	2793.08	6.64	5.5
18457	189.91	1260.25	6.64	5.26
13106	310.35	2054.29	6.62	5.71
11770	120.06	791.45	6.59	5.12
14080	62.14	403.37	6.49	4.04
11450	105.35	677.69	6.43	5.72
241612	27.28	170.95	6.27	3.51
14080	152.66	948.46	6.21	5.35
15439	130.25	758.36	5.82	5.26
20208	130.03	738.27	5.68	4.02
20208	289.02	1603.65	5.55	4.63
16846	46.27	256.12	5.54	4.19
11537	86.95	446.48	5.13	3.8
246049	112.12	573.51	5.11	3.69
18563	96.91	473.56	4.89	3.19
57262	27.91	132.96	4.76	3.14
217847	140.3	634.45	4.52	3.96
13363	76.34	342.96	4.49	2.82
11808	420.88	1837.2	4.37	3.82
12630	373.62	1629.33	4.36	3.79
	70.47	304.12	4.32	3.04
57435	576.02	2439.87	4.24	3.68
268482	41.68	172.46	4.14	2.85
11808	61.25	245.3	4	2.81
99000	39.75	154.89	3.9	3.1
112419	288.89	1124.56	3.89	3.19
235674	99.42	369.53	3.72	3.03
12491	226.91	801.02	3.53	3.18
54418	105.7	369.74	3.5	2.77
12702	804.45	2796.16	3.48	3.03
12182	62.85	216.55	3.45	2.95
15129 /// 15130	418.14	1439.69	3.44	3.04
12702	213.89	732.16	3.42	2.51
	76.8	262.28	3.42	2.11
12704	94.98	321.8	3.39	2.63
16569	45.41	152.78	3.36	1.56
228543	76.95	257.09	3.34	2.49
19662	368.94	1227.89	3.33	2.94
107513 /// 636537	165.39	539.01	3.26	2.59
16956	1460.67	4668.06	3.2	2.96
69097	170.18	540.9	3.18	2.48

	64.08	200.59	3.13	1.52
239618	416.8	1297.2	3.11	2.44
69065	522.75	1618.74	3.1	2.57
11722	109.84	339.17	3.09	2.68
18162	221.36	665.67	3.01	2.66
15122	103.41	307.69	2.98	2.43
84112	58.88	175.14	2.97	1.86
233529	288.78	847.16	2.93	2.5
241624	97.9	286.83	2.93	2.31
14939	185.06	533.48	2.88	2.16
74570	86.2	247.92	2.88	1.4
13166	60.85	172.37	2.83	1.88
	108.98	307.13	2.82	1.97
218454	155.64	422.65	2.72	1.97
330369	58.9	159.68	2.71	2.24
	70.32	188.46	2.68	1.82
552908	299.58	794.84	2.65	2.21
17777	497.95	1315.5	2.64	2.09
241490	124.64	328.22	2.63	2.21
192285	63.87	167.71	2.63	1.32
77462	88.52	231.67	2.62	2.37
19384	81.75	210.75	2.58	1.93
76969	66.62	172.16	2.58	1.72
12350	3463.89	8915.03	2.57	2.25
77041	88.33	225.66	2.55	2.09
73442	66.4	168.31	2.53	1.45
	132.35	333.75	2.52	2.07
278507	136.24	341.85	2.51	1.99
76768	636.9	1595.41	2.5	2.15
59020	376.12	942.13	2.5	2.02
654820	132.05	330.13	2.5	1.95
20776	115.11	286.23	2.49	1.56
15122	104.3	256.87	2.46	2.03
11767	464.36	1119.57	2.41	1.77
21743	81.45	195.08	2.4	1.93
170439	78.89	189.18	2.4	1.52
21832	133.1	318.67	2.39	2.07
14598	73.45	175.35	2.39	1.93
331063	105.01	250.54	2.39	1.93
67800	1525	3622.74	2.38	2.2
331063 /// 74548	164.77	391.13	2.37	1.83
69368	104.74	248.23	2.37	1.76
22041	685.96	1611.52	2.35	2.15
67896	770.11	1812	2.35	1.98
12491	116.16	271.97	2.34	2.08
58185	374.37	877.4	2.34	1.88
71760	81.08	188.96	2.33	1.47
11806	1507.83	3437.93	2.28	1.91
23966	86.55	197.7	2.28	1.77
77059	133.7	301.92	2.26	1.38
17005	96.73	215.47	2.23	1.93
94071	943.66	2091.84	2.22	2.09

11806	1844.86	4088.35	2.22	1.81
74131	251.01	558.21	2.22	1.66
11806	1848.18	4092.43	2.21	1.81
72315	167.46	370.2	2.21	1.49
319448	206.74	455.72	2.2	1.78
237636	268.33	588.67	2.19	1.89
15957	95.58	208	2.18	1.52
109815	192.77	419.05	2.17	1.99
14132	223.33	484.04	2.17	1.92
677168	170.82	368.93	2.16	1.96
52793	100.32	213.16	2.12	1.9
13869	91.83	194.82	2.12	1.53
668462 /// 668581 //	798.97	1684.93	2.11	1.83
22361	370.74	781.32	2.11	1.75
11997	278.73	585.5	2.1	1.88
667373 /// 669129	1583.55	3306.35	2.09	1.96
56185	259.57	542.02	2.09	1.79
11813	482.14	1000.52	2.08	1.85
70617	735.83	1532.1	2.08	1.84
11770	3395.8	7038.28	2.07	1.68
243219	689.73	1419.42	2.06	1.84
19218	142.22	292.68	2.06	1.8
71776	128.61	265.19	2.06	1.56
	97.03	198.93	2.05	1.58
18996	233.94	480.67	2.05	1.43
219228	101.46	207.66	2.05	1.41
73720	108.19	221.94	2.05	1.37
93671	1163.43	2346.25	2.02	1.74
12180	853.27	1715.03	2.01	1.54
	154.77	310.05	2	1.85
70300	114.64	229.76	2	1.61
353047	270.72	541.66	2	1.42
50912	111.94	224.33	2	1.41
14825	206.94	413.06	2	1.36

upper bound of FC	difference of means
-100000000	-503.62
-100000000	-168.61
-213.36	-259.9
-139.26	-753.61
-26.52	-686.63
-100000000	-137.92
-100000000	-193.88
-100000000	-168.99
-205.4	-118.2
-20.29	-197.15
-15.22	-363.82
-100000000	-779.21
-12.23	-1098.83
-9.61	-1450.87
-12.59	-133.46
-20.14	-146.93
-20.59	-166.76
-8.1	-608.15
-117.71	-248.38
-8.52	-229.69
-8.35	-151.27
-5.71	-1039.58
-11.43	-225.79
-10.06	-191.36
-7.54	-171.65
-11.45	-111.25
-6.79	-130.12
-4.19	-2225.34
-4.67	-863.3
-4.74	-402.21
-100000000	-827.46
-4.66	-187.76
-4.23	-154.77
-4.46	-180.04
-4.64	-152.46
-3.42	-1782.33
-4.5	-1112.57
-16.47	-122.07
-4.75	-150.71
-3.49	-219.8
-3.37	-376.98
-4.54	-168.02
-3.48	-338.05
-4.09	-102.54
-3.88	-102.38
-3.09	-2790.8
-4.8	-101.34
-3.59	-233.3
-3.8	-221.61

-4.48	-104.56
-2.98	-1018.57
-2.95	-294.15
-3.25	-214.37
-5.54	-163.39
-5.25	-175.55
-2.99	-105.1
-4.71	-249.73
-3.08	-446.42
-3.06	-124.09
-2.94	-114.3
-2.68	-1072.49
-3.01	-680.06
-2.8	-189.83
-3.58	-241.34
-4.21	-158.57
-4.49	-180.71
-3.37	-213.78
-3.99	-101.54
-2.5	-243.49
-2.82	-121.51
-2.8	-407.17
-3.16	-117.63
-2.8	-110.27
-2.9	-187.58
-2.55	-397.75
-3.1	-364.11
-2.39	-158.79
-3.93	-133.18
-3.16	-168.58
-2.53	-128.33
-2.63	-832.44
-3.65	-103.14
-2.85	-163.29
-2.49	-134.43
-2.67	-103.27
-2.45	-225.72
-2.34	-170.73
-2.39	-173.33
-3.1	-201.32
-2.31	-1665.8
-2.86	-953.88
-2.61	-222.74
-3.03	-187.52
-2.73	-116.25
-2.74	-114.71
-2.87	-141.45
-2.65	-523.52
-2.4	-129.98

upper bound of FC	difference of means
100000000	4440.11
100000000	253.46
100000000	318.04
100000000	143.74
14.43	664.84
27.35	218.84
10.44	5240.41
11.88	230.61
10.34	487.66
9.37	611.47
8	2372.59
8.24	1070.34
7.63	1743.94
8.43	671.38
9.37	341.23
7.33	572.35
14.74	143.68
7.25	795.8
6.49	628.11
7.94	608.24
6.57	1314.63
7.29	209.86
7.58	359.53
8.19	461.38
7.46	376.65
7.32	105.05
5.24	494.14
8.86	266.62
5.05	1416.32
5.01	1255.71
5.82	233.65
4.98	1863.86
6.03	130.79
6.87	184.05
4.99	115.14
4.64	835.67
4.58	270.11
3.92	574.11
4.38	264.04
3.98	1991.72
4.02	153.71
3.93	1021.56
5.24	518.27
5.49	185.48
4.7	226.81
49.88	107.37
4.35	180.14
3.75	858.95
4.25	373.62
3.47	3207.39
4.42	370.71

7.57	136.5
3.84	880.4
3.68	1095.99
3.61	229.33
3.43	444.31
3.76	204.28
7.28	116.26
3.54	558.38
3.94	188.92
4.28	348.42
9.36	161.72
4.67	111.52
4.7	198.15
3.7	267.01
3.24	100.79
3.75	118.14
3.28	495.27
3.55	817.56
3.23	203.58
4.62	103.84
2.91	143.15
3.54	129
4.91	105.53
3	5451.15
3.18	137.33
6.51	101.91
3.17	201.41
3.05	205.61
2.94	958.52
3.06	566.01
3.39	198.08
5.35	171.12
3.1	152.57
3.2	655.21
3.02	113.63
4.13	110.3
2.81	185.58
2.96	101.89
3.07	145.52
2.56	2097.74
3.15	226.36
3.09	143.48
2.59	925.56
2.74	1041.89
2.66	155.81
2.93	503.03
5.31	107.88
2.76	1930.09
3.13	111.15
5.01	168.21
2.6	118.74
2.35	1148.18

2.8	2243.49
3.07	307.2
2.81	2244.26
3.49	202.73
2.67	248.98
2.56	320.34
2.94	112.42
2.38	226.28
2.45	260.7
2.38	198.11
2.39	112.85
2.74	102.99
2.48	885.96
2.6	410.58
2.37	306.77
2.23	1722.8
2.47	282.45
2.35	518.38
2.36	796.26
2.49	3642.48
2.33	729.69
2.39	150.46
2.82	136.58
2.86	101.9
3.42	246.73
2.8	106.2
2.83	113.76
2.39	1182.82
2.7	861.76
2.18	155.27
2.54	115.12
2.62	270.95
2.65	112.39
3.12	206.12