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An in vitro model of murine middle ear epithelium

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Summary statement: Development and systematic characterisation of an *in vitro* otopathogenic infection model of the murine middle ear epithelium as a tool to better understand the complex pathophysiology of Otitis media.

Abstract

Otitis media (OM) or middle ear inflammation is the most common pediatric disease and leads to significant morbidity. Although understanding of underlying disease mechanisms is hampered by complex pathophysiology it is clear that epithelial abnormalities underpin the disease. There is currently a lack of a well characterised in vitro model of the middle ear (ME) epithelium that replicates the complex cellular composition of the middle ear. Here we report the development of a novel in vitro model of mouse middle ear epithelial cells (mMECs) at air liquid interface (ALI) that recapitulates the characteristics of the native murine ME epithelium. We demonstrate that mMECs undergo differentiation into the varied cell populations seen within the native middle ear. Proteomic analysis confirmed that the cultures secrete a multitude of innate defence proteins from their apical surface. We could show that the mMECs supported the growth of the otopathogen, NTHi, suggesting that the model can be succesfully utilised to study host pathogen interactions in the middle ear. Overall, our mMEC culture system can help better understand the cell biology of the middle ear and improve our understanding of the pathophysiology of OM. The model also has the potential to serve as a platform for validation of treatments designed to reverse aspects of epithelial remodelling underpinning OM development.

Introduction:

Otitis Media (OM) or inflammation of the middle ear is the most prevalent childhood disease, a leading cause of surgery in developed countries, and a significant reason for pediatric mortality in developing countries. Eighty percent of children suffer from at least one episode of OM by three years of age (Bakaletz 2010, Woodfield & Dugdale 2008).

The middle ear epithelium is similar to the respiratory epithelium and is composed of ciliated cells, secretory cells, non secretory cells and basal cells. Secretory cells are responsible for the production of mucins and various anti-microbial proteins such as lactotransferrin, lysozyme, defensins and surfactants (Lim et al 2000, McGuire 2002). The epithelium, along with its secretions, is involved in maintaining homeostasis and sterility within the middle ear cavity (MEC). Epithelial remodelling, characterised by mucociliary metaplasia and infiltration of the MEC with inflammatory cells, is a common feature of OM (Straetemans et al 2001).

In most animals, the middle ear is a relatively inaccessible organ lined by a thin mucociliary epithelium and sampling of the mucosa is a terminal procedure. Human middle ear tissue can be acquired only during surgical procedures and this limits the amount of sample available for study of OM. Culturing of middle ear cells *in vitro* enables maximisation of the available material, allows the effect of modifying culture conditions to be studied more easily and also allows functional studies to be performed. Previously, attempts have been made to culture middle ear epithelial cells from a number of organisms including rats (Toyama et al 2004, Ueyama et al 2001, Vanblitterswijk et al 1986), mice (Tsuchiya et al 2005), chinchillas (Amesara et al 1992, Nakamura et al 1991), gerbils (Herman et al 1992, Portier et al 2005, Takeno 1990), rabbits (Schousboe et al 1995) and humans (Choi et al 2002, Chun et al 2002, Moon et al 2000). These studies have included organ and explant cultures, primary cell cultures and development of middle ear cell lines.

However, there remains a lack of a robust *in vitro* middle ear epithelial model that differentiates into the different epithelial cell types of the middle ear and is free of fibroblast contamination. This has greatly restricted the ability to identify the function of different cell types and their products within the middle ear and limits our understanding of the pathophysiology of OM development.

We report here the development of a novel *in vitro* primary model of the mouse middle ear epithelium using air liquid interface (ALI) culture and systematically

characterise the different cell types present in the middle ear. We also demonstrate, that this culture system can be utilised to study host pathogen interactions within the middle ear and thus has the potential to allow investigatation of the mechanisms of OM pathogenesis.

Results:

We established an air liquid interface (ALI) culture system to model the mouse middle ear epithelium *in vitro* (Fig. 1A). We performed a morphological analysis and systemactically characterised the various epithelial cell types expressed by our model in comparison to the native mouse middle ear epithelium.

Cell culture characteristics

The average number of epithelial cells isolated was 74,667±10,621 cells per MEC (n=12 batches). Primary culture of mMECs proceeded in two phases – a proliferative phase in submerged culture and a differentiation phase at air liquid interface (Fig. 1B-H). mMECs were seeded at a density of 1x10⁴ cells/ membrane and 3 days after seeding 16.8±2.6% adhered to the membrane and had started forming small epithelial islands (Fig. 1B). The attached cells began to elongate to establish contact with neighbouring cells and began to proliferate rapidly from day 5 to day 7 (Fig. 1C, D). Cells formed a confluent monolayer of flat, polygonal cells within 9 to 10 days in submerged culture (ALI Day 0) in the presence of Rho Kinase inhibitor, ROCKi (Fig. 1E). The morphology of the cells became more complex when transferred to ALI. At ALI Day 3, cells started changing in size and shape (Fig. 1F) and by ALI Day 7 two distinct sub populations of cells could be observed, the majority of which were flat polygonal cells, intersected with clusters of slightly elevated, more compactly arranged cells (Fig. 1G). Around ALI Day 9, ciliary beating could be seen in these 14 clusters under phase contrast microscope. ALI Day cells (1.2x10⁵cells/membrane) displayed a cobble-stone appearance with well-defined cell boundaries; and were a combination of flat polygonal cells and compactly clustered elevated cells with actively beating ciliated cells. (Fig. 1H, Movie S1). No cells with fibroblast-like morphology were seen in the mMEC ALI cultures. However, fibroblasts were isolated through differential adherence to plastic during the cell isolation process. (Fig. 1I).

Cell morphology

Electron microscopy analysis during ALI culture revealed the development of a mucociliary epithelium (Fig. 2). At ALI Day 0 (undifferentiated cells), scanning electron microscopy showed uniformly flat, large cells with microvilli on the apical surfaces (Fig. 2A). At ALI Day14, the cells exhibited a dome shaped appearance (Fig. 2B), areas of flatter polygonal and secretory cells with microvilli on the apical surfaces and areas abundant in ciliated cells (Fig. 2C). The morphology of the ALI Day 14 mMEC cultures resembled the *in vivo* middle ear epithelium (Fig. 2D). Transmission electron microscopy revealed that ALI Day 14 cells were polarized with desmosomes on the baso-lateral surfaces suggesting the formation of tight junctions, another feature of epithelial cells (Fig. 2E). The formation of tight junctions was further confirmed by uniform expression of ZO-1 in the cell membrane (Fig. 2F)

Expression of epithelial markers by mMEC cultures

The expression of a selected panel of genes, known to be expressed by the middle ear epithelium and upper airways, was analysed by RT- PCR of RNA from the original mMECs before seeding and compared to ALI Days 0 and 14 cells. Fibroblasts isolated by differential adherence were used as a negative control for epithelial markers (Fig. 3A). Bpifa1 and Bpifb1 are secreted, putative innate immune molecules expressed in the upper airways. Bpifa1 was expressed strongly in the original and ALI Day 14 cells, but lower in the undifferentated ALI Day 0 cells. Bpifb1 was detected only in the original cells, not in cultured cells. Tekt1 (a marker of ciliated cells) was detected at ALI Day 14. Analysis of Muc5ac and Muc5b expression, markers of goblet cells, suggested that Muc5ac was weakly expressed in mMEC original cells but was not detectable in the cultured cells, whereas Muc5b was expressed more strongly in the original cells and maintained this expression to ALI Day 14. We also studied the mucosal innate immune genes Lactotransferrin, Surfactant protein D (Stfpd) and Regenerating islet-derived protein 3 gamma (Reg3 γ). Lactotransferrin and Reg3 γ were detected in the mMEC original cells and at ALI Days 0 and 14, whereas expression of Stfpd was seen in the cells during ALI differentiation. As expected, the expression of *Keratin5*, a marker of basal cells, was reduced as cells differentiated from ALI Day 0 to Day 14. The expression of these epithelial markers in the mMEC cultures indicates that the cells differentiate in culture from ALI Day 0 to ALI Day 14 and the pattern of expression in the

differentiated cells is in line with that seen in the mMEC original cells isolated from the middle ear. The absence of *Vimentin* in ALI Day 14 cultures indicates that our mMEC cultures are devoid of fibroblast contamination. *Oaz1* was used as a housekeeping gene for all RT-PCR experiments.

MS analysis of the apical secretome of mMECs

To complement our gene expression studies, we also performed a global proteomic analysis of apical ALI Day 14 secretions of mMEC cells by Orbitrap mass spectrometry (MS). Table 1 lists the most abundant secreted proteins identified classified according to their emPAI score. The most abundant secreted protein was Lactotransferrin, with Serotransferrin, Reg3^v, Lipocalin2, Ceruloplasmin and Bpifa1 also being found at high levels. Multiple anti-proteinase and proteinase proteins were also found in the secretions including members of the WFDC family, WFDC2, WFDC18 (EXPI) and SLPI (WFDC4), as well as multiple cathepsins. We validated the secretion of Bpifa1, Lactotransferrin and Reg3^v in the apical washes from the differentiating mMEC cells using western blotting (Fig. 3B,C). The full list of proteins identified is given in Table S1.

Localisation of epithelial markers in mMEC cultures.

We used immunofluorescence confocal (IFC) microscopy to study the localisation of epithelial markers in ALI Day 0 and ALI Day 14 cultures, in order confirm the differentiation process at the protein level. ALI Day 0 cultures showed abundant staining of P63 (basal cell marker), scanty staining of Bpifa1 (Fig. 4A) and no staining of FoxJ1 (ciliated cells) and Muc5B (goblet cells) (Fig. 4C). However by ALI Day 14, the cells had differentiated into multiple cell types. These stained strongly for Bpifa1, had reduced levels of p63 (Fig. 4B) and were populated with ciliated and goblet cells (Fig. 4D). Bpifa1 was localised the non ciliated population in differentiated mMECs, consistent with that seen in the *in vivo* middle ear epithelium (Fig. S1). In keeping with proteomic and expression data, we also detected abundant cytosolic levels of Lactotransferrin and Reg3^v in the ALI Day 14 cultures (Fig. 4E, F). Staining of nuclei with DAPI and Z-slice imaging using confocal microscopy also demonstrated that the cells formed a flat monolayer at ALI Day 0, but showed a more complex reorganisation by ALI Day14 with a combination of flat

and elevated cells showing 2 or 3 different layers, with nuclei further away from the base of the membrane (Fig. 4G, H).

Relative abundance of secretory and ciliated cells

As noted above, phase contrast microscopy and SEM of mMEC cultures demonstrated that the ALI Day 14 cultures were composed of distinct anatomical areas of flat polygonal cells and patches of more elevated pseudostratified cells (Fig. 1H, Fig. 2B). To study this in more detail, we used IFC to determine the localisation of different epithelial markers within the two regions of cellular morphology. FoxJ1 positive ciliated cells and Muc5B positive goblet cells, were restricted to the elevated pseudostratified cell clusters, whilst Bpifa1 was more commonly seen in the flatter cells, although some staining was seen in the elevated cells, especially near the periphery (Fig. 5). This was consistent with our observation that ciliated cells could be seen beating in the elevated clusters of cells under light microscopy. We also confirmed Bpifa1 and Muc5B positive cells were not ciliated (Fig. 5A, B). Again this alaysis confirmed the existence of elevated pseudostratified cells within the cultures (Fig. 5C,D).

mMEC ALI cultures as an otopathogenic infection model

Having established a novel mMEC culture model, we wanted to evaluate its utility as a model system for the study of host pathogen interactions in the middle ear. We infected differentiated ALI Day 14 cells with a GFP-tagged *NTHI 375^{SR}*. IFC microscopy indicated that only a few cells were infected 24 hpi but by 48 hpi the infection rate had increased and the bacteria continued to spread laterally in culture infecting the majority of cells by 72 hpi (Fig. 6A-C). We confirmed this observation by quantifying the amount of green fluorescence using image analysis and observed that in every batch the amount of bacterial infection increased in a time dependent manner (Fig. 6E, Fig. S2).

We also studied the expression of the pro-inflammatory chemokine *Cxcl2/MIP2a* during the progression of infection. *Cxcl2* was significantly upregulated post infection at 24hpi (131 fold) and 48hpi (36 fold) compared to the 24hpi MOCK sample set as the reference (Fig. 6F). This data confirms that our mMEC model is capable of eliciting an inflammatory response.

Discussion:

We have developed a novel *in vitro* model for the culture and differentiation of primary mouse middle ear epithelial cells (mMECs) cultured at an air liquid interface. The ALI system has previously been used to culture respiratory epithelial cells (TBE) from several species (Clarke et al 1992, Davidson et al 2000, Yamaya et al 1992, You et al 2002) and more recently, it has been applied to the culture of murine nasal epithelial cells (Woodworth et al 2007). The exposure of apical cell surfaces to air and the supply of nutrients from the basal compartment mimics the *in vivo* upper airway epithelium and promotes differentiation. As the middle ear epithelium can be considered to be an extension of the upper airways and exhibits physiological similarities to the upper respiratory tract, we reasoned that we could extend the use of the ALI system for the successful culture of primary mMECs.

Our mMEC cultures can be used to study both proliferation and differentiation of middle ear cells. The medium used for the culture of mMECs was the same as that used for culture of mouse tracheal epithelial cells, mTECs (You et al 2002). The media supplements, epidermal growth factor, insulin, transferrin and cholora toxin enhance cell proliferation and ciliogeneisis whereas retinoic acid is important for differentiation of mucous cells (Lechner et al 1982, Wu et al 1997, You & Brody 2013, You et al 2002). The addition of ROCKi to the culture medium has been shown to enhance basal cell proliferation in airway epithelial cultures (Horani et al 2013). Using ROCKi, we could maximise the number of transwell-cultures established from the limited number of cells isolated from the thin middle ear epithelium without altering differentiation (Fig. S3); an observation that has been made in both human and mouse airway cells (Butler et al 2016, Horani et al 2013) and thus reduce the number of animals required for each batch of cells cultured. mMECs grown at ALI differentiated into different cell types by ALI Day 14, a typical timescale used for the differentiation of murine respiratory epithelial cells (You et a 2002). We extended this culture period to ALI Day 18 without any morphological differences (data not shown); however further time points were not tested.

An important and common problem identified in a number of previous attempts to grow middle ear epithelial cells was the contamination and overgrowth of fibroblasts in the cultures (Nakamura et al 1991, Tsuchiya et al 2005, Vanblitterswijk et al 1986). By adding a differential adherence step, in which fibroblasts adhere to plastic in preference to the epithelial cells, we were able to eliminate fibroblasts from our

mMEC cultures, generating a pure epithelial population, as shown by a lack of expression of the fibroblast marker, *Vimentin*, in ALI Day 14 cultures.

The mMEC cultures model the native middle ear epithelium as the cells exhibit characteristic epithelial features such as a cobble-stoned morphology, formation of tight junctions, apical-basal polarised, presence of desmosomes and apical microvilli. The cultures contained a combination of single layers of flatter polygonal cells and clusters of pseudostratified, dome shaped, elevated cells, with some of these having beating cilia. The dome shaped appearance of cells can be attributed to an active ion transport mechanism (Chun et al 2002, Nakamura et al 1991).

A major limitation of previous middle ear epithelial cultures has been the lack of differentiation into distinct epithelial cell types. The middle ear epithelium, like the upper airway epithelium, is composed of ciliated cells, basal cells, goblet cells and other secretory cells. Previous studies outlined difficulties in maintaining ciliated cells in culture (Nakamura et al 1991, Ueyama et al 2001). To our knowledge, the development of ciliated cells has only been described in one study of human middle ear cultures (Choi et al 2002), also grown at the ALI. Our cultures clearly show the presence of actively beating cilia and also stain for nuclear FoxJ1 protein. We found that the distribution of cilia in our cultures partially mimics that seen in the native middle ear epithelium by SEM (Fig. S4). Parts of the middle ear epithelium are populated with tracts of dense cilia, parts with interspersed ciliated and non-ciliated cells and some parts with a simple epithelium composed of flat non-ciliated polygonal cells. This distribution of cell types within the middle ear cavity is supported by previous studies (Lim 1979, Thompson & Tucker 2013) which show that the native epithelium of the hypotympanum and towards the opening of the ET is densely packed with cilitaed cells, whereas the mesotympanum contains ciliated islands amongst other non ciliated cells. The epitympnaum is composed primarily of of flat squamous cells which may also contribute to the secretory defence system. On this basis, we believe that our mMECs, which are a combination of flat non ciliated, secretory and ciliated cells closely model the morphology of the native middle ear epithelium.

Mucins are products of goblet cells, unique to the mucosal epithelia, and are essential in the maintenance of mucosal innate defence. A number of attempts have been described to utilise the available middle ear models to study mucin gene expression at the transcriptional level (Kerschner et al 2010, Liu et al 2016, Moon et

al 2000, Tsuchiya et al 2005). However, these studies were limited by the lack of production and localisation of detectable amounts of mucins at the protein level. Goblet cells (as shown by MUC5B positivity) were seen in the elevated pseudostratified clusters of our ALI Day 14 mMEC cells in close association with ciliated cells. It has previously been shown that in the native middle ear epithelium the abundance of mucous secreting cells is in parallel to the distribution of ciliated cells, reiterating that our model mimics the *in vivo* middle ear epithelium (Lim et al 1973). MUC5B is the predominant mucin in COM effusions (Preciado et al 2010). It has been shown to be indispensible for airway mucocilary clearance and maintainence of mucosal homeostasis and Muc5b^{-/-} mice develop OM (Roy et al 2014) Expression of MUC5B at proteomically detectable levels as demonstrated by IFC enables our model to be potentially utilised for further study of the role of mucins in the middle ear epithelium. Moreover, treating primary tracheal and bronchial epithelial cells grown at ALI with IL-13 has been shown to induce goblet cell hyperplasia(Atherton et al 2003, Kondo et al 2002). Our model provides a platform to study middle ear mucous hypersecretory phenotypes in vitro. The lack of MUC5AC or MUC5B in our MS analysis may be due of the loss of these high molecular weight glycoproteins during the sample processing and preparation.

In addition to ciliated and goblet cells, our cultures also produce a range of other secretory cell products. Bpifa1 is the most widely studied member of the BPI fold (BPIF) containing family of putative host defence proteins (Bingle et al 2011) and we have previously shown it to be an abundant product of the murine upper respiratory tract and nasopharynx (Musa et al 2012). BPIFA1 was identified as a candidate for OM in a recent GWAS study (Rye et al 2012) and loss of the protein has been implicated in the development of OM in aged mice (Bartlett et al 2015). Bpifa1 was readily detectable in mMEC cultures and apical secretions from the cells. Bpifa1 was localised to non-ciliated cells in the cultures, which is consistent with studies from the respiratory tract (Barnes et al 2008, Kim et al 2006, Musa et al 2012). Bpifb1, on the other hand, was detectable in the original cells, but not in the ALI Day 14 cultures and was absent from the list of proteins identified in the proteomic analysis performed using mass spectrometry. Bpifb1 protein is localised to goblet cells and minor glands associated with the Eustaschian tube (ET), but its expression is limited in the middle ear epithelium (data not shown). This observation suggests that our mMEC cultures model the middle ear epithelium rather than the ET epithelium.

Proteomic analysis by MS identified a number of secretory proteins in the apical washes from the cells. The most abundant proteins comprised a variety of secretory host defence proteins with anti-microbial roles, proteins involved in cellular proliferation, wound repair, stress response, compliment activity and maintenance of cellular homeostasis. It is notable that our MS data contains a number of proteins also identified in a proteomic analysis of ear exudates from Chronic OM patients such as Lactotransferrin, Bpifa1, Lipocalin, Lysozyme and various cathespins and complement proteins (Val et al 2016). Lactotransferrin was the most abundant protein identified in our study. It is an innate immune protein secreted by airway mucosal surfaces that has been shown to play a role in maintenance of middle ear immunity (Bernstei.Jm et al 1974, Lim et al 2000, Moon et al 2002). It prevents colonisation of mucosal surfaces by scavenging environmental iron, thus limiting its availability for bacterial growth. Human milk Lactotransferrin has been shown to attenuate the pathogenic potential of Haemophilus influenzae by proteolytically cleaving two important colonisation factors found on the bacterial surface (Hendrixson et al 2003) and the administration of Apolactoferrin has been shown to reduce bacterial counts in chinchilla middle ears with pneumococcal induced OM (Schachern et al 2010). Surfactant protein D was originally identified as a lung surfactant associated protein but is also expressed in the ME and the ET and has a suggested role in enhancing opsonisation and phagocytosis of bacteria (Lim et al 2000, van Rozendaal et al 2001, Wright 1997). Reg3y is C-type lectin produced by Paneth cells and secreted into the intestinal lumen with a suggested bactericidal activity against gram-positive bacteria by binding to peptidoglycan in the bacterial cell wall (Cash et al 2006). It has been shown to spatially regulate the separation of microbiota from the host small intestinal epithelium (Vaishnava et al 2011). Reg3 $\gamma^{-/-}$ mice have an altered mucosal distribution and increased inflammatory response in the small intestine (Loonen et al 2014). *Reg3y* has also been shown to be involved in pulmonary innate immunity as it is induced by Stat3 during methicillin resistant Staphalococcus aureus (MRSA) infection in lung and inhibits MRSA growth in vitro (Choi et al 2013). This is the first study reporting the expression of this protein in the ME and it is possible that Reg3y performs a similar function in the middle ear.

It was noticeable that the proteomic data unexpectedly contained many intracellular proteins. We also detected the presence of the membrane tethered mucins Muc1, Muc4 and Muc18. (Table S1). This can be reasoned to be the content of secreted

exosomes. Exosomes are small membrane bound units released by the fusion of endosomal micro vesicular bodies with the apical plasma membrane. They have been suggested to be involved in stimulating immune responses, modulating secretory activities and engaging in cell communication by packaging and delivering microRNAs to other cells (Keller et al 2006, Valadi et al 2007). Exosomes are released by epithelial cells (Kapsogeorgou et al 2005, van Niel et al 2001), found in bronchoalveolar lavage fluids (Admyre et al 2003) and also in the apical secretions from human bronchioalveloar epithelial (HBE) cultures (Kesimer et al 2009, Pillai et al 2014). The presence of exosome associated proteins in our mMEC secretome further adds to the potential utility of this model to study the role of exosomes in middle ear biology and OM pathogenesis.

Infection of mMECs using the human otopathogen, NTHi demonstrates that our culture system can be effectively utilised for the study of host pathogen interactions within the middle ear. Our studies show that, NTHi initially infected a small number of cells in culture and the infection spread laterally over time. Since the advent of pneumococcal vaccines, NTHi is the most common pathogen in OM (Casey & Pichichero 2004). Epithelial remodelling, one of the most common features of OM, is characterised by mucous metaplasia. The chemokine $Cxcl2/MIP2\alpha$ is the murine homologue of IL-8 and is a key mediator of overproduction of mucin (Juhn et al 2008). NTHi infection is known to stimulate Cxcl2 upregulation in several murine tissues including the middle ear, lungs and the inner ear (Gaschler et al 2009, Lim et al 2007, Woo et al 2012). Moreover, Cxcl2 was identified as the most upregulated gene when mice were trans-tympanically injected with NTHi and on infection of the mouse middle ear epithelial cell line (mMEEC) with NTHi (Preciado et al 2013). Our data shows that Cxcl2 was significantly upregulated on NTHi infection at 24 and 48 hours post infection, suggesting that our mMECs respond to bacterial infection in a manner similar to the native middle ear epithelium. This opens up a new avenue to utilise this system to study the response of middle ear cells to different insults, injuries and infections. Our mMEC cultures can potentially be utilised to study the interaction of host middle ear epithelial cells with a variety of bacterial as well as viral otopathogens.

Previous studies have demonstrated successful gene silencing in primary HBE cells and mTECs using lentiviral transduction systems (Horani et al 2012, Vladar & Stearns 2007). Recently the CRISPR/CAS9 genome editing system was utilised in HBE cells and mouse tracheal organoid cultures in order to study genes involved in the regeneration of basal cells into mucocilliary cells of the airway epithelium (Gao et al 2015). The physiological similarities of mMEC cells with airway epithelial cells and the use of similar culture conditions for their growth means that the mMEC model might be amenable to gene editing studies in future.

It is known that primary airway epithelial cells cultured from cystic fibrosis and asthma patients maintain the disease phenotype in culture (Davies et al 2003, Matsui et al 1998). A number of mouse models are available for the study of OM. These include mice deficient in innate immunity genes such as *Evi1*, *Fbxo11*, *TLRs* and *Myd*88, cilliary development genes such as *Dnahc5* (Rye et al 2011) and goblet cells (Roy et al 2014). It will be interesting to see if our mMEC culture system can be utilised to reproduce the OM phenotype of these mouse mutants in vitro and enable comparative studies between unaffected and diseased cultures. OM often involves complex responses involving the middle ear epithelium, sub-epithelial mesenchyme, inflammatory cells and middle ear effusion, making it challenging to identify epithelial cell-specific responses. The mMEC culture system will provide us with the ability to isolate and assay responses of specific sub poupulations of epithelial cells. Our cell isolation method eliminates the influence of explants and excludes fibroblasts from culture, which were two of the most important confounding factors showin in previous studies of primary middle ear epithelial cells. It is a 3-dimentional model of the middle ear epithelium and hence mimics the in vivo physiology more closely compared to cell lines. The possibility of replicating the phenotype of the genetic models of OM, the capacity to easily manipulate differentiation of cells by modifying culture conditions and the ability to infect cells with various ototpathogens makes our model widely applicable to the wider OM community. The availability of such a wellcharacterised model of the middle ear epithelium can help better understand the cell biology of the middle ear and improve our understanding of the pathogenesis of OM.

Materials and Methods

Ethics statement

Humane care and animal procedures were carried out in accordance to the appropriate UK Home Office Project licence. Randomised male and female 8-10 week old C57BL/6 and C3H/HeH mice, housed in individually ventilated cages (Techniplast UK Ltd) under specific pathogen free (SPF) conditions were obtained from MRC Harwell, UK.

Dissection of the middle ear cavity

The detailed protocol used for dissection is outlined in Fig S5. Mice were euthanized by terminal intra-peritoneal injection of 100µL pentobarbital (50mg/ml, Henry Schein®) and exsanguinated by cutting the inferior vana cava. Mice were decapitated, the skin at the nape of the neck was incised, bisected anteriorly and removed entirely to expose the bony surface of the skull and the lower jaw was detached under direct visualization. Under a dissecting microscope (Olympus SZx10), the skullcap was gently opened with a pair of fine forceps and the brain was removed. The head was bisected at midline and oriented with the opening of the ear facing upwards. Any muscle, soft tissue and remnant hair surrounding the ear were removed using fine dissecting scissors and forceps, leaving the MEC (bulla), still attached to the outer ear canal (OEC) and the inner ear (IE). The bony shell of the bulla was further cleaned free of any attached extraneous tissue. The OEC, which appears a shade lighter than the MEC, was gently broken away from the MEC using stork bill forceps. The tympanic membrane and the ossicles usually detached from the MEC along with the OEC. Alternatively, they were physically removed using fine stork bill forceps. Lastly, the cup-shaped MEC was carefully lifted away from the inner ear.

Isolation and differentiation of middle ear epithelial cells at air liquid interface

The protocol for primary culture and differentiation of mouse middle ear epithelial cells (mMECs) was adapted from a previously described method for mTECs; (You & Brody 2013, You et al 2002). For each batch of cells, bullae from approximately six mice (12 bullae) were pooled in a tube containing pronase (1.5 mg/ml) in *'mMEC basic media'*: DMEM/F-12 HAMs media (Life Technology, Cat No-

31330-038) supplemented with penicillin (100 µg /ml) and streptomycin (100 µg/ml) (Life Technology, Cat No- 15070-063) and subjected to overnight proteolysis at 4°C. The pronase was neutralised by the addition of 10% foetal bovine serum (FBS) and the bullae were gently agitated by inverting the tube 25 times. The bullae were then transferred to 2ml of fresh mMEC basic 10% FBS media, the tube was inverted again 25 times and this process was repeated 3 times. The combined proteolytic and mechanical actions led to dissociation of the middle ear cells from the bullae (Fig. S6). Media from the three tubes was combined and centrifuged at 500g for 10 minutes at 10°C. The pelleted cells were re-suspended in 1ml of media containing 1 mg/ml bovine serum albumin (BSA) and 0.5 mg/ml DNase I (Sigma-Aldrich, Cat No-DN25). Cell viability and number were assessed using trypan blue staining and a haemocytometer. Cells were centrifuged at 500g for 5 minutes at 10°C and the pellet re-suspended in 5ml of mMEC basic 10% FBS media. In order to seperate contaminating fibroblasts from epithelial cells a differential adherence step was performed by plating the cells on 60mm surface treated tissue culture dishes at 37°C in a 5% CO2 incubator for 3-4 hours. Fibroblasts attached to the plastic whilst the non-adherent epithelial cells were collected, centrifuged at 500 xg for 5 minutes at 10°C and re-suspended in 1ml of 'mMEC plus' media: mMEC basic media supplemented with 5% FBS, 30 µg/ml bovine pituitary extract (Life Technology, Cat No- 13028-014), 10 µg/ml of insulin (Sigma-Aldrich, Cat No- I1882), 25 ng/ml of mouse epidermal growth factor (BD Biosciences, Cat No- 354001), 5 µg/ml of transferrin (Sigma-Aldrich, Cat No- T1147), 0.1 µg/ml of cholera toxin (Sigma-Aldrich, Cat No- C8052) and 0.01 µM of freshly added retinoic acid (Sigma-Aldrich, Cat No- R2625).

For optimisation of culture conditions, cells were plated on either tissue culture plastic or sterile, 0.4µm pore sized transparent PET (Polyethylene Terephthalate) membranes coated with 150µL (50µg/ml) of rat-tail collagen type I (BD Biosciences, Cat No- 354236) in a 24-well supported transwell format (Falcon, Cat No-353095). mMECs were seeded at an initial density of 1 x 10⁴ and 2 x 10⁴ cells/well in the presence or absence of 10 µM of Rho Kinase inhibitor, Y-27632 dihydrochloride (ROCKi, Tocris bioscience, Cat- 1254) and 5 x 10⁴ cells/well without ROCKi. A seeding density of 1 x 10⁴ cells/well with ROCKi on transwell membranes was identified as optimum and therefore used for culturing all following batches of cells (Fig. S3A-G). An average of 671,788 ± 59,790 (n=9 batches) cells were obtained

from 12 pooled bullae. 30 to 35 transwells were typically seeded at this density and the remaining cells were lysed in Trizol reagent (Sigma- Aldrich, Cat No- T2494) to give freshly isolated mMEC original cells for comparison with the cultured cells.

Cells were initially cultured, submerged, in 'mMEC plus-proliferation media' with 300 µL of media in the top chamber and 700 µL in the bottom chamber. Media was changed every 48 hours, until the cells were completely confluent, thereafter media from the apical chamber was removed and media in the basal chamber was replaced with 'mMEC SF- differentiation media': DMEM/F-12 media supplemented with 1 mg/ml BSA (Life Technology, Cat No- 31330-038), 5 µg/ml insulin, 30 µg/ml bovine pituitary extract, 5 µg/ml transferrin, 5 ng/ml mouse epidermal growth factor, 0.025 µg/ml cholera toxin and freshly added 0.01 µM retinoic acid to induce ALI culture. This system of culture promotes differentiation of cells by mimicking the in vivo situation. Cells were differentiated at ALI for 14 days and media was changed every 48 hours. Cells were lysed in 250 µL of Trizol reagent for RNA extraction and apical washes were collected in 200 µL of sterile HBSS at ALI Day 0 (submerged Day 10), Day 3, Day 7 and Day 14. Figure 2A gives a brief overview of the complete cell culture system. Doubling time for cells seeded at 5 x 10⁴ cells/well in *mMEC*-*Plus media* without ROCKi was determined by trypsinizing the cells and using the formula: $PD = t \times Log2/(LogC2-LogC1)$ [PD = Population doubling, t = 48 hours, Log = 10 based Log, C1 = initial cell count, C2 = final cell count].

Immunoflorescence microscopy

Transwell membranes at ALI Day 0 and Day 14 were fixed with 10% phosphate buffered formalin at $37^{\circ}C$ for 30 minutes. Cells were permeabilised using 0.5% Triton X-100. Non-specific binding was blocked using 10% goat serum in PBS and incubation for 1 hour at 80rpm on an orbital shaker at room temperature. The membranes were washed 3 times with PBS for 5 minutes at 150 rpm and incubated with the following primary antibodies: anti-Bpifa1 (1:200), anti-Foxj1 (1:300), anti-p63 (1:100), anti-MUC5B (1:100), anti-ZO1 (1:200), anti-lacto transferrin (1:200) or anti-Reg3 γ (1:200), overnight at 4°C at 80rpm on the shaker. All antibody details are given in Table S2. The following day, membranes were washed 3 times with PBS for 5 minutes at 150 rpm and bound primary antibodies detected using 1:200 dilution of the appropriate fluorophore-tagged secondary antibody (Alexa-Fluor 568 Goat anti-rabbit antibody (Cat No- A11011) or Alexa Fluor 488 Goat anti-mouse antibody (Cat

No- A11001)) incubated for 1 hour at room temperature at 80 rpm in the dark. Membranes were washed 3 times as above, detached carefully from their transwell support with a fine scalpel and placed on a glass microscope slide with the cells facing upwards. Nuclei were counterstained using Vectashield DAPI mounting medium (Vector laboratories) and the cells imaged using an Olympus Fluoview 1000 Confocal microscope.

Scanning Electron Microscopy

ALI Day 0 and Day 14 cell membranes were washed free of culture media with sterile, warm HBSS and fixed in 3% Glutaraldehyde in 0.1M Sodium Cacodylate buffer overnight at 4°C. The membranes were washed 2 times with 0.1M Cacodylate buffer for 5 minutes each, detached from their transwell support as described above and postfixed in 2% aqueous Osmium tetroxide. The specimens were washed briefly in water, dehydrated in a graded ethanol series, dried in a 1:1 mixture of 100% ethanol: Hexamethyldisilazane (HEX) before final drying in 100% HEX. The membranes were placed overnight in a fume hood, mounted onto a pin-stub using a Leit-C sticky tab, (gold sputter coated) and examined using a Philips XL-20 SEM at 15kV.

Transmission Electron Microscopy

ALI Day 14 cell membranes were fixed with 3% Glutaraldehyde in 0.1M Sodium Cacodylate buffer, washed and post fixed in 2% Osmium Tetroxide as mentioned above. The detached membranes were washed briefly in water and dehydrated through graded ethanols, cleared in Epoxy-propane (EPP) and infiltrated in 1:1 mixture of araldite resin: EPP mixture overnight on a rotor. This mixture was replaced in two changes with fresh araldite resin mixture over an 8-hour period, before being embedded and cured at 60°C in an oven for 48-72 hours. Ultrathin sections (approximately 85nm thick) were cut on a Leica UC 6 ultra-microtome onto 200 mesh copper grids, stained for 30 minutes with saturated aqueous Uranyl Acetate followed by Reynold's Lead Citrate for 5minutes. Sections were examined using a FEI Tecnai Transmission Electron Microscope at an accelerating voltage of 80Kv. Electron micrographs were recorded using a Gatan Orius 1000 digital camera and Digital Micrograph software.

Non-typeable Haemophilus influenzae infections

mMECs were cultured in antibiotic-free media for 48 hours prior to infection. A GFP tagged, streptomycin resistant strain of the clinical OM isolate, NTHi-375 (NTHi 375^{SR}), derived from a Finnish pneumococcal vaccine study on children undergoing tympanocentesis in 1994-95 was used for all bacterial challenge experiments (Cody et al 2003). *NTHi* 375^{SR} was grown from glycerol stocks on brain heart infusion (BHI) agar plates supplemented with 2µg/ml Nicotinamide adenine dinucleotide hydrate (NAD, Sigma Aldrich, Cat No- N7004), 2µg/ml hemin (Sigma Aldrich, Cat No-H9039) and 200µg/ml streptomycin sulphate (Melford Laboratories, Cat No- S0148) at 37°C, 5% CO₂ overnight. The following day colonies were transferred into BHI broth supplemented with NAD and hemin and incubated for 3 hr at 37°C, 5% CO₂, 250rpm. The optical density (OD490) of 1ml of liquid culture was spectrophotometrically determined (Jenway 6300) and the culture diluted with PBS to give a concentration of 1x10⁹ bacteria/ml. An appropriate amount of culture in antibiotic-free mMEC-SF media was added to the apical chamber of ALI Day 14 mMEC cultures such that the membranes were infected at a multiplicity of infection (MOI) of 1:100 (mMECs: bacteria). An equal volume of sterile PBS was added to generate MOCK infected controls. The membranes were incubated at 37°C, 5% CO₂ for 1 hr and washed 3 times with sterile HBSS to remove non-adherent bacteria. Media was replaced in the basal chamber and cultures were incubated for 4, 24, 48 and 72 hours post infection (hpi). At each time point apical washes were collected, cells were lysed in Trizol reagent for RNA extraction and membranes were fixed for IFC as described. Membranes were visualized using confocal microscopy and infection was quantified by measuring the mean integrated fluorescence intensity of four central 10x fields at each time point, in each batch, using Image J software.

Reverse transcription PCR (RT-PCR)

For end-point RT- PCR, total RNA was extracted from at least 3 batches of freshly mMEC original cells and mMECS at ALI days 0 and 14 lysed in Trizol. RNA yield was determined using NanoDrop-1000 (Thermoscientific). Residual genomic DNA was digested by DNase I treatment (Promega, Cat No-M6101) and 200ng of RNA was reverse transcribed using AMV Reverse Transcriptase (Promega, Cat No-M9004). RT-PCR was performed with 1µl of template cDNA and Maxima Hot Start Green PCR Master Mix (ThermoFisher Scientific, Cat No- K1061). The cycling

conditions were: 95°C for 5 minutes; denaturation: 94°C for 1 minute (25-35 cycles); annealing: 60°C for 1 minute; extension: 72°C for 1 minute; final extension: 72°C for 7 minutes (MJ Research PTC-200). The primer pairs used are described in Table S3. The amplified PCR products were run on a 2% agarose gel containing 0.5μ g/ml ethidium bromide (Dutscher scientific, Cat No- 4905006) and bands visualised using a Biorad ChemiDocTM XRS+.

Real time quantitative PCR (RT-qPCR)

For RT-qPCR, total RNA was extracted from at least three independent batches of ALI Day 14 cultures infected with NTHi-375^{SR} for 24 and 48 hours and their corresponding MOCK infected cultures. Residual genomic DNA was removed using the DNA-free[™] kit (Life technologies, Cat No- AM1906), RNA quantified using NanoDrop-8000 (Thermoscientific) and integrity checked on an Agilent Bioanalyzer 2100 instrument using RNA 6000 Nano kit. 400ng of total RNA was reverse transcribed into cDNA in a 20µL reaction volume using a Superscript™ III First Strand Synthesis kit (Invitrogen, cat No- 11752-050), in accordance with manufacturer's instructions. RT-qPCR was performed using an Applied Biosystems TaqMan gene expression assay for Cxcl2 (Mm00436450_m1) on a 7500 Fast Real Time PCR system (Applied Biosystems), with 2x Taqman Fast Universal Master Mix (Applied Biosystems, Cat No- 4352042). 10ng cDNA was added to each reaction and three technical replicates were performed for each assay in each batch. Genetic expression levels of Cxcl2 were normalized to three endogenous controls: ATP5B, CyC1 (Primerdesign geNorm[™] Reference Gene Selection Kit) and Ppia (TaqMan assay (Mm02342429_g1) and analysed using ABI 7500 software v2.0.1 using the 2- $\Delta\Delta Ct$ method. Data is presented as mean Relative quantification (RQ) and error bars represent standard error of mean.

Western Blotting

Standard western blotting technique was used to detect secreted proteins in apical HBSS washes collected at ALI D0, D3, D7 and D14 from the mMEC cultures. An equal volume of 2x SDS loading buffer (20% SDS, 1M DTT, Glycerol, Tris-HCl 0.5M pH6.8, 0.2% bromophenol blue, protease inhibitors) was added to the wash and denatured at 95°C for 5 minutes. 40 μ L of total sample was resolved on a 12% polyacrylamide gel, transferred to a PVDF membrane using a semi dry blotting

system (Biorad Trans-blot turbo) and probed with primary antibodies (anti-Bpifa1 (1:200), anti-lacto transferrin (1:2000), and anti-REG3γ (1:5000) overnight at 4°C. The primary antibody was detected using a polyclonal goat anti-rabbit secondary antibody (Dako P0448) conjugated with HRP (1:2000). Primary antibody details are described in Table S1. Protein bands were visualised using the EZ system (Geneflow, Cat No- 30500500B).

Immunohistochemistry

Formalin fixed, paraffin embedded serial sections of the mouse head passing through the middle ear cavity were deparaffinised and dewaxed in a 100% Xylene (Sigma, UK) and rehydrated in 100% Ethanol (Fischer Scientific, UK). Endogenous peroxidase activity was blocked using 0.3 % H₂O₂-Methanol (Fischer Scientific, UK). Non-specific binding was blocked by incubating the sections in 100% goat serum for 30 minutes at room temperature. Sections were washed and incubated in anti-Bpifa1 primary antibody (1:750) overnight in a humified chamber at 4°C. The following day, sections were washed twice in PBS, incubated in 0.5% biotinylated polyclonal goat anti-rabbit secondary antibody (Vectastain^R Elite^R ABC kit, Cat No- PK-6101) for 30 minutes at room temperature, followed by incubation with the ABC reagent for signal amplification for 30 minutes. NovaRed mixture (Vector, Cat No- SK 4800) was used for colour development. Sections were counterstained using Harris's Haematoxylin (Thermo Scientific), differentiated in acid alcohol, treated with Scott's tap water, dehydrated in 95% to absolute ethanol, cleared using Xylene (Leica, Cat No- ST 4020), mounted in DPX (Leica Biosystems) and visualized and imaged using a light microscope (Olympus BX61).

Sample preparation for Mass Spectrometry

Apical wash secretions from 6 batches of ALI Day 14 mMEC cultures were pooled. TCA precipitation was performed (30% TCA in acetone) at -20°C for 2 hours to precipitate soluble proteins (50 µg). Proteins were pelleted at 12,000 g for 10 minutes (4°C) and pellets washed three times with ice-cold acetone, air dried and. re-suspended in 50 mM ammonium bicarbonate, 0.1 % RapiGest SF (waters). Samples were heated at 80°C for 10 minutes, reduced with 3 mM DTT at 60°C for 10 minutes, cooled and then alkylated with 9 mM iodoacetamide (Sigma) for 30 minutes. All steps were performed with intermittent vortexing. Proteomic-grade

trypsin (Sigma) was added at a protein: trypsin ratio of 50:1 and incubated at 37°C overnight. The samples were precipitated using 1% TFA at 37°C for 2 hr and centrifuged at 12,000 g for 1 hr (4°C) to remove RapiGest SF. The peptide supernatant was desalted using C₁₈ reverse-phase stage tips (Thermo Scientific Pierce) according to the manufacturer's instructions, dried and re-suspended in 3% (v/v) acetonitrile, 0.1% (v/v) TFA for analysis by Mass Spectrometry (MS).

NanoLC-MS ESI MS/MS analysis

Peptides were analysed by on-line nanoflow LC using the Thermo EASY-nLC 1000 LC system (Thermo Fisher Scientific) coupled with Q-Exactive mass spectrometer (Thermo Fisher Scientific). Samples were loaded onto an Easy-Spray C₁₈ column (50cm, i.d. 75 μ m), fused to a silica nano-electrospray emitter (Thermo Fisher Scientific). Chromatography was performed at 35°C with a buffer system consisting of 0.1% formic acid (buffer A) and 80% acetonitrile in 0.1% formic acid (buffer B). The peptides were separated over a 97 minute linear gradient of 3.8 – 50% buffer B at a flow rate of 300 nL/min. The Q-Exactive was operated in data-dependent mode with dynamic exclusion and survey scans acquired at a resolution of 70,000. The 10 most abundant isotope patterns with charge states +2, +3 and/or +4 from the survey scan were selected with an isolation window of 2.0Th and fragmented by higher energy collisional dissociation with normalized collision energies of 30. The maximum ion injection times for the survey scan and the MS/MS scans were 250 and 100ms, respectively, and the ion target value was set to 1E6 for survey scans and 1E4 for the MS/MS scans.

Protein Identification and Quantification

Thermo RAW files were imported into Progenesis LC–MS (version 4.1, Nonlinear Dynamics) and only peaks with a charge state between +2 and +7 were picked. Spectral data were exported for peptide identification using the Mascot (version 2.3.02, Matrix Science) search engine. Tandem MS data were searched against translated ORFs from the mouse genome (Uniprot release 2015_02; 16,868 sequences; 9,451,355 residues). The search parameters were as follows: precursor mass tolerance was set to 10 ppm and fragment mass tolerance was set to 0.8 or 0.01Da and two missed tryptic cleavages were permitted. Carbamidomethylation (cysteine) was set as a fixed modification and oxidation (methionine) set as variable

modification. Mascot search results were further validated using the machinelearning algorithm Percolator embedded within Mascot. The Mascot decoy database function was utilised and the false discovery rate was <1%, while individual percolator ion scores> 13 indicated identities or extensive homology (p <0.05). Mascot search results were imported into Progenesis LC–MS for relative quantification using non- conflicting peptides.

Statistics

A paired two tailed Students t-test was used to compare relative Cxcl2 expression between the MOCK infected and *NTHi* infected samples at each time point. Data was presented using GraphPad Prism version 6.0.

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Competing interests

The authors declare that they have no competing or financial interests.

Author contributions

A.M.and C.D.B. concieved and designed the experiments and interpreted results.

A.M. and C.D.B. wrote the paper.

A.M. performed majority of the experiments and analysed data.

K.M.A. and L.B. provided technical expertise on cell culture work.

D.W. helped with design and analysis of qPCR experiments.

H.A. performed RT-PCRs on differentiation samples and western blots for LTF and REG3γ.

C.R. performed IHC for Bpifa1 on Wt middle ear sections.

S.A. and J.P.S. performed and analysed the proteomics data.

D.H. provided the NTHi strain and helped concieve infection experiments.

K.M.A., S.D.M.B. and L.B. assisted critical evaluation and drafting of the manuscript.

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Figures



Fig. 1: Primary culture of mouse middle ear epithelial cells. Timeline for culture of mMECs is shown above (A). Bullae were dissected, treated with pronase for dissociation of the middle ear epithelial cells and fibroblasts were excluded from culture by differential adherence to plastic. Epithelial cells were grown in submerged culture till confluence, before ALI was induced. Samples for transcriptional and proteomic analysis were collected at regular time points. Phase contrast images showing cells in culture under 10x magnification (B-I). In the proliferative submerged conditions, a small number of cells attached to form epithelial islands 3 days after seeding (B) The cells proliferated faster from day 5 (C) through day 7 (D) and formed a confluent monolayer at day 9. This was termed as ALI Day 0 (E) Morphology of cells changed from ALI Day 3 (F) and clusters of compactly arranged cells started forming at ALI Day 7 (G). ALI Day 14 cultures were composed of flat polygonal and compactly clustered pseudo stratified cells with active cilia. White arrowheads mark

elevated ciliated cells and asterisk mark flatter polygonal cells. (H). Fibroblasts cultured on plastic plates through differential adhesion method (I) Scale bar = 200 μ m.



Fig. 2: Electron microscopy of mMEC cultures. Scanning electron microscopy of ALI Day 0 mMEC cultures showing large flat polygonal cells with apical microvilli (A) ALI Day 14 cultures showing dome shaped cells at higher magnification (B) and combination of interspersed flat polygonal and densely ciliated cell populations a lower magnification (C) resembling the morphology of native middle ear epithelium (D). Cracks developed in the membrane are due to processing of samples for SEM. White arrowheads mark elevated ciliated cells and asterix mark flatter polygonal cells. Transmission electron microscopy of ALI Day 14 mMEC cultures showing adjacent ciliated and secretory cells and formation of tight junctions demonstrated by presence of desmosomes (asterisk). Arrow shows cilia (E). Immunoconfocal microscopy image showing formation of tight junctions marked by ZO1 positive staining. Area beyond white lines indicates a cross section through the membrane. (n=3) Scale bar =10 μ m (A,C,D) ; 5 μ m (B); 1 μ m (E); 50 μ m (F).



Fig. 3: Expression of epithelial markers in mMEC cultures. End-point RT-PCR showing expression of a selected panel of upper airway associated genes in mMEC original cells isolated from the middle ear cavity, ALI D0 cells and ALI D14 cells. Expression profile of ALI Day 14 cells was similar to mMEC original cells isolated from the middle ear for most genes (A). Detection of Bpifa1 (B), Lactotransferrin (C) and Reg3Y (D) in the apical washes from differentiating cells using Western blotting technique. Data is representative of three independent cultures.



epithelial Fig. 4: Localisation of markers in mMEC cultures. Immunofluorescence confocal images (representative of three independent batches) showing abundant expression of the basal cell marker, P63; limited expression of the secretory protein, Bpifa1 (A), no expression of goblet cell marker, Muc5B and the ciliated marker, FoxJ1 (C) in undifferentiated ALI Day 0 mMEC cultures (A and C). Differentiated mMEC ALI Day 14 (B, D, E, F) cultures showing expression of secretory cells positive for Bpifa1 (B), Lactotransferrin (E), Reg3y (F), goblet cells positive for Muc5B and ciliated cells positive for FoxJ1 (D). Area beyond white lines indicates a cross section through the membrane. High magnification. Z-stack cross sections of nuclei stained with DAPI shows that ALI Day 0 cells form a flat monolayer (G) whereas ALI Day 14 cells are a combination of pseudostratified elevated cells (arrowheads) and flatter cells (asterisk) (H). Scale bar =50µm.



Fig. 5: Distribution of cell types in mMEC cultures. Low powered (20x) immunofluorescence confocal images representative of 3 independent batches of ALI Day 14 mMEC cultures showing a combination of flat cells and clusters of elevated cells at slightly different foci. Ciliated cells (FoxJ1) do not co-localise with and goblet cells (Muc5B) (A) and Bpifa1 expressing cells (B) Scale bar =100µm. High magnification (60x) cross section Z-stack confocal images showing that FoxJ1 and Muc5B expression is mostly restricted to the elevated cell types (arrowheads) (C) and Bpifa1 is predominantly expressed by flatter cells (asterisk) (D) Scale bar = $50\mu m$ (C,D).



Fig. 6: mMEC cultures serve as an otopathogenic infection model. High magnification (60x) confocal images representative of n=3 independent batches of ALI Day 14 mMECs infected with GFP tagged NTHi-375SR for 24 hours (A), 48 hours (B) and 72 hours (C) showing a time dependent increase in bacterial infection. Control ALI Day 14 mMECs mock infected for 72 hrs (D) Scale bar =50µm. Increase in mean green fluorescence intensity from 24hpi to 72hpi quantified from lower images from three independent batches, magnification (10x) suggesting demonstrating an increase is the amount of bacteria infecting the cultures with time (E). Relative gene expression of Cxcl2 (F) in infected mMEC cultures compared with MOCK infected cultures at 24 and 48 hours post infection. Expression of target gene Cxcl2 was normalized to 3 endogenous controls: ATP5B, CyC1 and Ppia and plotted relative to the 24hpi MOCK sample. Data is analysed using two tailed student's ttest and represented as mean relative quantification (RQ) ± SEM for three independent batches of cultures. *: p<0.05 **p<0.01 ***p<0.005.

Accession Number	Protein	Peptide count	emPAI Score	Biological reference
P08071	Lactotransferrin	47	202.53	Anti-microbial iron chelation
Q921I1	Serotransferrin	40	29.58	Iron chelation, cell proliferation
O09049	Regenerating islet-derived protein 3-gamma	8	11.38	Anti-microbial
P11672	Neutrophil gelatinase- associated lipocalin	9	10.45	Iron trafficking, innate immunity
Q61147	Ceruloplasmin	55	10.1	Copper transport, antioxidant defence
P97361	A member 1	9	4.74	Suggested role in innate immunity
P01027	Complement C3	111	4.68	Activates complement system
P06797	Cathepsin L1	12	3.11	Lysosomal protein degradation
Q61805	Lipopolysaccharide- binding protein	19	2.96	Antimicrobial activity through bacterial LPS binding
P10605	Cathepsin B	15	2.47	Intracellular protein degradation and turnover
Q61362	Chitinase-3-like protein 1	22	2.16	Tissue remodelling in response to environmental stress, activation of NfkB signalling pathway
P25785	Metalloproteinase inhibitor 2	14	2.08	Inactivates metalloenzymes
Q9CQV3	Serpin B11	24	1.97	Serine protease inhibitor
P18242	Cathepsin D	20	1.67	Intracellular protein breakdown
O88312	Anterior gradient protein 2 homolog	10	1.53	Regulates giblet cell differentiation and mucous secretion.
P50404	Pulmonary surfactant- associated protein D	23	1.5	Innate immunity response, respiratory gaseous exchange and homeostasis
P10810	Monocyte differentiation antigen CD14	16	1.41	TLR2 mediated innate immune and inflammatory response to bacterial LPS in concert with LBP
Q62426	Cystatin-B	6	1.24	Proteinase inhibitor
Q9ER10	Brain-specific serine protease 4	15	1.11	Protease activity
P08905	Lysozyme	10	1.05	Bacteriolytic activity
Q9DAU7	WAP four-disulfide core domain protein 2	6	0.91	Protease inhibitor
P11214	Tissue-type plasminogen activator	31	0.89	Tissue remodelling, cell migration
Q9WUU7	Cathepsin Z	14	0.89	Peptidase activity
O88593	Peptidoglycan recognition protein 1	10	0.81	Triggers apoptosis of Gram positive bacteria
Q60854	Serpin B6	23	0.81	Regulation of serine proteases
P34884	Macrophage migration inhibitory factor	5	0.61	Pro inflammatory cytokine involved in regulation o macrophage activity
Q9R118	Serine protease HTRA1	21	0.54	Cell proliferation and viral response

Table 1: Most abundant secreted proteins identified in apical ALI Day 14 washes from mMEC cultures
P12032	Metalloproteinase inhibitor 1	12	0.49	Cell differentiation and wound healing
P97430	Antileukoproteinase	11	0.49	Anti microbial host defence
P61939	Thyroxine-binding globulin	22	0.4	Thyroid hormone transport
P25085	Interleukin-1 receptor antagonist protein	7	0.35	Inhibits IL-1 activity
P70124	Serpin B5	22	0.35	Morphogenesis of epithelium
Q61703	Inter-alpha-trypsin inhibitor heavy chain H2	48	0.24	Hyaluronan metabolic process
P97290	Plasma protease C1 inhibitor	25	0.12	Blood coagulation, complement activation.

NB- Acession numbers and biological functions from the UniProt database

Supplementary Data for

An in vitro model of murine middle ear epithelium

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Fig. S1: Expression of Bpifa1 in WT middle ear epithelium

Bpifa1 expression was detected in the Wt mouse middle ear using immunohistochemistry. Low magnification image showing expression of Bpifa1 all along the middle ear epithelium (A). Bpifa1 is secreted by non-ciliated cells of the middle ear epithelium (B), but it coats the surface of the cilia (C). Negative controls showed absence of staining (D, E).



Fig. S2: Progression of NTHi infection in mMEC cultures

Low power confocal images of ALI Day 14 mMEC cells infected with GFP tagged NTHI-375^{SR} at 24 hpi (A), 48hpi (B) and 72 hpi (C). Each panel represents 4 central fields at 10x magnification spanning approximately 50% of the membrane. The amount of infection at each time point was quantified as a measure of the mean integrated intensity for green fluorescence in these panels from 3 independent batches of infections. Scale bar = $200 \,\mu\text{m}$



Fig. S3: Optimisation of seeding density for mMEC culture

Phase contrast images of ALI Day 14 mMEC cells in culture seeded at various densities in absence and presence of ROCKi (A-G). On plastic, both in absence (A) and presence (E) of ROCKi, cells did not exhibit the typical epithelial morphology, started forming vacuoles, detached from the surface and did not achieve confluence. In absence of ROCKi, at 1 x10⁴ cells/ well (B) cells grew in small epithelial clusters but did not form a confluent monolayer, at 2 x10⁴ cells/ well (C) a proportion of wells formed a confluent monolayer at submerged Day 15 and at 5 x10⁴ cells (D), cells formed a confluent monolayer around submerged Day 9. The cells differentiated on ALI and demonstrated a typical cobbled stone appearance. In presence of ROCKi, both at 1 x10⁴ cells/ well (F) and 2 x10⁴ cells/ well (G), the cells formed a confluent monolayer in submerged culture by day 9-10. By ALI Day 14, the cells differentiated into various cell types and exhibited cobbled stone appearance typical of epithelial cells. Scale bar = 200 μ m. White arrowheads indicate elevated clusters of cells with actively beating cilia and asterisk denote flatter polygonal cells. RT PCR of ALI Day 14 samples (H) demonstrating that adding ROCKi to the culture medium did not affect differentiation into secretory and ciliated cell types. Seeding density of 1 x10⁴

cells/ well with ROCKi was selected in order to maximize the utility of the isolated cells.



Fig. S4: Distribution of ciliated cells in WT native mouse middle ear cavity

Scanning electron microscopy images showing the distribution of ciliated cells in various parts of the native WT middle ear cavity. Areas near the Eustachian tube were lined with tracts of dense ciliated cells (A). The central part of the middle consists of flat polygonal cells interspersed with clusters of short and long ciliated cells (B), similar to what is seen in our mMEC cultures. Areas towards the tympanic membrane exhibit patches similar to central part of the middle ear as well as larger patches of flat polygonal cells with apical microvilli (C). Scale bar = 10 μ m



Fig. S5: Dissection and isolation of the mouse middle ear cavity

Wild type C57BI/6J or C3HeH mice (A) were decapitated (B). The head was skinned (C) and skull cap removed (D). Dorsal (E) and ventral (F) view of the head after removing the brain, showing the bullae or middle ear cavities, MECs. Bisected head showing the outer ear, OE (G). Under dissecting microscope the bullae were separated from surrounding tissue (H). The MEC is attached to the outer ear cavity (OEC) at the tympanic membrane (Tm) and tapers towards the opening of the Eustachian tube (ET) near its posterior end (I). Removal of the OEC and the Tm reveals the cochlea of the inner ear on the ventral side of the MEC (J) The cup shaped MEC was detached from the inner ear (K) Tissue was dissected along the dotted lines.



Fig. S6: Effect of Pronase treatment on the middle ear epithelial cells

Haematoxylin and eosin staining of dissected middle ear cavities showing intact middle ear epithelium before proteolytic treatment with pronase (A) and loss of the epithelial lining after treatment with pronase and mechanical agitation (B) during the cell isolation process suggesting that our isolation protocol successfully dissociates middle ear epithelial cells for culture. Scale bar = 200µm



Movie S1: Time lapse images showing ciliary beating in live ALI Day 14 mMEC cells

Table S1: Proteins identified by MS analysis of ALI Day 14 mMEC apical washes

Accession Number	Protein Name	Gene name	Peptide count	emPAI
P08071	Lactotransferrin	TRFL_MOUSE	47	202.53
Q921I1	Serotransferrin	TRFE_MOUSE	40	29.58
O09131	Glutathione S-transferase omega-1	GSTO1_MOUSE	17	28.64
P60710	Actin, cytoplasmic 1	ACTB_MOUSE	23	24.63
P50446	Keratin, type II cytoskeletal 6A	K2C6A_MOUSE	35	19.14
P08074	[NADPH] 2	CBR2_MOUSE	16	19.04
Q9Z331	Keratin, type II cytoskeletal 6B	K2C6B_MOUSE	36	18.18
P62806	Histone H4	H4_MOUSE	6	17.1
P10107	Annexin A1	ANXA1_MOUSE	24	14.87
P11679	Keratin, type II cytoskeletal 8	K2C8_MOUSE	33	14.67
P19001	Keratin, type I cytoskeletal 19	K1C19_MOUSE	35	14.07
Q61781	Keratin, type I cytoskeletal 14	K1C14_MOUSE	35	13
O09049	Regenerating islet-derived protein 3-gamma	REG3G_MOUSE	8	11.38
P35700	Peroxiredoxin-1	PRDX1_MOUSE	14	11.13
Q06890	Clusterin	CLUS_MOUSE	22	10.47
P11672	Neutrophil gelatinase- associated lipocalin	NGAL_MOUSE	9	10.45
Q61147	Ceruloplasmin	CERU_MOUSE	55	10.1
Q61414	Keratin, type I cytoskeletal 15	K1C15_MOUSE	32	9.92
P17742	Peptidyl-prolyl cis-trans isomerase A	PPIA_MOUSE	14	9.81
Q922U2	Keratin, type II cytoskeletal 5	K2C5_MOUSE	34	8.7
Q9QWL7	Keratin, type I cytoskeletal	K1C17_MOUSE	33	7.76
O08709	Peroxiredoxin-6	PRDX6_MOUSE	15	7.38
P10649	Glutathione S-transferase Mu 1	GSTM1_MOUSE	18	6.68
P07356	Annexin A2	ANXA2_MOUSE	23	6.63
P52480	Pyruvate kinase PKM	KPYM_MOUSE	36	6.57
P68372	Tubulin beta-4B chain	TBB4B_MOUSE	21	6.13
Q6ZWY9	Histone H2B type 1-C/E/G	H2B1C_MOUSE	11	6.12
Q8BND5	Sulfhydryl oxidase 1	QSOX1_MOUSE	43	6.1
Q9DCV7	Keratin, type II cytoskeletal 7	K2C7_MOUSE	35	6.01
P16858	Glyceraldehyde-3- phosphate dehydrogenase	G3P_MOUSE	21	5.87
Q61398	Procollagen C-	PCOC1_MOUSE	23	5.56

	endopeptidase enhancer 1			
	Keratin, type II cytoskeletal			
P07744	4	K2C4_MOUSE	31	5.43
O35639	Annexin A3	ANXA3_MOUSE	24	5.16
P68033	1	ACTC_MOUSE	23	5.06
P26040	Ezrin	EZRI_MOUSE	39	5.04
P99024	Tubulin beta-5 chain	TBB5_MOUSE	21	4.94
Q05816	Fatty acid-binding protein, epidermal	FABP5_MOUSE	9	4.92
P08240	Malate dehydrogenase,		24	4 77
1 00245	BPI fold-containing family A			
P97361	member 1	BPIA1_MOUSE	9	4.74
P01027	Complement C3	CO3_MOUSE	111	4.68
Q8BGZ7	Keratin, type II cytoskeletal	K2C75_MOUSE	36	4.22
P56480	ATP synthase subunit beta, mitochondrial	ATPR MOUSE	30	4 18
P62830	60S ribosomal protein L23	RI 23 MOUSE	10	4.08
1 02030	Keratin, type I cytoskeletal			4.00
P05784	18	K1C18_MOUSE	32	3.99
P50543	Protein S100-A11	S10AB_MOUSE	5	3.92
P28654	Decorin	PGS2_MOUSE	23	3.86
P40142	Transketolase	TKT_MOUSE	34	3.7
Q9QUI0	Transforming protein RhoA	RHOA_MOUSE	13	3.7
Q8VCT4	Carboxylesterase 1D	CES1D_MOUSE	27	3.69
Q9Z2K1	Keratin, type I cytoskeletal	K1C16_MOUSE	35	3.64
P57780	Alpha-actinin-4	ACTN4_MOUSE	61	3.61
P17563	Selenium-binding protein 1	SBP1_MOUSE	31	3.49
P17182	Alpha-enolase	ENOA_MOUSE	28	3.36
P17182 P11499	Alpha-enolase Heat shock protein HSP 90-beta	HS90B_MOUSE	28 40	3.36 3.31
P17182 P11499	Alpha-enolase Heat shock protein HSP 90-beta Keratin, type I cytoskeletal	HS90B_MOUSE	28 40 31	3.36 3.31
P17182 P11499 P08730 P14211	Alpha-enolase Heat shock protein HSP 90-beta Keratin, type I cytoskeletal 13	HS90B_MOUSE K1C13_MOUSE	28 40 31 27	3.36 3.31 3.29 3.28
P17182 P11499 P08730 P14211 O9D379	Alpha-enolase Heat shock protein HSP 90-beta Keratin, type I cytoskeletal 13 Calreticulin Epovide bydrolase 1	HS90B_MOUSE K1C13_MOUSE CALR_MOUSE	28 40 31 27 29	3.36 3.31 3.29 3.28 3.27
P17182 P11499 P08730 P14211 Q9D379	Alpha-enolase Heat shock protein HSP 90-beta Keratin, type I cytoskeletal 13 Calreticulin Epoxide hydrolase 1 Heat shock cognate 71 kDa	HS90B_MOUSE K1C13_MOUSE CALR_MOUSE HYEP_MOUSE	28 40 31 27 29 27	3.36 3.31 3.29 3.28 3.27 2.24
P17182 P11499 P08730 P14211 Q9D379 P63017	Alpha-enolase Heat shock protein HSP 90-beta Keratin, type I cytoskeletal 13 Calreticulin Epoxide hydrolase 1 Heat shock cognate 71 kDa protein	HS90B_MOUSE K1C13_MOUSE CALR_MOUSE HYEP_MOUSE HSP7C_MOUSE	28 40 31 27 29 37 29	3.36 3.31 3.29 3.28 3.27 3.24 2.47
P17182 P11499 P08730 P14211 Q9D379 P63017 Q00493 P00707	Alpha-enolase Heat shock protein HSP 90-beta Keratin, type I cytoskeletal 13 Calreticulin Epoxide hydrolase 1 Heat shock cognate 71 kDa protein Carboxypeptidase E	HS90B_MOUSE K1C13_MOUSE CALR_MOUSE HYEP_MOUSE HSP7C_MOUSE CBPE_MOUSE	28 40 31 27 29 37 28 42	3.36 3.31 3.29 3.28 3.27 3.24 3.17
P17182 P11499 P08730 P14211 Q9D379 P63017 Q00493 P06797	Alpha-enolase Heat shock protein HSP 90-beta Keratin, type I cytoskeletal 13 Calreticulin Epoxide hydrolase 1 Heat shock cognate 71 kDa protein Carboxypeptidase E Cathepsin L1	HS90B_MOUSE HS90B_MOUSE K1C13_MOUSE CALR_MOUSE HYEP_MOUSE HSP7C_MOUSE CBPE_MOUSE CATL1_MOUSE DD054_MOUSE	28 40 31 27 29 37 28 12 2	3.36 3.31 3.29 3.28 3.27 3.24 3.17 3.11
P17182 P11499 P08730 P14211 Q9D379 P63017 Q00493 P06797 P62962	Alpha-enolase Heat shock protein HSP 90-beta Keratin, type I cytoskeletal 13 Calreticulin Epoxide hydrolase 1 Heat shock cognate 71 kDa protein Carboxypeptidase E Cathepsin L1 Profilin-1 Protein disulfide-isomerase	ENOA_MOUSE HS90B_MOUSE K1C13_MOUSE CALR_MOUSE HYEP_MOUSE HSP7C_MOUSE CBPE_MOUSE CATL1_MOUSE PROF1_MOUSE	28 40 31 27 29 37 28 12 9	3.36 3.31 3.29 3.28 3.27 3.24 3.17 3.11 3.1
P17182 P11499 P08730 P14211 Q9D379 P63017 Q00493 P06797 P62962 P27773	Alpha-enolaseHeat shock protein HSP90-betaKeratin, type I cytoskeletal13CalreticulinEpoxide hydrolase 1Heat shock cognate 71 kDaproteinCarboxypeptidase ECathepsin L1Profilin-1Protein disulfide-isomeraseA3	HS90B_MOUSE K1C13_MOUSE CALR_MOUSE HYEP_MOUSE HSP7C_MOUSE CBPE_MOUSE CATL1_MOUSE PROF1_MOUSE PDIA3_MOUSE	28 40 31 27 29 37 28 12 9 35	3.36 3.31 3.29 3.28 3.27 3.24 3.17 3.11 3.1 3.05
P17182 P11499 P08730 P14211 Q9D379 P63017 Q00493 P06797 P62962 P27773 P62908	Alpha-enolaseHeat shock protein HSP90-betaKeratin, type I cytoskeletal13CalreticulinEpoxide hydrolase 1Heat shock cognate 71 kDaproteinCarboxypeptidase ECathepsin L1Profilin-1Protein disulfide-isomeraseA340S ribosomal protein S3	ENOA_MOUSE HS90B_MOUSE K1C13_MOUSE CALR_MOUSE HYEP_MOUSE HSP7C_MOUSE CBPE_MOUSE CATL1_MOUSE PROF1_MOUSE PDIA3_MOUSE RS3_MOUSE	28 40 31 27 29 37 28 12 9 35 19	3.36 3.31 3.29 3.28 3.27 3.24 3.17 3.11 3.1 3.05 3.05
P17182 P11499 P08730 P14211 Q9D379 P63017 Q00493 P06797 P62962 P27773 P62908 Q7TMM9	Alpha-enolaseHeat shock protein HSP90-betaKeratin, type I cytoskeletal13CalreticulinEpoxide hydrolase 1Heat shock cognate 71 kDaproteinCarboxypeptidase ECathepsin L1Profilin-1Protein disulfide-isomeraseA340S ribosomal protein S3Tubulin beta-2A chain	ENOA_MOUSE HS90B_MOUSE K1C13_MOUSE CALR_MOUSE HYEP_MOUSE HSP7C_MOUSE CBPE_MOUSE CATL1_MOUSE PROF1_MOUSE PDIA3_MOUSE TBB2A_MOUSE	28 40 31 27 29 37 28 12 9 35 19 21	3.36 3.31 3.29 3.28 3.27 3.24 3.17 3.11 3.1 3.05 3.05 3.03
P17182 P11499 P08730 P14211 Q9D379 P63017 Q00493 P06797 P62962 P27773 P62908 Q7TMM9 Q8VDD5	Alpha-enolaseHeat shock protein HSP90-betaKeratin, type I cytoskeletal13CalreticulinEpoxide hydrolase 1Heat shock cognate 71 kDaproteinCarboxypeptidase ECathepsin L1Profilin-1Protein disulfide-isomeraseA340S ribosomal protein S3Tubulin beta-2A chainMyosin-9	ENOA_MOUSE HS90B_MOUSE K1C13_MOUSE CALR_MOUSE HYEP_MOUSE HSP7C_MOUSE CBPE_MOUSE CATL1_MOUSE PROF1_MOUSE PDIA3_MOUSE RS3_MOUSE TBB2A_MOUSE MYH9_MOUSE	28 40 31 27 29 37 28 12 9 35 19 21 127	3.36 3.31 3.29 3.28 3.27 3.24 3.17 3.11 3.1 3.05 3.05 3.03
P17182 P11499 P08730 P14211 Q9D379 P63017 Q00493 P06797 P62962 P27773 P62908 Q7TMM9 Q8VDD5 P09103	Alpha-enolaseHeat shock protein HSP90-betaKeratin, type I cytoskeletal13CalreticulinEpoxide hydrolase 1Heat shock cognate 71 kDaproteinCarboxypeptidase ECathepsin L1Profilin-1Protein disulfide-isomeraseA340S ribosomal protein S3Tubulin beta-2A chainMyosin-9Protein disulfide-isomerase	ENOA_MOUSE HS90B_MOUSE K1C13_MOUSE CALR_MOUSE HYEP_MOUSE HYEP_MOUSE CBPE_MOUSE CATL1_MOUSE PROF1_MOUSE PDIA3_MOUSE TBB2A_MOUSE MYH9_MOUSE PDIA1_MOUSE	28 40 31 27 29 37 28 12 9 35 12 9 35 19 21 127 39	3.36 3.31 3.29 3.28 3.27 3.24 3.17 3.11 3.11 3.11 3.05 3.05 3.03 3.03 3.02
P17182 P11499 P08730 P14211 Q9D379 P63017 Q00493 P06797 P62962 P27773 P62908 Q7TMM9 Q8VDD5 P09103 P68368	Alpha-enolaseHeat shock protein HSP90-betaKeratin, type I cytoskeletal13CalreticulinEpoxide hydrolase 1Heat shock cognate 71 kDaproteinCarboxypeptidase ECathepsin L1Profilin-1Protein disulfide-isomeraseA340S ribosomal protein S3Tubulin beta-2A chainMyosin-9Protein disulfide-isomeraseTubulin alpha-4A chain	ENOA_MOUSE HS90B_MOUSE K1C13_MOUSE CALR_MOUSE HYEP_MOUSE HYEP_MOUSE CBPE_MOUSE CATL1_MOUSE PROF1_MOUSE PDIA3_MOUSE RS3_MOUSE TBB2A_MOUSE MYH9_MOUSE PDIA1_MOUSE TBA4A_MOUSE	28 40 31 27 29 37 28 12 9 35 19 21 127 39 22	3.36 3.31 3.29 3.28 3.27 3.24 3.17 3.11 3.1 3.05 3.05 3.03 3.02 2.99

Q8VED5	Keratin, type II cytoskeletal 79	K2C79 MOUSE	32	2.98
Q61805	Lipopolysaccharide-binding protein	LBP_MOUSE	19	2.96
Q9Z1Q5	Chloride intracellular channel protein 1	CLIC1_MOUSE	17	2.96
Q07797	Galectin-3-binding protein	LG3BP_MOUSE	26	2.94
Q61646	Haptoglobin	HPT_MOUSE	20	2.94
Q7TPR4	Alpha-actinin-1	ACTN1_MOUSE	60	2.92
P62821	Ras-related protein Rab-1A	RAB1A_MOUSE	18	2.9
O88844	Isocitrate dehydrogenase [NADP] cytoplasmic	IDHC_MOUSE	30	2.87
P08228	Superoxide dismutase [Cu-	SODC MOUSE	11	2.78
P18760	Cofilin-1	COF1 MOUSE	12	2.72
	Glutathione S-transferase			
P19157	P 1	GSTP1_MOUSE	10	2.71
Q9R0P5	Destrin	DEST_MOUSE	13	2.7
P06151	L-lactate dehydrogenase A chain	LDHA_MOUSE	23	2.62
P20029	78 kDa glucose-regulated	GRP78 MOUSE	30	2.61
P10126	Flongation factor 1-alpha 1	EE1A1 MOUSE	23	2.54
060605	Myosin light polypeptide 6		11	2.54
062159	Rho-related GTP-binding	RHOC MOUSE	11	2.5
09D1G1	Ras-related protein Rah-1R	RAB1B MOUSE	17	2.5
P07901	Heat shock protein HSP 90-alpha	HS90A MOUSE	39	2.47
P10605	Cathepsin B	CATB MOUSE	15	2.47
P68040	Guanine nucleotide-binding protein subunit beta-2-like	GBLP MOUSE	22	2.47
P97429	Annexin A4	ANXA4 MOUSE	22	24
Q64669	NAD(P)H dehydrogenase [quinone] 1	NQO1 MOUSE	15	2.38
O54974	Galectin-7	LEG7 MOUSE	10	2.33
P33267	Cytochrome P450 2F2	CP2F2 MOUSE	29	2.3
P47739	Aldehyde dehydrogenase, dimeric NADP-preferring	AL3A1_MOUSE	25	2.29
Q61508	Extracellular matrix protein	ECM1_MOUSE	37	2.29
Q9R0P3	S-formylglutathione hydrolase	ESTD_MOUSE	16	2.28
P47738	Aldehyde dehydrogenase, mitochondrial	ALDH2_MOUSE	26	2.24
P17751	I riosephosphate isomerase	TPIS_MOUSE	18	2.19
P63038	bu KDa heat shock protein, mitochondrial	CH60_MOUSE	36	2.16
Q61362	Chitinase-3-like protein 1	CH3L1_MOUSE	22	2.16
P25785	Metalloproteinase inhibitor 2	TIMP2_MOUSE	14	2.08
Q8BFZ3	Beta-actin-like protein 2	ACTBL_MOUSE	24	2.08

00000444	Electron transfer			2.00
	navoprotein subunit beta		14	2.08
P63101	14-3-3 protein zeta/delta	1433Z_MOUSE	19	2.07
Q03265	alpha, mitochondrial	ATPA_MOUSE	36	2.07
Q62266	Cornifin-A	SPR1A_MOUSE	13	2.01
P21956	Lactadherin	MFGM_MOUSE	23	1.99
P15626	Glutathione S-transferase	GSTM2 MOUSE	18	1 07
00001/3	Sorpin B11	SDR11 MOUSE	24	1.97
Q9CQV3	Fukarvotic translation	SPBIT_WOUSE	24	1.97
P63242	initiation factor 5A-1	IF5A1_MOUSE	8	1.93
P11276	Fibronectin	FINC_MOUSE	108	1.84
P51881	ADP/ATP translocase 2	ADT2_MOUSE	21	1.84
P05064	Fructose-bisphosphate		23	1.82
F 03004	Transitional endoplasmic	ALDOA_INOUSL	23	1.02
Q01853	reticulum ATPase	TERA_MOUSE	45	1.82
	Phosphatidylethanolamine-			
P70296	binding protein 1	PEBP1_MOUSE	9	1.81
P21981	glutamyltransferase 2	TGM2 MOUSE	36	1.79
P26043	Radixin	RADI MOUSE	39	1.79
	40S ribosomal protein			
P62245	S15a	RS15A_MOUSE	11	1.76
Q6IFX2	Keratin, type I cytoskeletal	K1C42_MOUSE	33	1.75
	6-phosphogluconate			
000000	dehydrogenase,	ADCD MOUSE	20	1 74
Q9DCD0	Ervthrocyte band 7 integral		23	1.74
P54116	membrane protein	STOM_MOUSE	17	1.71
P26041	Moesin	MOES_MOUSE	40	1.7
P18242	Cathepsin D	CATD_MOUSE	20	1.67
Q923D2	Flavin reductase (NADPH)	BLVRB_MOUSE	13	1.65
Q9R0Q7	Prostaglandin E synthase 3	TEBP_MOUSE	9	1.65
	Purine nucleoside			
P23492	phosphorylase	PNPH_MOUSE	19	1.64
Q60932	selective channel protein 1	VDAC1 MOUSE	18	1.64
	Pyruvate dehydrogenase			
	E1 component subunit			
Q9D051	beta, mitochondrial	ODPB_MOUSE	18	1.63
Q9WVA4	Transgelin-2	TAGL2_MOUSE	14	1.63
P37040	reductase	NCPR MOUSE	35	1.6
P61027	Ras-related protein Rab-10	RAB10 MOUSE	17	1.6
P67778	Prohibitin	PHB MOUSE	18	1.58
	ATP synthase subunit			
P56382	epsilon, mitochondrial	ATP5E_MOUSE	4	1.56
088312	Anterior gradient protein 2	AGR2 MOUSE	10	1.53
	Peptidyl-prolyl cis-trans			
P24369	isomerase B	PPIB_MOUSE	15	1.51

P50404	Pulmonary surfactant- associated protein D	SETPD MOUSE	23	1.5
P/8678	Prelamin-A/C	LMNA MOUSE	16	1.0
P61750			12	1.47
P01750	ADF-IDOSVIATION TACION 4	ARF4_MOUSE	12	1.47
	Arginase-1	ARGIT_MOUSE	18	1.47
Q8BSL/	ADP-ribosylation factor 2	ARF2_MOUSE	12	1.45
P62827	protein Ran	RAN_MOUSE	11	1.44
P10810	antigen CD14	CD14_MOUSE	16	1.41
Q9CBB3	5-nydroxyisourate	HIUH MOUSE	7	1 41
	Rab GDP dissociation			
Q61598	inhibitor beta	GDIB_MOUSE	28	1.4
P24549	Retinal dehydrogenase 1	AL1A1_MOUSE	23	1.39
D00047	Stress-70 protein,	ODDZE MOUIOE	40	4.00
P38647	mitochondriai	GRP/5_MOUSE	43	1.39
Q61468	Mesothelin Chlorido introcollulor	MSLN_MOUSE	29	1.38
Q9QYB1	channel protein 4	CLIC4 MOUSE	17	1.38
dodibi	Glutathione S-transferase			1.00
P24472	A4	GSTA4_MOUSE	12	1.36
Q8BFU2	Histone H2A type 3	H2A3_MOUSE	7	1.36
O35129	Prohibitin-2	PHB2_MOUSE	26	1.35
000500	Heterogeneous nuclear		4.0	1.00
088569	ribonucleoproteins A2/B1	ROA2_MOUSE	19	1.33
P47911	60S ribosomal protein L6	RL6_MOUSE	19	1.33
Q9CPV4	containing protein 4	GLOD4 MOUSE	22	1.33
Q9WV54	Acid ceramidase	ASAH1 MOUSE	21	1.33
	EGF-containing fibulin-like			
	extracellular matrix protein		~ 7	
Q8BPB5	1 Sodium/potossium	FBLN3_MOUSE	27	1.31
	transporting ATPase			
Q8VDN2	subunit alpha-1	AT1A1_MOUSE	53	1.27
P97461	40S ribosomal protein S5	RS5_MOUSE	10	1.25
	Voltage-dependent anion-			
Q60931	selective channel protein 3	VDAC3_MOUSE	17	1.25
064433	TU KDa neat shock protein,	CH10 MOUSE	10	1 25
062426	Cystatin-B	CYTB MOUSE	6	1.20
Q02420	Electron transfer		0	1.24
	flavoprotein subunit alpha,			
Q99LC5	mitochondrial	ETFA_MOUSE	23	1.24
D47757	F-actin-capping protein		20	1 22
F4//5/	Long-chain specific acyl-	CAPZD_IVIOUSE	20	1.22
	CoA dehydrogenase,			
P51174	mitochondrial	ACADL_MOUSE	24	1.21
P08207	Protein S100-A10	S10AA_MOUSE	5	1.2
P84228	Histone H3.2	H32_MOUSE	6	1.2
Q9CR57	60S ribosomal protein L14	RL14_MOUSE	12	1.2
P09411	Phosphoglycerate kinase 1	PGK1 MOUSE	29	1.18
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O6IEZ6	Keratin, type II cytoskeletal	K2C1B MOUSE	30	1 18
P62267	40S ribosomal protein S23	RS23 MOUSE	8	1.10
1 02207	ATP synthase subunit	1025_WOUGE	0	1.15
Q91VR2	gamma, mitochondrial	ATPG_MOUSE	17	1.15
Q9CQI6	Coactosin-like protein	COTL1_MOUSE	10	1.14
Q9D8N0	Elongation factor 1-gamma	EF1G_MOUSE	27	1.14
P10639	Thioredoxin	THIO_MOUSE	7	1.12
P84084	ADP-ribosylation factor 5	ARF5_MOUSE	13	1.12
P62264	40S ribosomal protein S14	RS14_MOUSE	6	1.11
Q9ER10	Brain-specific serine protease 4	BSSP4_MOUSE	15	1.11
Q91VI7	Ribonuclease inhibitor	RINI_MOUSE	27	1.1
P62702	40S ribosomal protein S4, X isoform	RS4X_MOUSE	20	1.09
P62717	60S ribosomal protein L18a	RL18A_MOUSE	14	1.09
P97351	40S ribosomal protein S3a	RS3A_MOUSE	19	1.08
Q99JY9	Actin-related protein 3	ARP3_MOUSE	21	1.08
P17879	Heat shock 70 kDa protein 1B	HS71B_MOUSE	37	1.07
P60766	Cell division control protein 42 homolog	CDC42_MOUSE	9	1.06
O88342	WD repeat-containing protein 1	WDR1_MOUSE	37	1.05
P08905	Lysozyme C-2	LYZ2_MOUSE	10	1.05
	Ornithine			
P29758	aminotransferase, mitochondrial	OAT_MOUSE	23	1.05
P15532	Nucleoside diphosphate kinase A	NDKA_MOUSE	12	1.03
P48036	Annexin A5	ANXA5_MOUSE	25	1.03
P51410	60S ribosomal protein L9	RL9_MOUSE	11	1.03
Q61171	Peroxiredoxin-2	PRDX2_MOUSE	10	1.03
Q01768	Nucleoside diphosphate kinase B	NDKB_MOUSE	12	1.02
Q8K354	Carbonyl reductase	CBR3 MOUSE	20	1 02
Q91XV3	Brain acid soluble protein 1	BASP1 MOUSE	10	1.02
P62889	60S ribosomal protein L30	RI 30 MOUSE	8	1.01
P99029	Peroxiredoxin-5, mitochondrial	PRDX5_MOUSE	15	1.01
Q00612	Glucose-6-phosphate 1- dehydrogenase X	G6PD1_MOUSE	35	1.01
P14152	Malate dehydrogenase, cytoplasmic	MDHC MOUSE	21	1
Q68FD5	Clathrin heavy chain 1	CLH1 MOUSE	95	0.99
Q6ZWN5	40S ribosomal protein S9	RS9 MOUSE	14	0.99
P60843	Eukaryotic initiation factor 4A-I	IF4A1_MOUSE	23	0.98
Q60930	Voltage-dependent anion- selective channel protein 2	VDAC2_MOUSE	16	0.98
Q9WV32	Actin-related protein 2/3 complex subunit 1B	ARC1B_MOUSE	19	0.98

P16110	Galectin-3	LEG3_MOUSE	10	0.97
P35441	Thrombospondin-1	TSP1_MOUSE	70	0.97
P05202	Aspartate aminotransferase, mitochondrial	AATM MOUSE	29	0.95
P48962	ADP/ATP translocase 1	ADT1 MOUSE	22	0.95
P58252	Elongation factor 2	EF2 MOUSE	52	0.95
P62281	40S ribosomal protein S11	RS11 MOUSE	10	0.94
P62874	Guanine nucleotide-binding protein G(I)/G(S)/G(T) subunit beta-1	GBB1_MOUSE	13	0.94
Q9WTY4	Aquaporin-5	AQP5_MOUSE	8	0.94
Q9Z0K8	Pantetheinase	VNN1_MOUSE	23	0.94
O70435	Proteasome subunit alpha type-3	PSA3_MOUSE	14	0.93
P53994	Ras-related protein Rab-2A	RAB2A_MOUSE	15	0.93
0311/17	Keratin, type II cytoskeletal	K220 MOUSE	30	0.93
P35486	Pyruvate dehydrogenase E1 component subunit alpha, somatic form, mitochondrial	ODPA MOUSE	25	0.92
	Glutathione reductase,			
P47791	mitochondrial	GSHR_MOUSE	25	0.91
Q9DAU7	domain protein 2	WFDC2_MOUSE	6	0.91
P24452	protein	CAPG_MOUSE	14	0.9
P62259	14-3-3 protein epsilon	1433E_MOUSE	18	0.9
Q9CZM2	60S ribosomal protein L15	RL15_MOUSE	14	0.9
DIADIA	Tissue-type plasminogen		04	0.00
P11214		TPA_MOUSE	31	0.89
P62242	405 ribosomal protein 58	RS8_MOUSE	10	0.89
Q8BG05	ribonucleoprotein A3	ROA3_MOUSE	20	0.89
Q9WUU7	Cathepsin Z	CATZ_MOUSE	14	0.89
P04104	Keratin, type II cytoskeletal 1	K2C1_MOUSE	35	0.88
Q61581	Insulin-like growth factor- binding protein 7	IBP7 MOUSE	18	0.88
P08113	Endoplasmin	ENPL MOUSE	47	0.87
Q3THE2	Myosin regulatory light chain 12B	ML12B_MOUSE	11	0.86
Q6ZWV3	60S ribosomal protein L10	RL10_MOUSE	13	0.86
P27659	60S ribosomal protein L3	RL3 MOUSE	22	0.85
P62331	ADP-ribosylation factor 6	ARF6_MOUSE	9	0.85
P63323	40S ribosomal protein S12	RS12_MOUSE	7	0.85
P67984	60S ribosomal protein L22	RL22_MOUSE	6	0.85
Q80X90	Filamin-B	FLNB MOUSE	146	0.85
Q99LX0	Protein DJ-1	PARK7 MOUSE	13	0.85
P06745	Glucose-6-phosphate isomerase	G6PI_MOUSE	32	0.84

P16460	Argininosuccinate synthase	ASSY_MOUSE	25	0.84
	Carbonyl reductase			
P48758	[NADPH] 1	CBR1_MOUSE	19	0.84
Q6ZWU9	40S ribosomal protein S27	RS27_MOUSE	5	0.83
	diphosphooligosaccharide			
	protein glycosyltransferase			
Q91YQ5	subunit 1	RPN1_MOUSE	40	0.83
Q9CXW4	60S ribosomal protein L11	RL11_MOUSE	9	0.83
O35640	Annexin A8	ANXA8_MOUSE	23	0.82
P56395	Cytochrome b5	CYB5_MOUSE	8	0.82
	Cytochrome b-c1 complex			
Q9CZ13	subunit 1, mitochondrial	QCR1_MOUSE	22	0.82
O88593	protein 1	PGRP1 MOUSE	10	0.81
Q60854	Serpin B6	SPB6 MOUSE	23	0.81
P14069	Protein S100-A6	S10A6 MOUSE	5	0.79
P61358	60S ribosomal protein I 27	RI 27 MOUSE	6	0.79
D62925	Pas related protein Pap 1A	PADIA MOUSE	10	0.79
F02033	Guanine nucleotide-binding	KAFTA_MOUSE	10	0.79
	protein G(I)/G(S)/G(T)			
P62880	subunit beta-2	GBB2_MOUSE	13	0.79
Q9QXC1	Fetuin-B	FETUB_MOUSE	17	0.79
D11092	T-complex protein 1		25	0.79
P11903		TCFA_MOUSE	30	0.70
P14206	Serine/threonine-protein	RSSA_MOUSE	15	0.78
	phosphatase PP1-alpha			
P62137	catalytic subunit	PP1A_MOUSE	19	0.78
P62911	60S ribosomal protein L32	RL32_MOUSE	8	0.77
	Na(+)/H(+) exchange			
P70441	RE1	NHRE1 MOUSE	27	0.77
P61514	60S ribosomal protein L 37a	RI 37A MOUSE	6	0.75
101014	Heterogeneous nuclear		0	0.70
P61979	ribonucleoprotein K	HNRPK_MOUSE	27	0.75
Deadea	Dynein light chain 1,			0.75
P63168		DYL1_MOUSE	5	0.75
P68254	14-3-3 protein theta	14331_MOUSE	18	0.75
P0CG49	Polyubiquitin-B	UBB_MOUSE	28	0.74
P40124	associated protein 1	CAP1 MOUSE	30	0.74
	26S protease regulatory			
P62196	subunit 8	PRS8_MOUSE	28	0.74
Q9CQV8	14-3-3 protein beta/alpha	1433B_MOUSE	17	0.74
0070¥1	Heterogeneous nuclear		22	0.74
Q9ZZA I	Proteasome subunit beta		22	0.74
O55234	type-5	PSB5_MOUSE	16	0.73
D / 0 0 0 0	Eukaryotic initiation factor			
P10630	4A-II	IF4A2_MOUSE	23	0.73
P47962	60S ribosomal protein L5	RL5_MOUSE	17	0.73
P08752	protein G(i) subunit alpha-2	GNAI2 MOUSE	20	0.72
			1	

	ATP synthase F(0)			
090007	mitochondrial	AT5F1 MOUSE	18	0.72
Q9DBH5	Vesicular integral- membrane protein VIP36	LMAN2 MOUSE	21	0.72
P14602	Heat shock protein beta-1	HSPB1 MOUSE	12	0.71
	Proteasome subunit beta			
P99026	type-4	PSB4_MOUSE	11	0.71
P80316	subunit epsilon	TCPE_MOUSE	36	0.7
P12970	60S ribosomal protein L7a	RL7A_MOUSE	16	0.69
P51150	Ras-related protein Rab-7a	RAB7A_MOUSE	17	0.69
P63330	Serine/threonine-protein phosphatase 2A catalytic subunit alpha isoform	PP2AA_MOUSE	18	0.69
P97494	Glutamatecysteine ligase catalytic subunit	GSH1_MOUSE	34	0.69
Q8BTM8	Filamin-A	FLNA_MOUSE	143	0.69
Q99KI0	Aconitate hydratase, mitochondrial	ACON_MOUSE	41	0.69
O54734	Dolichyl- diphosphooligosaccharide protein glycosyltransferase 48 kDa subunit	OST48_MOUSE	22	0.68
O70570	Polymeric immunoglobulin receptor	PIGR_MOUSE	38	0.68
P47955	60S acidic ribosomal protein P1	RLA1_MOUSE	4	0.68
Q8BP67	60S ribosomal protein L24	RL24_MOUSE	10	0.68
Q91V41	Ras-related protein Rab-14	RAB14_MOUSE	15	0.68
Q9D154	Leukocyte elastase inhibitor A	ILEUA_MOUSE	23	0.68
	Alcohol dehydrogenase	AK1A1 MOUSE	23	0.68
De0122		RUVB1 MOUSE	23	0.66
00122			22	0.66
Q93092	Heterogeneous nuclear	TALDO_WOUSL	23	0.00
Q99020	ribonucleoprotein A/B	ROAA_MOUSE	14	0.66
Q9D2Q8	Protein S100-A14	S10AE_MOUSE	8	0.66
P09405	Nucleolin	NUCL_MOUSE	48	0.65
P25444	40S ribosomal protein S2	RS2_MOUSE	22	0.65
Q61503	5~-nucleotidase	5NTD_MOUSE	32	0.65
Q6IME9	Keratin, type II cytoskeletal	K2C72_MOUSE	34	0.65
	Transmembrane emp24			
Q9D1D4	10	TMEDA MOUSE	10	0.65
P30115	Glutathione S-transferase A3	GSTA3 MOUSE	14	0.64
P54071	Isocitrate dehydrogenase [NADP], mitochondrial	IDHP_MOUSE	31	0.64
P80317	T-complex protein 1 subunit zeta	TCPZ_MOUSE	29	0.64
Q64727	Vinculin	VINC_MOUSE	84	0.64
Q9WTM5	RuvB-like 2	RUVB2_MOUSE	28	0.64

Q9CZU6 Citrate synt	hase, al	CISY MOUSE	21	0.63
Cytochrome P00405 Subunit 2	e c oxidase	COX2 MOUSE	6	0.62
Q61937 Nucleophos	min	NPM MOUSE	16	0.62
Q62186 protein sub	associated	SSRD_MOUSE	8	0.62
Q69ZN7 Myoferlin		MYOF MOUSE	131	0.62
O55142 60S ribosor	nal protein L35a	RL35A_MOUSE	9	0.61
P21107 Tropomyosi	n alpha-3 chain	TPM3_MOUSE	24	0.61
Macrophage	e migration		_	
P34884 inhibitory fa	ctor	MIF_MOUSE	5	0.61
Q8VEM8 mitochondri	al	MPCP_MOUSE	21	0.61
Q9Z2U1 type-5	e subunit alpha	PSA5_MOUSE	11	0.6
Q8BMS1 Trifunctiona subunit alph mitochondri	l enzyme na, al	ECHA_MOUSE	40	0.59
3-hydroxyad 008756 dehydroger	cyl-CoA ase type-2	HCD2_MOUSE	16	0.58
P35550 rRNA 2~-O- methyltrans	ferase fibrillarin	FBRL_MOUSE	24	0.58
P49312 Heterogene ribonucleop	ous nuclear rotein A1	ROA1_MOUSE	19	0.58
Heterogene	ous nuclear		05	0.57
		HINKHI_MOUSE	25	0.57
D60967 405 ribosor	ein sigma	14335_MOUSE	19	0.57
Small nucle	nai protein 520 ar	RS20_MOUSE	6	0.57
P62315 ribonucleop	rotein Sm D1	SMD1_MOUSE	5	0.57
P62855 40S ribosor	nal protein S26	RS26_MOUSE	6	0.57
Q9QUM9 type-6	e subunit alpha	PSA6_MOUSE	13	0.57
Protein disu	Ilfide-isomerase	PDIA4 MOUSE	13	0.56
POCOS6 Histore H2	۸ 7	H2AZ MOUSE	43 6	0.56
P61982 14-3-3 prote		1433G MOUSE	18	0.56
P68510 14-3-3 prote		1433E_MOUSE	18	0.56
Asparagine	tRNA ligase,		10	0.00
Q8BP47 cytoplasmic	-	SYNC_MOUSE	30	0.56
Q9.IM76 Complex su	d protein 2/3 bunit 3	ARPC3 MOUSE	12	0.56
P45376 Aldose redu	ictase	ALDR MOUSE	20	0.55
Rab GDP d	issociation			
P50396 inhibitor alp	ha	GDIA_MOUSE	26	0.55
P62754 40S ribosor	nal protein S6	RS6_MOUSE	13	0.55
P62852 40S ribosor	nal protein S25	RS25_MOUSE	6	0.55
O78IK2	d during scle growth	USMG5 MOUSE	3	0.55
P12265 Beta-alucur	onidase	BGLR MOUSE	33	0.54
P35980 60S ribosor	nal protein I 18	RL18 MOUSE	9	0.54
P50580 Proliferation	n-associated	PA2G4 MOUSE	24	0.54

	protein 2G4			
Q9R118	Serine protease HTRA1	HTRA1_MOUSE	21	0.54
O54990	Prominin-1	PROM1_MOUSE	41	0.53
P09803	Cadherin-1	CADH1_MOUSE	33	0.53
P63001	Ras-related C3 botulinum toxin substrate 1	RAC1_MOUSE	9	0.53
Q921F2	TAR DNA-binding protein 43	TADBP_MOUSE	17	0.53
P14685	26S proteasome non- ATPase regulatory subunit	PSMD3 MOUSE	39	0.52
P62082	40S ribosomal protein S7	RS7 MOUSE	11	0.52
Q8BF79	Frlin-2	FRI N2 MOUSE	24	0.52
Q8BHN3	Neutral alpha-glucosidase AB	GANAB_MOUSE	51	0.52
Q9D8W5	26S proteasome non- ATPase regulatory subunit 12	PSD12_MOUSE	34	0.52
O55143	Sarcoplasmic/endoplasmic reticulum calcium ATPase 2	AT2A2_MOUSE	54	0.51
Q60668	Heterogeneous nuclear ribonucleoprotein D0	HNRPD_MOUSE	15	0.51
Q9R0Q3	Transmembrane emp24 domain-containing protein 2	TMED2_MOUSE	11	0.51
008807	Peroxiredoxin-4	PRDX4_MOUSE	16	0.5
P29341	protein 1	PABP1_MOUSE	38	0.5
P30412	isomerase C	PPIC_MOUSE	11	0.5
Q99K51	Plastin-3	PLST_MOUSE	40	0.5
Q9CYN9	Renin receptor	RENR_MOUSE	17	0.5
P12032	Metalloproteinase inhibitor 1	TIMP1_MOUSE	12	0.49
P14148	60S ribosomal protein L7	RL7_MOUSE	18	0.49
P57776	Elongation factor 1-delta	EF1D_MOUSE	20	0.49
P97430	Antileukoproteinase	SLPI_MOUSE	11	0.49
Q99PT1	Rho GDP-dissociation inhibitor 1	GDIR1_MOUSE	14	0.49
Q9DB20	ATP synthase subunit O, mitochondrial	ATPO_MOUSE	14	0.49
Q9WU78	interacting protein	PDC6I_MOUSE	52	0.49
P10493	Nidogen-1	NID1_MOUSE	46	0.48
P17225	Polypyrimidine tract-binding protein 1	PTBP1_MOUSE	22	0.48
P28474	class-3	ADHX_MOUSE	20	0.48
P62814	V-type proton ATPase subunit B, brain isoform	VATB2_MOUSE	29	0.48
P62849	40S ribosomal protein S24	RS24_MOUSE	7	0.48
P14733	Lamin-B1	LMNB1_MOUSE	39	0.47
P41105	60S ribosomal protein L28	RL28_MOUSE	11	0.47

P47963	60S ribosomal protein L13	RL13_MOUSE	14	0.47
	T-complex protein 1			
P80315	subunit delta	TCPD_MOUSE	37	0.47
Q8BFR5	mitochondrial	EFTU MOUSE	31	0.47
	Endoplasmic reticulum-			0
	Golgi intermediate			
Q9DC16	compartment protein 1	ERGI1_MOUSE	13	0.47
O70251	Elongation factor 1-beta	EF1B_MOUSE	14	0.46
P14824	Annexin A6	ANXA6_MOUSE	49	0.46
P14131	40S ribosomal protein S16	RS16_MOUSE	12	0.45
P24270	Catalase	CATA_MOUSE	35	0.45
P24527	Leukotriene A-4 hvdrolase	LKHA4 MOUSE	39	0.45
	T-complex protein 1			
P42932	subunit theta	TCPQ_MOUSE	39	0.45
P53026	60S ribosomal protein L10a	RL10A_MOUSE	15	0.45
0791\/5	Mitochondrial carrier	MTCH2 MOUSE	14	0.45
	Cutochromo hE typo P		0	0.45
QUQAZ	Dolichyl-		0	0.45
	diphosphooligosaccharide			
	protein glycosyltransferase			
Q9DBG6	subunit 2	RPN2_MOUSE	26	0.45
	GIP:AMP			
Q9W/TP7	mitochondrial	KAD3 MOUSE	19	0.45
Gowin	Glutathione S-transferase		10	0.10
P13745	A1	GSTA1_MOUSE	14	0.44
	60S acidic ribosomal			
P14869	protein P0	RLA0_MOUSE	18	0.44
P49722	type-2	PSA2 MOUSE	13	0 44
1 101 22	Ubiquitin-conjugating		10	
P61079	enzyme E2 D3	UB2D3_MOUSE	5	0.44
	Platelet-activating factor			
061206	acetyinydrolase IB subunit	PA1B2 MOUSE	10	0.44
Q01200	Actin-related protein 2/3	FAID2_WOUGL	10	0.44
Q9CVB6	complex subunit 2	ARPC2_MOUSE	22	0.44
	Sodium/potassium-			
D14004	transporting ATPase		10	0.42
P14094		ATTBT_WOUSE	10	0.43
P61089	enzyme E2 N	UBE2N_MOUSE	11	0.43
P61255	60S ribosomal protein L26	RL26 MOUSE	9	0.43
	26S protease regulatory			
P62334	subunit 10B	PRS10_MOUSE	27	0.43
Q8K183	Pyridoxal kinase	PDXK_MOUSE	16	0.43
	3-ketoacyl-CoA thiolase A,			
Q921H8	peroxisomal	THIKA_MOUSE	22	0.43
	binding protein 2			
Q9D0S9	mitochondrial	HINT2_MOUSE	11	0.43
	NADP-dependent malic			
P06801	enzyme	MAOX_MOUSE	32	0.42
P16675	Lysosomal protective	PPGB_MOUSE	22	0.42

	protein			
Q62267	Cornifin-B	SPR1B_MOUSE	15	0.42
0.01/07.0	Succinate dehydrogenase [ubiquinone] flavoprotein			
Q8K2B3	subunit, mitochondrial	SDHA_MOUSE	36	0.42
Q9JJI8	60S ribosomal protein L38	RL38_MOUSE	3	0.42
P35979	60S ribosomal protein L12	RL12_MOUSE	11	0.41
P62270	40S ribosomal protein S18	RS18_MOUSE	12	0.41
Q8R081	Heterogeneous nuclear ribonucleoprotein L	HNRPL_MOUSE	24	0.41
Q9JJ00	Phospholipid scramblase 1	PLS1_MOUSE	13	0.41
D 45050	Medium-chain specific acyl- CoA dehydrogenase,			
P45952	mitochondrial	ACADM_MOUSE	22	0.4
P60335	Poly(rC)-binding protein 1	PCBP1_MOUSE	20	0.4
P61939	Thyroxine-binding globulin	THBG_MOUSE	22	0.4
000704	Vitamin K-dependent		07	
Q08761	protein S	PROS_MOUSE	37	0.4
Q61753	dehydrogenase	SERA_MOUSE	22	0.4
Q9D8E6	60S ribosomal protein L4	RL4_MOUSE	27	0.4
0.00411/22	Vesicle-associated membrane protein-			
Q9WV55	Associated protein A	VAPA_MOUSE	15	0.4
Q9Z2U0	type-7	PSA7_MOUSE	16	0.4
O09167	60S ribosomal protein L21	RL21_MOUSE	9	0.39
P63325	40S ribosomal protein S10	RS10_MOUSE	13	0.39
Q8VEK3	Heterogeneous nuclear ribonucleoprotein U	HNRPU_MOUSE	43	0.39
Q922R8	Protein disulfide-isomerase A6	PDIA6_MOUSE	25	0.39
Q9DB77	Cytochrome b-c1 complex subunit 2, mitochondrial	QCR2_MOUSE	22	0.39
D12507	Intercellular adhesion		26	0.20
P13097			20	0.30
P17710	Hexokinase-1	HXK1_MOUSE	57	0.38
P70333	ribonucleoprotein H2	HNRH2_MOUSE	25	0.38
Q02053	activating enzyme 1	UBA1_MOUSE	55	0.38
Q6URW6	Myosin-14	MYH14_MOUSE	126	0.38
Q9DBJ1	Phosphoglycerate mutase	PGAM1_MOUSE	15	0.38
P26443	Glutamate dehydrogenase 1, mitochondrial	DHE3_MOUSE	35	0.37
P48771	Cytochrome c oxidase subunit 7A2, mitochondrial	CX7A2_MOUSE	5	0.37
P59999	Actin-related protein 2/3 complex subunit 4	ARPC4_MOUSE	13	0.37
P70195	Proteasome subunit beta type-7	PSB7_MOUSE	12	0.37
Q62425	Cytochrome c oxidase subunit NDUFA4	NDUA4_MOUSE	6	0.37
Q9D6R2	Isocitrate dehydrogenase	IDH3A_MOUSE	21	0.37

	[NAD] subunit alpha, mitochondrial			
Q9R1P0	Proteasome subunit alpha type-4	PSA4_MOUSE	14	0.37
O55023	Inositol monophosphatase	IMPA1_MOUSE	17	0.36
P42208	Septin-2	SEPT2_MOUSE	21	0.36
Q8CAQ8	MICOS complex subunit Mic60	MIC60_MOUSE	55	0.36
Q99JY0	Trifunctional enzyme subunit beta, mitochondrial	ECHB_MOUSE	28	0.36
P04186	Complement factor B	CFAB_MOUSE	42	0.35
P13020	Gelsolin	GELS_MOUSE	34	0.35
P25085	Interleukin-1 receptor antagonist protein	IL1RA_MOUSE	7	0.35
P61211	ADP-ribosylation factor-like protein 1	ARL1_MOUSE	10	0.35
P70124	Serpin B5	SPB5 MOUSE	22	0.35
Q9QZ88	Vacuolar protein sorting- associated protein 29	VPS29_MOUSE	11	0.35
008749	Dihydrolipoyl dehydrogenase, mitochondrial		26	0.34
000749	Fatty aldehyde		20	0.34
P47740	dehydrogenase	AL3A2_MOUSE	21	0.34
P97384	Annexin A11	ANX11_MOUSE	22	0.34
P97449	Aminopeptidase N	AMPN_MOUSE	51	0.34
P97807	Fumarate hydratase, mitochondrial	FUMH_MOUSE	32	0.34
Q62465	Synaptic vesicle membrane protein VAT-1 homolog	VAT1 MOUSE	18	0.34
	Non-POU domain-			
Q99K48	containing octamer-binding protein	NONO_MOUSE	23	0.34
000000	Protein transport protein			0.04
<u>Q9CQS8</u>	Sec61 subunit beta	SC61B_MOUSE	6	0.34
Q9DC53	Copine-8	CPNE8_MOUSE	33	0.34
Q9JKR6	protein 1	HYOU1_MOUSE	58	0.34
	Membrane-associated			
055022	component 1	PGRC1 MOUSE	10	0.33
007813	Apoptosis regulator BAX	BAX MOUSE	10	0.33
	ADP-ribosylation factor-like	ARI 8B MOUSE	12	0.33
	Transmembrane protein 43	TMM43_MOUSE	21	0.33
	GTP-binding protein		21	0.00
P36536	SAR1a	SAR1A_MOUSE	11	0.32
P46978	Dolichyl- diphosphooligosaccharide protein glycosyltransferase subunit STT3A	STT3A_MOUSE	36	0.32
P80314	T-complex protein 1 subunit beta	TCPB_MOUSE	36	0.32
Q7TMK9	Heterogeneous nuclear ribonucleoprotein Q	HNRPQ_MOUSE	39	0.32

	Dihydrolipoyllysine-residue acetyltransferase			
	component of pyruvate			
Q8BMF4	mitochondrial	ODP2_MOUSE	33	0.32
Q8JZU2	Tricarboxylate transport protein, mitochondrial	TXTP_MOUSE	17	0.32
Q922B2	AspartatetRNA ligase, cytoplasmic	SYDC_MOUSE	38	0.32
Q9CQN1	Heat shock protein 75 kDa, mitochondrial	TRAP1_MOUSE	50	0.32
Q9DCN2	reductase 3	NB5R3_MOUSE	17	0.32
O70503	Estradiol 17-beta- dehydrogenase 12	DHB12_MOUSE	18	0.31
O88310	Intelectin-1a	ITL1A_MOUSE	17	0.31
	Aspartate			
P05201	aminotransferase,	AATC MOUSE	27	0.31
1 00201	Transcriptional activator		21	0.01
P42669	protein Pur-alpha	PURA_MOUSE	15	0.31
P62305	Small nuclear ribonucleoprotein E	RUXE_MOUSE	4	0.31
Q02257	Junction plakoglobin	PLAK_MOUSE	44	0.31
	Endoplasmic reticulum			
Q9D1Q6	resident protein 44	ERP44_MOUSE	24	0.31
Q9JKF1	protein IQGAP1	IQGA1_MOUSE	93	0.31
Q9QWR8	Alpha-N- acetylgalactosaminidase	NAGAB MOUSE	27	0.31
	Proteasome subunit beta			
Q9R1P1	type-3	PSB3_MOUSE	9	0.31
P19253	60S ribosomal protein L13a	RL13A_MOUSE	17	0.3
	Small nuclear			
P27048	ribonucleoprotein- associated protein B	RSMB_MOUSE	14	0.3
P35279	Ras-related protein Rab-6A	RAB6A_MOUSE	17	0.3
D00040	T-complex protein 1		00	
P80313	T-complex protein 1	TCPH_MOUSE	33	0.3
P80318	subunit gamma	TCPG_MOUSE	38	0.3
P97467	Peptidyl-glycine alpha- amidating monooxygenase	AMD_MOUSE	51	0.3
Q62167	ATP-dependent RNA helicase DDX3X	DDX3X_MOUSE	40	0.3
Q8K3J9	G-protein coupled receptor family C group 5 member C	GPC5C_MOUSE	11	0.3
Q9CQD1	Ras-related protein Rab-5A	RAB5A_MOUSE	14	0.3
Q9ESW8	Pyroglutamyl-peptidase 1	PGPI_MOUSE	10	0.3
Q9JIW9	Ras-related protein Ral-B	RALB_MOUSE	10	0.3
	Succinyl-CoA ligase			
Q9WUM5	alpha, mitochondrial	SUCA MOUSE	15	0.3
Q9Z0L8	Gamma-glutamvl hvdrolase	GGH MOUSE	17	0.3
009547	Vesicle-trafficking protein		12	0.20
006047	3E6220	JUZZD_IVIUUSE	13	0.29

O70554	Small proline-rich protein 2B	SPR2B_MOUSE	5	0.29
P00493	Hypoxanthine-guanine phosphoribosyltransferase	HPRT_MOUSE	13	0.29
P46638	Ras-related protein Rab- 11B	RB11B_MOUSE	16	0.29
Q07076	Annexin A7	ANXA7_MOUSE	23	0.29
Q61879	Myosin-10	MYH10_MOUSE	125	0.29
Q8K353	Cysteine-rich and transmembrane domain- containing protein 1	CYTM1_MOUSE	2	0.29
Q8VDW0	ATP-dependent RNA helicase DDX39A	DX39A_MOUSE	26	0.29
0000/110	Redox-regulatory protein		4.4	0.00
Q9CTH2	FAMZ13A	FZI3A_MOUSE	14	0.29
Q9D103	transmembrane protein 1	IFM1_MOUSE	2	0.29
	Sulfide:quinone			
000440	oxidoreductase,			0.00
Q9R112	mitochondrial	SQRD_MOUSE	32	0.29
O54782	mannosidase	MA2B2_MOUSE	44	0.28
P30416	isomerase FKBP4	FKBP4 MOUSE	30	0.28
P47964	60S ribosomal protein L36	RL36_MOUSE	7	0.28
	High mobility group protein	—		
P63158	B1	HMGB1_MOUSE	11	0.28
Q3UQ28	Peroxidasin homolog	PXDN_MOUSE	86	0.28
Q60692	Proteasome subunit beta type-6	PSB6_MOUSE	11	0.28
Q62093	Serine/arginine-rich splicing factor 2	SRSF2_MOUSE	11	0.28
Q6IRU5	Clathrin light chain B	CLCB_MOUSE	10	0.28
Q78IS1	Transmembrane emp24 domain-containing protein 3	TMED3_MOUSE	13	0.28
Q91WS0	CDGSH iron-sulfur domain- containing protein 1	CISD1_MOUSE	6	0.28
	Bifunctional purine		25	0.00
<u> 7</u> 9C 10 19	Ras-related protein Rab-	PUR9_MOUSE	30	0.20
Q9ERI2	27A	RB27A_MOUSE	13	0.28
	EGF-containing fibulin-like			
000000	extracellular matrix protein		22	0.00
<u>Ganna</u>	Z Proteasome subunit beta	FBLIN4_WOUSE	23	0.28
O09061	type-1	PSB1_MOUSE	14	0.27
P51807	Dynein light chain Tctex-	DYLT1 MOUSE	7	0.27
	Dolichyl-			
	diphosphooligosaccharide			
P61804	subunit DAD1	DAD1 MOUSE	5	0.27
P83882	60S ribosomal protein L36a	RL36A MOUSE	6	0.27
Q8R3G9	Tetraspanin-8	TSN8 MOUSE	8	0.27
091852			15	0.27
SCINOL			10	0.21

00\//114.2	PhenylalaninetRNA ligase	SVER MOUSE	11	0.27
Q900A2			41	0.27
Q9ZZVVU	D-dopachrome	DINPEP_IVIOUSE	24	0.27
O35215	decarboxylase	DOPD MOUSE	9	0.26
	Transcription elongation			
P62869	factor B polypeptide 2	ELOB_MOUSE	9	0.26
Q61656	Probable ATP-dependent RNA helicase DDX5	DDX5_MOUSE	35	0.26
Q64437	Alcohol dehydrogenase class 4 mu/sigma chain	ADH7 MOUSE	23	0.26
Q6PCW6	Cornifelin	CNFN MOUSE	2	0.26
	Aflatoxin B1 aldehyde			
Q8CG76	reductase member 2	ARK72_MOUSE	17	0.26
Q8C194	Glycogen phosphorylase, brain form	PYGB MOUSE	53	0.26
	ATP-dependent RNA	—		
Q91VR5	helicase DDX1	DDX1_MOUSE	46	0.26
Q91YD6	Villin-like protein	VILL_MOUSE	51	0.26
	EF-hand domain-containing			
Q9D8Y0	protein D2	EFHD2_MOUSE	16	0.26
P12023	Amyloid beta A4 protein	A4_MOUSE	39	0.25
	Thioredoxin-dependent			
P20108	mitochondrial	PRDX3 MOUSE	15	0.25
D46725			70	0.25
P40735			70	0.25
Q02248	Catenin beta-1	CINB1_MOUSE	42	0.25
Q571E4	sulfatase	GALNS_MOUSE	25	0.25
Q8BK67	Protein RCC2	RCC2 MOUSE	32	0.25
Q8K2I4	Beta-mannosidase	MANBA MOUSE	49	0.25
	26S proteasome non-			
	ATPase regulatory subunit			
Q8VDM4	2	PSMD2_MOUSE	56	0.25
0011/12	Cytosolic acyl coenzyme A		20	0.25
			20	0.25
Q9D0F3	Protein ERGIC-53	LMAN1_MOUSE	25	0.25
Q9D1R9	60S ribosomal protein L34	RL34_MOUSE	6	0.25
	Protein arginine N-	ANM1 MOUSE	20	0.25
	4F2 cell-surface antigen		20	0.25
P10852	heavy chain	4F2_MOUSE	30	0.24
	Small nuclear			
P62320	ribonucleoprotein Sm D3	SMD3_MOUSE	9	0.24
P70349	Histidine triad nucleotide- binding protein 1	HINT1 MOUSE	8	0.24
	Short-chain specific acyl-		-	
	CoA dehydrogenase,			
Q07417	mitochondrial	ACADS_MOUSE	21	0.24
Q61703	heavy chain H2	ITIH2_MOUSE	48	0.24
	Brain-specific angiogenesis			
0.051011	inhibitor 1-associated			
Q8BKX1	protein 2	BAIP2_MOUSE	42	0.24
Q8BT60	Copine-3	CPNE3_MOUSE	26	0.24

0000145	Thioredoxin domain-			0.24
Q9CQIVI5	Containing protein 17	TXD17_MOUSE	8	0.24
Q9DCV5	254	TM254_MOUSE	4	0.24
O09172	Glutamatecysteine ligase regulatory subunit	GSH0_MOUSE	12	0.23
P23953	Carboxylesterase 1C	EST1C_MOUSE	25	0.23
P28656	Nucleosome assembly	NP1L1 MOUSE	13	0.23
P62000	60S ribosomal protein 31		7	0.23
F 02300	Heat shock 70 kDa protein		/	0.23
Q61316	4	HSP74_MOUSE	57	0.23
Q64310	Surfeit locus protein 4	SURF4 MOUSE	9	0.23
	UDP-			
	glucuronosyltransferase 1-			
Q64435	6	UD16_MOUSE	25	0.23
Q9CX86	Heterogeneous nuclear ribonucleoprotein A0	ROA0_MOUSE	15	0.23
Q9D0T1	NHP2-like protein 1	NH2L1 MOUSE	9	0.23
	DnaJ homolog subfamily A			
Q9QYJ0	member 2	DNJA2_MOUSE	23	0.23
P16045	Galectin-1	LEG1_MOUSE	9	0.22
	Cation-dependent			
D0 4000	mannose-6-phosphate		10	0.00
P24668	receptor	MPRD_MOUSE	18	0.22
P50247	Adenosylhomocysteinase	SAHH_MOUSE	27	0.22
P63276	40S ribosomal protein S17	RS17_MOUSE	6	0.22
Q02496	Mucin-1	MUC1_MOUSE	11	0.22
Q7TQI3	Ubiquitin thioesterase OTUB1	OTUB1_MOUSE	15	0.22
Q8BH95	Enoyl-CoA hydratase, mitochondrial	ECHM_MOUSE	18	0.22
Q8VC85	U6 snRNA-associated Sm- like protein LSm1	LSM1 MOUSE	11	0.22
099.170	Syntenin-2	SDCB2 MOUSE	16	0.22
000020	AcvI-coenzyme A		10	0.22
Q9CQR4	thioesterase 13	ACO13_MOUSE	10	0.22
	Interferon-induced			
Q9CQW9	transmembrane protein 3	IFM3_MOUSE	4	0.22
		PUR6 MOUSE	27	0.22
	Major yoult protoin		51	0.22
Q9EQK5	Acidic leucipe-rich puclear	MVP_MOUSE	51	0.22
	phosphoprotein 32 family			
Q9EST5	member B	AN32B_MOUSE	12	0.22
	Mitochondrial dicarboxylate			
Q9QZD8	carrier	DIC_MOUSE	17	0.22
	Ectonucleoside			
09\//179	diphosphobydrolase 5	ENTPS MOUSE	24	0.22
008002	Syntenin_1		15	0.21
000992			10	0.21
055131	26S protesse regulatory	SEPI7_MOUSE	20	0.21
O88685	subunit 6A	PRS6A MOUSE	26	0.21
P10923	Osteopontin	OSTP MOUSE	12	0.21
1 10020			14	5.21

D12787	Cytochrome c oxidase	COX5A MOUSE	10	0.21
F 12707	Spectrin alpha chain, non-		10	0.21
P16546	erythrocytic 1	SPTN1_MOUSE	189	0.21
P20352	Tissue factor	TF_MOUSE	17	0.21
P21460	Cystatin-C	CYTC MOUSE	10	0.21
P35564	Calnexin	CALX MOUSE	27	0.21
	Enoyl-CoA delta isomerase			
P42125	1, mitochondrial	ECI1_MOUSE	18	0.21
P47753	F-actin-capping protein subunit alpha-1	CAZA1_MOUSE	17	0.21
P47754	F-actin-capping protein subunit alpha-2	CAZA2 MOUSE	15	0.21
	V-type proton ATPase			
P50516	catalytic subunit A	VATA_MOUSE	37	0.21
DE0106	Thiosulfate		10	0.21
P52190	V-type proton ATPase 16		10	0.21
P63082	kDa proteolipid subunit	VATL_MOUSE	3	0.21
Q00915	Retinol-binding protein 1	RET1 MOUSE	12	0.21
	2-oxoglutarate			
000507	dehydrogenase,		50	0.04
Q60597	mitochondriai	ODO1_MOUSE	52	0.21
Q6ZQI3	Malectin	MLEC_MOUSE	18	0.21
Q8C1B7	Septin-11	SEP11_MOUSE	25	0.21
Q8R164	Valacyclovir hydrolase	BPHL_MOUSE	23	0.21
Q99K30	receptor kinase substrate 8-like protein 2	ES8L2_MOUSE	40	0.21
Q9D2G2	Dihydrolipoyllysine-residue succinyltransferase component of 2- oxoglutarate dehydrogenase complex, mitochondrial	ODO2 MOUSE	23	0.21
Q9EQU5	Protein SET	SET MOUSE	12	0.21
	Acyl-coenzyme A			
	thioesterase 9,		20	0.01
Q9R0X4	Succipyl-CoA ligase IADP-	ACOT9_MOUSE	30	0.21
097219	forming] subunit beta, mitochondrial	SUCB1 MOUSE	31	0.21
P62204	Calmodulin	CALM MOUSE	9	0.2
P97326	Cadherin-6	CADH6 MOUSE	40	0.2
002810		NUCB1 MOUSE	20	0.2
Q02019	Hvdroxvacvl-coenzyme A		23	0.2
	dehydrogenase,			
Q61425	mitochondrial	HCDH_MOUSE	19	0.2
Q7TNG5	Echinoderm microtubule- associated protein-like 2	EMAL2_MOUSE	31	0.2
Q80UW2	F-box only protein 2	FBX2_MOUSE	14	0.2
Q80VQ0	Aldehyde dehydrogenase family 3 member B1	AL3B1_MOUSE	24	0.2
Q80Y14	Glutaredoxin-related protein 5, mitochondrial	GLRX5_MOUSE	8	0.2

	Calcium-binding mitochondrial carrier			
Q8BMD8	protein SCaMC-1	SCMC1_MOUSE	29	0.2
	NADH dehydrogenase			
Q9ERS2	subcomplex subunit 13	NDUAD MOUSE	10	0.2
Q9WVJ3	Carboxypeptidase Q	CBPQ MOUSE	21	0.2
	Eukaryotic translation			
007014	initiation factor 2 subunit 3,		24	0.0
			24	0.2
Qazilda	I -lactate dehydrogenase B	STVC_NOUSE	00	0.2
P16125	chain	LDHB_MOUSE	21	0.19
P27046	Alpha-mannosidase 2	MA2A1_MOUSE	63	0.19
P70372	ELAV-like protein 1	ELAV1_MOUSE	17	0.19
0041/07	Microsomal glutathione S-			0.40
Q91VS7	transferase 1 Epithelial cell adhesion	MGST1_MOUSE	6	0.19
Q99JW5	molecule	EPCAM_MOUSE	14	0.19
	ATP-dependent (S)-			
000742	NAD(P)H-hydrate		10	0.10
Q9C242	Drotoin SEC12 homolog	SEC12 MOUSE	10	0.19
Q9D11VIO	ATP synthase subunit	3EC13_1003E	13	0.19
Q9D3D9	delta, mitochondrial	ATPD_MOUSE	5	0.19
Q9DAW9	Calponin-3	CNN3_MOUSE	18	0.19
000750	Eukaryotic translation			0.40
Q9QZD9	Initiation factor 3 subunit I	EIF3I_MOUSE	22	0.19
O35685	nudC	NUDC_MOUSE	23	0.18
O70492	Sorting nexin-3	SNX3_MOUSE	10	0.18
Q61990	Poly(rC)-binding protein 2	PCBP2_MOUSE	18	0.18
	Transmembrane channel-			
Q7TQ65	like protein 4	TMC4_MOUSE	33	0.18
Q80YX1	Tenascin	TENA_MOUSE	105	0.18
Q8JZM8	Mucin-4	MUC4_MOUSE	115	0.18
Q8K023	1 member C18	AKC1H MOUSE	22	0.18
	DnaJ homolog subfamily C			
Q91YW3	member 3	DNJC3_MOUSE	32	0.18
	Tubulin polymerization-			
Q9CRB6	member 3	TPPP3 MOUSE	14	0.18
	Heterogeneous nuclear			
Q9D0E1	ribonucleoprotein M	HNRPM_MOUSE	52	0.18
	ATP synthase subunit d, mitochondrial	ATP5H MOUSE	11	0.18
	Peroxisomal acyl-			0.10
Q9QXD1	coenzyme A oxidase 2	ACOX2_MOUSE	41	0.18
	Mitochondrial import			
Q9QYA2	homolog	TOM40 MOUSE	12	0.18
	Transmembrane protein			
B2RWJ3	240	TM240_MOUSE	6	0.17
P20060	Beta-hexosaminidase	HEXB MOUSE	30	0.17
1 20000	Sabarin botu			0.17

P26231	Catenin alpha-1	CTNA1_MOUSE	56	0.17
	Connective tissue growth			
P29268	factor	CTGF_MOUSE	24	0.17
P51660	enzyme type 2	DHB4_MOUSE	40	0.17
P63028	Translationally-controlled tumor protein	TCTP_MOUSE	7	0.17
P84104	Serine/arginine-rich splicing factor 3	SRSF3_MOUSE	10	0.17
Q64176	Carboxylesterase 1E	EST1E MOUSE	26	0.17
Q8BHL4	Retinoic acid-induced protein 3	RAI3 MOUSE	13	0.17
Q8BI08	Protein MAL2	MAL2 MOUSE	2	0.17
	Phosphoserine			
Q99K85	aminotransferase	SERC_MOUSE	27	0.17
Q99KK7	Dipeptidyl peptidase 3	DPP3_MOUSE	46	0.17
Q9CX00	IST1 homolog	IST1_MOUSE	12	0.17
	EH domain-containing			
Q9EQP2	protein 4	EHD4_MOUSE	32	0.17
Q9QUR6	Prolyl endopeptidase	PPCE_MOUSE	47	0.17
Q9QZQ8	Core histone macro-H2A.1	H2AY_MOUSE	19	0.17
P15864	Histone H1.2	H12_MOUSE	12	0.16
P23198	Chromobox protein homolog 3	CBX3_MOUSE	10	0.16
	Isocitrate dehydrogenase			
P70404	[NAD] subunit gamma 1, mitochondrial	IDHG1 MOUSE	17	0.16
170404	Cysteine and glycine-rich		17	0.10
P97315	protein 1	CSRP1_MOUSE	14	0.16
Q63844	Mitogen-activated protein kinase 3	MK03 MOUSE	22	0.16
	Serine/threonine-protein			
	phosphatase 2A 65 kDa			
076M73	isoform	2444 MOUSE	30	0.16
	Cvtoskeleton-associated			0.10
Q8BMK4	protein 4	CKAP4_MOUSE	40	0.16
	3-ketoacyl-CoA thiolase,			
Q8BWT1	mitochondrial	THIM_MOUSE	24	0.16
Q8VCC2	Liver carboxylesterase 1	EST1_MOUSE	23	0.16
Q9R0P6	Signal peptidase complex catalytic subunit SEC11A	SC11A_MOUSE	12	0.16
Q9R0Q6	Actin-related protein 2/3 complex subunit 1A	ARC1A_MOUSE	23	0.16
	Putative pre-mRNA-splicing factor ATP-dependent RNA	B. N/1		
035286	helicase DHX15	DHX15_MOUSE	43	0.15
P11438	membrane glycoprotein 1	LAMP1_MOUSE	18	0.15
P12815	Programmed cell death protein 6	PDCD6_MOUSE	10	0.15
P15535	Beta-1,4- galactosyltransferase 1	B4GT1 MOUSE	19	0.15
P61161	Actin-related protein 2		21	0.15
	Tumor susceptibility gene		<u> </u>	0.10
Q61187	101 protein	TS101_MOUSE	18	0.15

Q62261	Spectrin beta chain, non- erythrocytic 1	SPTB2 MOUSE	161	0.15
Q62470	Integrin alpha-3	ITA3 MOUSE	40	0.15
064324	Syntaxin-binding protein 2	STXB2 MOUSE	40	0.15
Q04024	High affinity copper uptake			0.10
Q8K211	protein 1	COPT1_MOUSE	3	0.15
Q8R2S8	CD177 antigen	CD177_MOUSE	37	0.15
Q8VCF1	Soluble calcium-activated nucleotidase 1	CANT1_MOUSE	27	0.15
Q9CZ30	Obg-like ATPase 1	OLA1_MOUSE	26	0.15
Q9DBP5	UMP-CMP kinase	KCY_MOUSE	11	0.15
Q9DCS2	UPF0585 protein C16orf13 homolog	CP013_MOUSE		0.15
Q9JHI5	Isovaleryl-CoA dehydrogenase, mitochondrial	IVD_MOUSE	24	0.15
Q9Z0F4	binding protein 1	CIB1_MOUSE	15	0.15
Q9Z261	Claudin-7	CLD7_MOUSE	4	0.15
O88668	Protein CREG1	CREG1_MOUSE	11	0.14
Daaaaa	Urokinase-type		05	
P06869	plasminogen activator	UROK_MOUSE	25	0.14
P10833	Ras-related protein R-Ras	RRAS_MOUSE	16	0.14
P28301	Protein-lysine 6-oxidase	LYOX_MOUSE	20	0.14
P30275	mitochondrial	KCRU_MOUSE	26	0.14
P35282	Ras-related protein Rab-21	RAB21_MOUSE	15	0.14
P35293	Ras-related protein Rab-18	RAB18_MOUSE	14	0.14
P45700	Mannosyl-oligosaccharide 1,2-alpha-mannosidase IA	MA1A1_MOUSE	37	0.14
P46467	Vacuolar protein sorting- associated protein 4B	VPS4B_MOUSE	27	0.14
P46471	26S protease regulatory subunit 7	PRS7 MOUSE	31	0 14
P54775	26S protease regulatory subunit 6B	PRS6B_MOUSE	22	0.14
P84099	60S ribosomal protein L19	RL19 MOUSE	9	0.14
Q501J6	Probable ATP-dependent RNA helicase DDX17	DDX17_MOUSE	35	0.14
060963	Platelet-activating factor	PAFA MOUSE	23	0 14
062165	Dystrodycan	DAG1_MOUSE	42	0.14
078107	Zinc transporter ZIP4	S3944 MOUSE	18	0.14
	26S proteasome non-	0000L	10	0.14
	ATPase regulatory subunit			
Q8BG32	11 Coll surface alveenratein	PSD11_MOUSE	32	0.14
Q8R2Y2	MUC18	MUC18_MOUSE	34	0.14
Q8VDP6	CDP-diacylglycerolinositol 3-phosphatidyltransferase	CDIPT_MOUSE	10	0.14
Q8VIJ6	Splicing factor, proline- and glutamine-rich	SFPQ_MOUSE	31	0.14
Q91VC3	Eukaryotic initiation factor	IF4A3_MOUSE	25	0.14

Q91W90	Thioredoxin domain- containing protein 5	TXND5 MOUSE	23	0.14
Q99LP6	GrpE protein homolog 1, mitochondrial	GRPE1 MOUSE	17	0.14
Q9DCT8	Cysteine-rich protein 2	CRIP2 MOUSE	11	0.14
	5~(3~)-			
	deoxyribonucleotidase,		11	0.14
				0.14
	Unconventional myosin-ic	MYUIC_MOUSE	66	0.14
Q9WTL2	Ras-related protein Rab-25	RAB25_MOUSE	14	0.14
Q9Z2I8	forming] subunit beta, mitochondrial	SUCB2_MOUSE	30	0.14
P09671	Superoxide dismutase [Mn], mitochondrial	SODM MOUSE	13	0.13
P35762	CD81 antigen	CD81 MOUSE	7	0.13
P40240	CD9 antigen	CD9 MOUSE	6	0.13
P51855	Glutathione synthetase	GSHB MOUSE	29	0.13
	Hepatoma-derived growth			
P51859	factor	HDGF_MOUSE	16	0.13
P63073	Eukaryotic translation initiation factor 4E	IF4E_MOUSE	14	0.13
P70202	Latexin	LXN_MOUSE	6	0.13
	26S proteasome non-			
007/07	ATPase regulatory subunit		50	0.40
Q31X57	1 Platelet-activating factor	PSMD1_MOUSE	50	0.13
	acetylhydrolase IB subunit			
Q61205	gamma	PA1B3_MOUSE	17	0.13
	Glycerol-3-phosphate			
064521	denydrogenase, mitochondrial	GPDM MOUSE	48	0.13
	Clarin-3	CLRN3 MOUSE	6	0.13
	Bleomycin hydrolase		28	0.13
QONUTO	Epidermal growth factor		20	0.13
	receptor kinase substrate			
Q8R5F8	8-like protein 1	ES8L1_MOUSE	41	0.13
	Occludin/ELL domain-		11	0.13
0000119	Placenta-expressed		14	0.13
Q8VEN2	transcript 1 protein	PLET1_MOUSE	4	0.13
000/T4	Carbonyl reductase family			0.40
Q91V14	member 4	CBR4_MOUSE	14	0.13
Q91VU0	Protein FAM3C	FAM3C_MOUSE	16	0.13
Q99JR5	antigen-like	TINAL MOUSE	22	0.13
Q99LJ1	Tissue alpha-L-fucosidase	FUCO MOUSE	25	0.13
Q9D8T0	Protein FAM3A	FAM3A MOUSE	15	0.13
00010	Ubiquitin carboxyl-terminal			
Q9JKB1	hydrolase isozyme L3	UCHL3_MOUSE	13	0.13
000150	Lysosomal alpha-		47	0.12
009159	Fukarvotic translation		4/	0.12
O55135	initiation factor 6	IF6_MOUSE	10	0.12
P12382	ATP-dependent 6-	PFKAL_MOUSE	38	0.12

	phosphofructokinase, liver			
P29533	Vascular cell adhesion protein 1	VCAM1_MOUSE	39	0.12
	Carcinoembryonic antigen-			
P31809	molecule 1	CEAM1_MOUSE	18	0.12
P32921	TryptophantRNA ligase, cytoplasmic	SYWC_MOUSE	33	0.12
P62918	60S ribosomal protein L8	RL8 MOUSE	14	0.12
P97290	Plasma protease C1 inhibitor	IC1_MOUSE	25	0.12
Q5SYD0	Unconventional myosin-Id	MYO1D_MOUSE	74	0.12
Q61553	Fascin	FSCN1_MOUSE	29	0.12
Q6IRU2	Tropomyosin alpha-4 chain	TPM4_MOUSE	20	0.12
	Serine/arginine-rich splicing			
Q6PDM2	factor 1	SRSF1_MOUSE	15	0.12
Q91ZJ5	UTPglucose-1-phosphate uridylyltransferase	UGPA_MOUSE	30	0.12
	von Willebrand factor A			
OOOKC8	domain-containing protein	VMA5A MOUSE	15	0.12
033100	Pre-mRNA-processing		40	0.12
Q99KP6	factor 19	PRP19_MOUSE	19	0.12
0001/01	Nicotinamide			0.40
Q99KQ4	phosphoribosyltransferase	NAMPT_MOUSE	29	0.12
Q99LF4	homolog	RTCB MOUSE	29	0.12
		AMPL MOUSE	32	0.12
			50	0.12
Q902D3	Succinvl-CoA:3-ketoacid	310_MOUSE	50	0.12
Q9D0K2	coenzyme A transferase 1, mitochondrial	SCOT1_MOUSE	25	0.12
	NADH dehydrogenase			
	[ubiquinone] flavoprotein 2, mitochondrial	NDUV2 MOUSE	16	0.12
Q9D030			10	0.12
QaDioi	Brain-specific angiogenesis	UPR3L_INIOUSE	10	0.12
	inhibitor 1-associated			
Q9DBJ3	protein 2-like protein 1	BI2L1_MOUSE	38	0.12
0000000	Persulfide dioxygenase		47	0.40
			17	0.12
Q9WUM3	Coronin-1B	COR1B_MOUSE	22	0.12
G3X9C2	F-box only protein 50	FBX50_MOUSE	18	0.11
O09117	Synaptopnysin-like protein	SYPL1_MOUSE	10	0.11
	Complement component 1			
035658	Q subcomponent-binding	CIOBR MOUSE	1/	0.11
00000	Proliferating cell nuclear		17	0.11
P17918	antigen	PCNA_MOUSE	18	0.11
B 22020	Non-specific lipid-transfer		20	0.11
P32020			30	0.11
P40237	Acidic leucine-rich nuclear		11	0.11
P97822	phosphoprotein 32 family	AN32E_MOUSE	10	0.11

	member E			
	Tyrosine-protein kinase			
Q04736	Yes	YES_MOUSE	28	0.11
Q0P557	Mitochondria-eating protein	MIEAP_MOUSE	29	0.11
Q3TEA8	binding protein 3	HP1B3_MOUSE	33	0.11
Q3U1J4	protein 1	DDB1_MOUSE	53	0.11
Q60864	Stress-induced- phosphoprotein 1	STIP1_MOUSE	41	0.11
Q61792	LIM and SH3 domain protein 1	LASP1_MOUSE	16	0.11
Q8BFR4	N-acetylglucosamine-6- sulfatase	GNS_MOUSE	28	0.11
Q8BL97	Serine/arginine-rich splicing factor 7	SRSF7_MOUSE	17	0.11
Q8JZQ9	Eukaryotic translation initiation factor 3 subunit B	EIF3B_MOUSE	39	0.11
Q8K2I3	Dimethylaniline monooxygenase [N-oxide- forming] 2	FMO2 MOUSE	30	0.11
40.12.0	Cytochrome b-c1 complex			
Q9CR68	subunit Rieske, mitochondrial	UCRI_MOUSE	16	0.11
Q9D0F9	Phosphoglucomutase-1	PGM1_MOUSE	37	0.11
	Alpha-aminoadipic semialdehyde dehydrogenase	ALZA1 MOUSE	24	0.11
090706		TOUR MOUSE	14	0.11
0907F2	Glynican-1	GPC1_MOUSE	33	0.11
09W/V/K4	EH domain-containing	EHD1_MOUSE	33	0.11
0972C6	Uroplakin-1b	UPK1B MOUSE	12	0.11
035295	Transcriptional activator protein Pur-beta	PURB MOUSE	16	0.1
O55029	Coatomer subunit beta~	COPB2 MOUSE	49	0.1
	Puromycin-sensitive	—		
Q11011	aminopeptidase	PSA_MOUSE	53	0.1
Q61490	CD166 antigen	CD166_MOUSE	29	0.1
Q61699	kDa	HS105_MOUSE	54	0.1
Q64704	Syntaxin-3	STX3_MOUSE	14	0.1
Q7TSV4	Phosphoglucomutase-2	PGM2_MOUSE	36	0.1
Q8BJZ3	Protein lifeguard 3	LFG3_MOUSE	6	0.1
Q8CI51	PDZ and LIM domain protein 5	PDLI5_MOUSE	33	0.1
Q922Q4	Pyrroline-5-carboxylate reductase 2	P5CR2_MOUSE	17	0.1
Q99MN1	LysinetRNA ligase	SYK_MOUSE	30	0.1
Q9CQA3	Succinate dehydrogenase [ubiquinone] iron-sulfur subunit, mitochondrial	SDHB MOUSE	24	0.1
	Mitochondrial 2- oxoglutarate/malate carrier			-
Q9CR62	protein	M2OM_MOUSE	21	0.1

0000050	Translocon-associated	SSRA MOUSE	10	0.1
0.90130	Armadillo repeat-containing		10	0.1
Q9D0L7	protein 10	ARM10_MOUSE	13	0.1
Q9D710	Thioredoxin-related transmembrane protein 2	TMX2_MOUSE	18	0.1
Q9D711	Pirin	PIR MOUSE	19	0.1
090819	Inorganic pyrophosphatase	IPYR MOUSE	20	0.1
000010	Phosphatidylinositide		20	0.1
Q9EP69	phosphatase SAC1	SAC1_MOUSE	35	0.1
	Estradiol 17-beta-			
Q9EQ06	dehydrogenase 11	DHB11_MOUSE	17	0.1
A2ASQ1	Agrin	AGRIN_MOUSE	89	0.09
O08677	Kininogen-1	KNG1_MOUSE	30	0.09
O35887	Calumenin	CALU MOUSE	17	0.09
	ATP-dependent RNA			
O70133	helicase A	DHX9_MOUSE	75	0.09
	PDZ and LIM domain			
070400	protein 1	PDLI1_MOUSE	19	0.09
P10518	dehydratase	HEM2_MOUSE	15	0.09
	26S proteasome non-			
DOGEAG	ATPase regulatory subunit		45	0.00
P26516	/	PSMD7_MOUSE	15	0.09
P45377	protein 2	ALD2_MOUSE	20	0.09
Q61768	Kinesin-1 heavy chain	KINH MOUSE	64	0.09
	N(4)-(beta-N-	_		
	acetylglucosaminyl)-L-			
Q64191	asparaginase	ASPG_MOUSE	17	0.09
Q6ZWX6	Eukaryotic translation initiation factor 2 subunit 1	IF2A_MOUSE	22	0.09
	Calcium-binding			
0001150	mitochondrial carrier			
Q8BH59	Activator of 90 kDa boat		38	0.09
	shock protein ATPase			
Q8BK64	homolog 1	AHSA1 MOUSE	24	0.09
	NHL repeat-containing			
Q8CCH2	protein 3	NHLC3_MOUSE	20	0.09
0.01/0.07	Procollagen	OTOFA MOUNT	10	
Q8K297	galactosyltransferase 1	GT251_MOUSE	40	0.09
Q8VCT3	Aminopeptidase B	AMPB_MOUSE	33	0.09
Q91V61	Sideroflexin-3	SFXN3_MOUSE	20	0.09
002208	Transmembrane protein	T132A MOUSE	11	0.09
Q322F0	Leucine-rich repeat-	1132A_10003L	44	0.03
Q922Q8	containing protein 59	LRC59 MOUSE	18	0.09
	Stomatin-like protein 2,		20	0.00
			20	0.09
Q99KP3	Lambda-crystallin homolog	URILI_MOUSE	20	0.09
Q99I 45	initiation factor 2 subunit 2	IF2B MOUSE	22	0.09
	2,4-dienoyl-CoA reductase.			0.00
Q9CQ62	mitochondrial	DECR_MOUSE	23	0.09
Q9CS42	Ribose-phosphate	PRPS2_MOUSE	19	0.09
-	· · ·	•		*
	pyrophosphokinase 2			
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	ArgininetRNA ligase,			
Q9D0I9	cytoplasmic	SYRC_MOUSE	42	0.09
Q9D0M3	Cytochrome c1, heme protein, mitochondrial	CY1_MOUSE	16	0.09
	2-oxoglutarate and iron-			
	dependent oxygenase			
000126	domain-containing protein		17	0.00
Q9D130	Cysteine and histidine-rich	OGFD3_MOUSE	17	0.09
	domain-containing protein			
Q9D1P4	1	CHRD1_MOUSE	23	0.09
	Peptidoglycan recognition			
A1A547	protein 3	PGRP3_MOUSE	19	0.08
O08529	Calpain-2 catalytic subunit	CAN2_MOUSE	38	0.08
	Mitogen-activated protein			
O08911	kinase 12	MK12_MOUSE	23	0.08
O54984	ATPase Asna1	ASNA_MOUSE	15	0.08
	H-2 class I			
Dataat	histocompatibility antigen,		10	0.00
P01901	K-B alpha chain	HA1B_MOUSE	19	0.08
P30677	protein subunit alpha-14	GNA14 MOUSE	24	0.08
1 00077	Dual specificity mitogen-		27	0.00
	activated protein kinase			
P31938	kinase 1	MP2K1_MOUSE	18	0.08
	V-type proton ATPase			
P51863	subunit d 1	VA0D1_MOUSE	17	0.08
	Heterogeneous nuclear			
000PI9	notein 2	HNRL2 MOUSE	41	0.08
004447	Creating kingge B type		21	0.00
Q04447	Calcium uniporter protein	KCKD_WOUSE	21	0.08
Q3UMR5	mitochondrial	MCU MOUSE	17	0.08
	Serum			
Q62087	paraoxonase/lactonase 3	PON3_MOUSE	17	0.08
	Radial spoke head protein			
Q8BYM7	4 homolog A	RSH4A_MOUSE	27	0.08
Oskoco	GDP-mannose 4,6	CMDS MOUSE	20	0.08
			30	0.00
Q8R1F1	Niban-like protein 1	NIBL1_MOUSE	44	0.08
Q8R5C5	Beta-centractin	ACTY_MOUSE	23	0.08
0011/145	RNA binding motif protein,		22	0.09
QUIVIND	Dna Lhomolog subfamily B		33	0.00
Q99KV1	member 11	DJB11 MOUSE	21	0.08
	NADH dehydrogenase			
	[ubiquinone] 1 alpha			
	subcomplex subunit 10,			
Q99LC3		NDUAA_MOUSE	23	0.08
0000006	Interleukin ennancer-	ILE2 MOUSE	21	0.08
	Protein transport protein		<u> </u>	0.00
Q9D662	Sec23B	SC23B MOUSE	39	0.08
	Minor histocompatibility			
Q9D8V0	antigen H13	HM13_MOUSE	15	0.08
Q9DC69	NADH dehydrogenase	NDUA9_MOUSE	27	0.08

	[ubiquinone] 1 alpha subcomplex subunit 9			
	mitochondrial			
Q9ERH8	Solute carrier family 28 member 3	S28A3_MOUSE	28	0.08
Q9JK81	UPF0160 protein MYG1, mitochondrial	MYG1_MOUSE	27	0.08
	26S proteasome non-			
Q9WVJ2	A I Pase regulatory subunit	PSD13_MOUSE	26	0.08
O08917	Flotillin-1	FLOT1_MOUSE	31	0.07
O88632	Semaphorin-3F	SEM3F_MOUSE	51	0.07
O88968	Transcobalamin-2	TCO2_MOUSE	23	0.07
P06909	Complement factor H	CFAH_MOUSE	69	0.07
P09055	Integrin beta-1	ITB1 MOUSE	39	0.07
	cAMP-dependent protein			
	kinase type II-alpha			
P12367	regulatory subunit	KAP2_MOUSE	21	0.07
P19324	Serpin H1	SERPH_MOUSE	25	0.07
P37889	Fibulin-2	FBLN2_MOUSE	50	0.07
P46664	Adenylosuccinate synthetase isozyme 2	PURA2_MOUSE	27	0.07
P48722	Heat shock 70 kDa protein 4L	HS74L_MOUSE	60	0.07
	Platelet-activating factor			
P63005	acetyinydrolase IB subunit	LIST MOUSE	27	0.07
1 00000	DnaJ homolog subfamily A		21	0.07
P63037	member 1	DNJA1_MOUSE	22	0.07
P70168	Importin subunit beta-1	IMB1_MOUSE	38	0.07
P81117	Nucleobindin-2	NUCB2_MOUSE	24	0.07
P84091	AP-2 complex subunit mu	AP2M1 MOUSE	29	0.07
Q04998	Inhibin beta A chain	INHBA MOUSE	30	0.07
	Epidermal growth factor			
Q08509	receptor kinase substrate 8	EPS8_MOUSE	45	0.07
OOTLICC	S-adenosylmethionine		04	0.07
Q31H50	Casein kinase II subunit	METK2_WOUSE	21	0.07
Q60737	alpha	CSK21_MOUSE	21	0.07
	KH domain-containing,			
	RNA-binding, signal			
060749	transduction-associated		22	0.07
00749	Histone-binding protein		22	0.07
Q60972	RBBP4	RBBP4_MOUSE	15	0.07
Q61092	Laminin subunit gamma-2	LAMC2_MOUSE	69	0.07
	Cullin-associated NEDD8-			
Q6ZQ38	dissociated protein 1	CAND1_MOUSE	66	0.07
Q80YQ8	Protein RMD5 homolog A	RMD5A_MOUSE	26	0.07
Q8BGX1	PC-esterase domain- containing protein 1B	PED1B_MOUSE	22	0.07
Q8BWN8	Acyl-coenzyme A thioesterase 4	ACOT4_MOUSE	21	0.07
Q8QZT1	Acetyl-CoA acetyltransferase,	THIL_MOUSE	23	0.07

	mitochondrial			
	Regulator of chromosome			
Q8VE37	Condensation	RCC1_MOUSE	18	0.07
	inhibitor 1 RNA-binding			
Q9CY58	protein	PAIRB_MOUSE	23	0.07
	CysteinetRNA ligase,	—		
Q9ER72	cytoplasmic	SYCC_MOUSE	51	0.07
Q9WVP1	AP-1 complex subunit mu-2	AP1M2_MOUSE	28	0.07
	Serine/threonine-protein			
Q9Z2W1	kinase 25	STK25_MOUSE	23	0.07
O08528	Hexokinase-2	HXK2_MOUSE	55	0.06
	116 kDa U5 small nuclear			
O08810	component	U5S1 MOUSE	56	0.06
000010	Lysosome membrane		00	0.00
O35114	protein 2	SCRB2_MOUSE	24	0.06
	Splicing factor U2AF 65			
P26369	kDa subunit	U2AF2_MOUSE	19	0.06
D26629	SerinetRNA ligase,	SVSC MOUSE	20	0.06
F20030		STOC_MOUSE	29	0.00
P30999	Catenin delta-1	CIND1_MOUSE	55	0.06
	bydroxymethyltransferase			
P50431	cytosolic	GLYC_MOUSE	28	0.06
	Ras GTPase-activating			
P97855	protein-binding protein 1	G3BP1_MOUSE	25	0.06
Dogoco	Bone morphogenetic		50	0.00
P98063	F3 ubiquitin-protein ligase	BIMPT_MOUSE	58	0.06
Q0MW30	NEURL1B	NEU1B MOUSE	23	0.06
	Keratin, type II cytoskeletal	—		
Q0VBK2	80	K2C80_MOUSE	33	0.06
000170	Transmembrane channel-	THOS MOULOS	10	0.00
Q32NZ6	IIKe protein 5		42	0.06
Q3TJD7	protein 7	PDLI7 MOUSE	28	0.06
05X.IY5	Coatomer subunit delta		29	0.06
061235	Beta-2-syntrophin	SNTB2 MOUSE	30	0.06
001235	ATP-dependent RNA	SINTB2_WOUSE	30	0.00
Q61655	helicase DDX19A	DD19A_MOUSE	34	0.06
	MethioninetRNA ligase,			
Q68FL6	cytoplasmic	SYMC_MOUSE	49	0.06
	Serine/threonine-protein		07	0.00
Q0P9R2	Rinase OSR I Res-related GTP-binding		21	0.06
Q7TT45	protein D	RRAGD MOUSE	23	0.06
	Calcium-activated chloride			
Q8BG22	channel regulator 2	CLCA2_MOUSE	45	0.06
	Sorting and assembly			
	machinery component 50	SAMED MOUSE	20	0.06
	AlaninetRNA ligase		29	0.00
Q8BGQ7	cytoplasmic	SYAC MOUSE	56	0.06
	V-type proton ATPase			
Q8BVE3	subunit H	VATH_MOUSE	27	0.06
Q8BXZ1	Protein disulfide-isomerase	TMX3_MOUSE	25	0.06

	TMX3			
	Endonuclease domain-			
Q8C522	containing 1 protein	ENDD1_MOUSE	23	0.06
	glutamate/prolinetRNA			
Q8CGC7	ligase	SYEP_MOUSE	94	0.06
000000	Lon protease homolog,	LONM MOUSE	50	0.06
			12	0.06
QORUE7	NADH dehvdrogenase	SERC2_WOUSE	12	0.06
	[ubiquinone] flavoprotein 1,			
Q91YT0	mitochondrial	NDUV1_MOUSE	27	0.06
Q99LD4	complex subunit 1	CSN1 MOUSE	26	0.06
Q9CWK8	Sorting nexin-2	SNX2 MOUSE	30	0.06
	Cytosolic non-specific			
Q9D1A2	dipeptidase	CNDP2_MOUSE	26	0.06
	Cysteine sulfinic acid		21	0.06
			47	0.00
Q9DBG3	AP-2 complex subunit beta	AP2B1_MOUSE	47	0.06
Q9DBX6	Cytochrome P450 2S1	CP2S1_MOUSE	28	0.06
Q9ES97	Reticulon-3	RTN3_MOUSE	45	0.06
Q9JHU4	heavy chain 1	DYHC1 MOUSE	283	0.06
Q9JIF7	Coatomer subunit beta	COPB MOUSE	56	0.06
	Fas-activated			
Q9JIX9	serine/threonine kinase	FASTK_MOUSE	30	0.06
	4- trimethylaminobutyraldebyd			
Q9JLJ2	e dehydrogenase	AL9A1_MOUSE	27	0.06
	Protein transport protein			
	Sec61 subunit alpha	S61A2 MOUSE	17	0.06
	Coronin 10	CORIC MOUSE	24	0.00
0900004	Fatty-acid amide hydrolase		24	0.00
O08914	1	FAAH1_MOUSE	26	0.05
070404	Eukaryotic translation		07	0.05
070194	Initiation factor 3 subunit D	EIF3D_MOUSE	27	0.05
089023	Tripeptidyl-peptidase 1	TPP1_MOUSE	22	0.05
P29416	subunit alpha	HEXA MOUSE	25	0.05
	Serine/threonine-protein			
D 40 450	phosphatase 2B catalytic		07	0.05
P48453	Subunit beta isoform	PP2BB_MOUSE	27	0.05
	acyl-CoA dehydrogenase,			
P50544	mitochondrial	ACADV_MOUSE	43	0.05
P51655	Glypican-4	GPC4_MOUSE	29	0.05
D07407	Dihydropyrimidinase-		22	0.05
P97427			33	0.05
000500			29	0.05
260298	Prolyl 4-hydroxylase	SKUD_NIUUSE	20	0.05
Q60715	subunit alpha-1	P4HA1_MOUSE	29	0.05
Q61545	RNA-binding protein EWS	EWS_MOUSE	23	0.05

Q64133	Amine oxidase [flavin- containing] A	AOFA MOUSE	30	0.05
Q64455	Receptor-type tyrosine-	PTPRJ MOUSE	54	0.05
Q6DFW4	Nucleolar protein 58	NOP58 MOUSE	27	0.05
	E3 ubiquitin-protein ligase			0.00
Q6PDK8	DTX4	DTX4_MOUSE	32	0.05
Q80UG5	Septin-9	SEPT9_MOUSE	40	0.05
Q8BH04	Phosphoenolpyruvate carboxykinase [GTP], mitochondrial	PCKGM_MOUSE	35	0.05
Q8BU30	IsoleucinetRNA ligase, cytoplasmic	SYIC_MOUSE	67	0.05
Q8CHT0	Delta-1-pyrroline-5- carboxylate dehydrogenase, mitochondrial	AL4A1_MOUSE	32	0.05
Q8CIE6	Coatomer subunit alpha	COPA_MOUSE	76	0.05
Q8R0X7	Sphingosine-1-phosphate lyase 1	SGPL1_MOUSE	30	0.05
Q8VCM8	Nicalin	NCLN_MOUSE	22	0.05
Q91V92	ATP-citrate synthase	ACLY_MOUSE	61	0.05
Q91WG0	Acylcarnitine hydrolase	EST2C MOUSE	17	0.05
0.001/0.7	Lipolysis-stimulated			
Q99KG5	lipoprotein receptor	LSR_MOUSE	26	0.05
Q9D6Z1	Nucleolar protein 56	NOP56_MOUSE	36	0.05
Q9DBG7	receptor subunit alpha	SRPR_MOUSE	41	0.05
Q9ET30	Transmembrane 9 superfamily member 3	TM9S3_MOUSE	21	0.05
Q9R069	molecule	BCAM MOUSE	35	0.05
E9Q557	Desmoplakin	DESP MOUSE	207	0.04
O35350	Calpain-1 catalytic subunit	CAN1 MOUSE	45	0.04
P19096	Fatty acid synthase	FAS MOUSE	131	0.04
	Eukaryotic translation			
P23116	initiation factor 3 subunit A	EIF3A_MOUSE	90	0.04
P23780	Beta-galactosidase	BGAL_MOUSE	28	0.04
P41216	Long-chain-fatty-acidCoA ligase 1	ACSL1_MOUSE	39	0.04
P70670	Nascent polypeptide- associated complex subunit alpha, muscle-specific form	NACAM_MOUSE	102	0.04
P82198	Transforming growth factor- beta-induced protein ig-h3	BGH3_MOUSE	42	0.04
Q3UPP8	Centrosomal protein of 63 kDa	CEP63_MOUSE	40	0.04
Q3UQ44	Ras GTPase-activating-like protein IQGAP2	IQGA2_MOUSE	96	0.04
Q62318	Transcription intermediary factor 1-beta	TIF1B_MOUSE	33	0.04
0001/40	N- acetylgalactosaminyltransfe		45	
Q80VA0		GALI7_MOUSE	45	0.04
Q8C1A5	I nimet oligopeptidase	IHOP1_MOUSE	45	0.04

090014	Complement C1s-A		20	0.04
	subcomponent		20	0.04
Q8JZQ2	AFG3-like protein 2	AFG32_MOUSE	53	0.04
Q8JZR0	ligase 5	ACSL5_MOUSE	37	0.04
Q8K0D5	Elongation factor G, mitochondrial	EFGM_MOUSE	45	0.04
Q8R146	Acylamino-acid-releasing enzyme	APEH_MOUSE	36	0.04
	Choline transporter-like			
Q91VA1	protein 4	CTL4_MOUSE	27	0.04
Q91VD9	oxidoreductase 75 kDa subunit, mitochondrial	NDUS1_MOUSE	42	0.04
Q91W50	Cold shock domain- containing protein E1	CSDE1 MOUSE	55	0.04
	Nuclear autoantigenic	—		
Q99MD9	sperm protein	NASP_MOUSE	43	0.04
Q9D0R2	ThreoninetRNA ligase, cytoplasmic	SYTC_MOUSE	47	0.04
000000	Acetoacetyl-CoA		20	0.04
Q9D2R0	Solute carrier family 15	AACS_MOUSE	38	0.04
Q9ES07	member 2	S15A2_MOUSE	29	0.04
Q9JHE3	Neutral ceramidase	ASAH2 MOUSE	25	0.04
	Calcium-binding			
	mitochondrial carrier			
Q9QXX4	protein Aralar2	CMC2_MOUSE	37	0.04
Q9QYB5	Gamma-adducin	ADDG_MOUSE	31	0.04
	Procollagen-lysine,2- oxoglutarate 5-dioxygenase	PLOD1 MOUSE	43	0.04
	Cadharin-13		33	0.04
QUIND	Delta-1-pyrroline-5-	CADIS_MOUSE	33	0.04
Q9Z110	carboxylate synthase	P5CS_MOUSE	48	0.04
097185	type-1	PADI1 MOUSE	36	0.04
Q02100	LETM1 and EF-hand		00	0.01
	domain-containing protein			
Q9Z2I0	1, mitochondrial	LETM1_MOUSE	45	0.04
B0\/2N1	Receptor-type tyrosine-	DTDDS MOUSE	96	0.03
	Kinagin like protoin KIE22		50	0.03
E9Q5G3	AP-2 complex subunit	KIF23_MOUSE	53	0.03
P17426	alpha-1	AP2A1 MOUSE	60	0.03
P39054	Dynamin-2	DYN2 MOUSE	56	0.03
P43406	Integrin alpha-V	ITAV MOUSE	61	0.03
1 10100	Ubiquitin carboxyl-terminal			0.00
P56399	hydrolase 5	UBP5_MOUSE	43	0.03
P56677	Suppressor of tumorigenicity 14 protein homolog	ST14 MOUSE	36	0.03
007139	Protein FCT2	FCT2 MOUSE	53	0.03
307133	Dolichyl-		33	0.00
	diphosphooligosaccharide			
	protein glycosyltransferase			
Q3TDQ1	subunit STT3B	STT3B_MOUSE	35	0.03

Q60997	Deleted in malignant brain tumors 1 protein	DMBT1 MOUSE	55	0.03
Q61739	Integrin alpha-6	ITA6 MOUSE	62	0.03
	Exportin 1	YPO1 MOUSE	52	0.03
QUESTS	Staphylococcal nuclease		50	0.03
Q78PY7	domain-containing protein	SND1_MOUSE	58	0.03
Q8BFY9	Transportin-1	TNPO1_MOUSE	38	0.03
Q8BIJ6	IsoleucinetRNA ligase, mitochondrial	SYIM MOUSE	49	0.03
	Piwi-like protein 2	PIWL2 MOUSE	66	0.03
000001	E3 ubiguitin-protein ligase		00	0.00
Q8CFI0	NEDD4-like	NED4L_MOUSE	54	0.03
Q8K310	Matrin-3	MATR3_MOUSE	46	0.03
Q8VDM6	Heterogeneous nuclear ribonucleoprotein U-like protein 1	HNRL1_MOUSE	45	0.03
Q99MR6	molecule homolog	SRRT MOUSE	49	0.03
Q99P72	Beticulon-4	RTN4 MOUSE	45	0.03
	Calsyntenin-1	CSTN1 MOUSE	10	0.03
QUEFLZ	Endoplasmic reticulum		41	0.03
Q9EQH2	aminopeptidase 1	ERAP1_MOUSE	52	0.03
Q9JIK5	Nucleolar RNA helicase 2	DDX21_MOUSE	52	0.03
Q9Z1T1	AP-3 complex subunit beta-	AP3B1_MOUSE	64	0.03
121915	Receptor-type tyrosine-	DTDDE MOUSE	105	0.02
R2R0LJ			07	0.02
DZKA34	Plexin-b2 Plasma membrane	PLADZ_WOUSE	97	0.02
G5E829	calcium-transporting ATPase 1	AT2B1_MOUSE	65	0.02
P02468	Laminin subunit gamma-1	LAMC1_MOUSE	83	0.02
P11087	Collagen alpha-1(I) chain	CO1A1 MOUSE	82	0.02
P59509	A disintegrin and metalloproteinase with thrombospondin motifs 19	ATS19_MOUSE	63	0.02
Q05793	Basement membrane- specific heparan sulfate proteoglycan core protein	PGBM_MOUSE	151	0.02
Q3UPL0	Protein transport protein Sec31A	SC31A_MOUSE	53	0.02
Q3UTJ2	Sorbin and SH3 domain- containing protein 2	SRBS2_MOUSE	70	0.02
Q5ND34	WD repeat-containing protein 81	WDR81_MOUSE	91	0.02
Q61001	Laminin subunit alpha-5	LAMA5_MOUSE	158	0.02
Q61595	Kinectin	KTN1_MOUSE	90	0.02
Q62315	Protein Jumonji	JARD2 MOUSE	68	0.02
Q64511	DNA topoisomerase 2-beta	TOP2B MOUSE	100	0.02
Q68FD7	Folliculin-interacting protein	FNIP1_MOUSE	68	0.02
Q6F3F9	G-protein coupled receptor 126	GP126_MOUSE	45	0.02

	UDP-glucose:glycoprotein	LIGGG1 MOUSE	84	0.02
QUF JL4	Intraflagellar transport	00001_00002	04	0.02
Q6VH22	protein 172 homolog	IF172_MOUSE	116	0.02
Q7TPV4	Myb-binding protein 1A	MBB1A_MOUSE	78	0.02
Q80XI3	Eukaryotic translation initiation factor 4 gamma 3	IF4G3_MOUSE	97	0.02
Q8BMJ2	LeucinetRNA ligase, cytoplasmic	SYLC_MOUSE	61	0.02
Q921M3	Splicing factor 3B subunit 3	SF3B3_MOUSE	60	0.02
Q9QXS1	Plectin	PLEC_MOUSE	342	0.02
Q07113	Cation-independent mannose-6-phosphate receptor	MPRI_MOUSE	124	0.01
Q61789	Laminin subunit alpha-3	LAMA3_MOUSE	181	0.01
Q6P4T2	U5 small nuclear ribonucleoprotein 200 kDa helicase	U520_MOUSE	122	0.01
Q8C6K9	Collagen alpha-6(VI) chain	CO6A6_MOUSE	127	0.01
Q8R0W0	Epiplakin	EPIPL_MOUSE	388	0.01
Q99PV0	Pre-mRNA-processing- splicing factor 8	PRP8_MOUSE	145	0.01
Q9D952	Envoplakin	EVPL_MOUSE	126	0.01
Q9QZZ4	Unconventional myosin-XV	MYO15_MOUSE	188	0.01
A2AAJ9	Obscurin	OBSCN_MOUSE	495	0

Primary Antibody	Туре	Manufacturer
Anti- Bpifa1	Rabbit polyclonal	Prepared in lab (Musa et al 2012)
Anti- FoxJ1	Mouse monoclonal	Affymetrix eBioscience (2A5 Cat no 14-9965)
Anti- Muc5B	Rabbit polyclonal	Santa Cruz Biotechnology (H-300, Sc-20119)
Anti- P63	Mouse monoclonal	Santa Cruz Biotechnology (4A4, Cat no Sc- 8431)
Anti- Lactotransferrin	Rabbit antisera	Milipore (Cat no 07- 685)
Anti- Reg3γ	Rabbit polyclonal	Kindly provided by Professor Lora Hooper (Cash et al 2006)
Anti- Zo1	Rabbit polyclonal	Life Technologies (Cat no 40-2200)

Table S2: Antibodies used for immuno-detection studies

Assay	Primer type	Sequence (5'-3')	Product size (bp)
Bpifa1	Forward Reverse	ACAGAGGAGCCGACGTCTAA CCAAGAAAGCTGAAGGTTC	127
Tekt1	Forward Reverse	CAGTGCGAAGTGGTAGACG TTCACCTGGATTTCCTCCTG	373
Muc5B	Forward Reverse	ATGGTGACCAAGAGCCAAAA CAGGACTGTTCACCCAGGTT	178
Muc5AC	Forward Reverse	GCACCAAAGACAGCAGATCA CTGAGAGGTTTGCAGCTCCT	167
Lactotransferrin	Forward Reverse	TCTGTCCCTGTGTATTGGT GTTTCCGGGTGTCATCAAGG	237
SP-D	Forward Reverse	AAGCAGGGGAACATAGGACC GCCTTTTGCCCCTGTAGATC	109
Bpifb1	Forward Reverse	CCCTGACCAAGATCCTTGAA GAGGCTGGAGTGAGCTTGAG	148
Keratin 5	Forward Reverse	ACCTTCGAAACACCAAGCAC TGACTGGTCCAACTCCTTCC	337
Vimentin	Forward Reverse	CAAGCAGGAGTCAAACGAG CCTGTAGGTGGCGATCTCAA	273
Reg3 ^γ	Forward Reverse	CCTGTCCTCCATGATCAAAGC GCAGACATAGGGTAACTCTAAGT	250
Oaz1	Forward Reverse	ACAGAGGAGCCGACGTCTAA CCAAGAAAGCTGAAGGTTC	274

Table S3: Primer sequences and amplicon sizes for end point reversetranscription PCR