RESEARCH ARTICLE

The EDA-deficient mouse has Zymbal’s gland hypoplasia and acute otitis externa
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ABSTRACT

In mice, rats, dogs and humans, the growth and function of sebaceous glands and eyelid Meibomian glands depend on the ectodysplasin signalling pathway. Mutation of genes encoding the ligand EDA, its transmembrane receptor EDAR and the intracellular signal transducer EDARADD leads to hypohidrotic ectodermal dysplasia, characterised by impaired development of teeth and hair, as well as cutaneous glands. The rodent ear canal has a large auditory sebaceous gland, the Zymbal’s gland, the function of which in the health of the ear canal has not been determined. We report that EDA-deficient mice, EDAR-deficient mice and EDARADD-deficient rats have Zymbal’s gland hypoplasia. EdaTa−/− mice have 25% prevalence of otitis externa at postnatal day 21 and treatment with agonist anti-EDAR antibodies rescues Zymbal’s glands. The aetiopathogenesis of otitis externa involves infection with Gram-positive cocci, and dosing pregnant and lactating EdaTa females and pups with enrofloxacin reduces the prevalence of otitis externa. We infer that the deficit of sebum is the principal factor in predisposition to bacterial infection, and the EdaTa−/− mouse is a potentially useful microbial challenge model for human acute otitis externa.

KEY WORDS: Hypohidrotic ectodermal dysplasia, Sparse and wavy hair rat, EDARADD, FBXO11, MECOM, EDAR, Tabby mouse

INTRODUCTION

Human acute otitis externa (AOE) is a common microbial infection of the ear canal (external acoustic meatus), with an estimated 1.72 million cases in the US in 2014 and an estimated annual treatment cost of $564 million (Collier et al., 2021). The ear canal is the only cul-de-sac keratinising skin surface in the body and is self-cleansing through the production of cerumen (wax), which is a mixture of desquamated keratinocytes and glandular secretions (Guest et al., 2004). The human ear canal has sebaceous glands associated with hair follicles (pilosebaceous units) that produce lipid-rich sebum (Bortz et al., 1990; Guest et al., 2004), and ceruminous glands, which are modified apocrine glands that secrete fluid rich in antibacterial peptides (Main and Lim, 1976; Stoeckelhuber et al., 2006). Wetting of the ear canal through swimming, bathing or high environmental humidity compromises these defences and predisposes the canal to microbial infection, giving rise to its common name ‘swimmer’s ear’.

Animal models of AOE generally involve disrupting the ear canal epithelial barrier by mechanical abrasion, sustained wetting or chemical irritants and inoculation of a microbial pathogen. Animal models include rats (Emgård and Hellström, 1997, 2001; Emgård et al., 2005; Demirel et al., 2018), guinea pigs (Wright and Dineen, 1972; King and Estrem, 1990; Zhai et al., 2014) and mice (Wright et al., 2000). However, spontaneous otitis externa is not a notable disease of laboratory rats, guinea pigs or mice in lab animal medicine texts (Fox et al., 2015).

Rodents lack the apocrine ceruminous glands that are found in human, dog, goat and pig ear canals (Wang et al., 2021), but have a specialised large multilobulated auditory sebaceous gland, also known as the Zymbal’s gland (Rudmann et al., 2012), which opens via a duct into the ear canal close to the tympanic membrane. The Zymbal’s gland is also called the ear-wax gland (glandula ceruminosa) (Grüneberg, 1971) or ceruminous gland (Berry et al., 1971). Hereafter, we use the term Zymbal’s gland to draw a distinction between this specialised sebaceous holocrine gland and the human apocrine ceruminous gland. The ear canal carries sound waves received by the outer ear, the pinna, towards the tympanic membrane, and in the mouse the canal is formed by a short osseous part, the ectotympanic ring, and a short annular cartilage (Navarro et al., 2017). Vibrations of the tympanic membrane are transmitted through the air-filled auditory bulla (middle ear) to the inner ear by the ossicular chain.

Sebaceous glands develop either as paired outgrowths of the hair follicle or in specialised glands, such as the Meibomian, preputial, clitoral and Zymbal’s glands, which develop independently of the hair follicle (Dhouflaiy and Oftedal, 2016). Preputial glands develop at embryonic day (E)14.5 as placodes in the epidermis of the genital tubercle, and the Meibomian gland placodes develop at the fused eyelid margins at E18.5. The embryological development of the Zymbal’s gland is less well studied (Dhouflaiy and Oftedal, 2016) but the primordium is reported to be present at E15 (Grüneberg, 1971).

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The growth and function of hair follicle sebaceous glands and Meibomian glands are dependent on the ectodysplasin signalling pathway, which comprises a TNF-like ligand ectodysplasin (EDA), its transmembrane receptor EDAR and the intracellular signal transducer EDARADD. The loss of signalling due to mutation of genes in this linear pathway leads to hypohidrotic ectodermal dysplasia (HED) and impairs the development of teeth and hair, as well as cutaneous glands (Kowalczyk-Quintas and Schneider, 2014). The Tabby (EdaTa) mouse is a model of X-linked HED (XLHED) and is deficient in EDA. Treatment of adult EdaTa mice with agonist anti-EDAR antibody (Kowalczyk et al., 2011; Kowalczyk-Quintas et al., 2014) restores sebaceous gland growth and sebum production, and sustained treatment of EdaTa and wild-type mice heightens sebum production (Kowalczyk-Quintas et al., 2015). In addition, prenatal correction of XLHED with a recombinant protein that includes the receptor binding domain of EDA restores Meibomian gland growth in humans (Schneider et al., 2018) and dogs (Margolis et al., 2019). Adult mouse sebaceous glands express the EDAR gene, suggesting the action of the agonist anti-EDAR antibody treatment is directly on the sebaceous glands (Kowalczyk-Quintas et al., 2015). The rat Zymbal’s gland expresses the EDARADD gene (del-Pozo et al., 2019a), so it is also likely to be stimulated directly by EDAR signalling.

Growth retardation of Meibomian glands causes dry eye, keratitis, ulceration and corneal neovascularization in EdaTa mice. The aetiopathogenesis is related to loss of lipid secretion altering tear film stability, a reduction in goblet cell density and increased susceptibility to eyelid and conjunctival inflammation through loss of EDA function (Cui et al., 2005; Wang et al., 2016). Furthermore, the Meibomian gland secretes EDA, and this is important for corneal epithelial homeostasis (Li et al., 2017). EDA signalling is also important for lacrimal gland development and function, and also contributes to corneal homeostasis and repair (Kuony et al., 2019).

The Zymbal’s gland in the rat and mouse is known principally from toxicology studies as a target for chemical carcinogens (Gold et al., 2001; Rudmann et al., 2012), and its normal physiological role is not documented. Postnatal day (P)10 EdaTa mice have Zymbal’s glands that are ∼5% of normal size and fewer ear canal hair follicles, which have smaller sebaceous glands than wild-type mice (Grüneberg, 1971), but the functional consequences of these deficits have not been investigated. In this study, we have investigated Zymbal’s gland growth and the health of the ear canal in the EdaTa mouse and in the EDARADD-deficient short and wavy hair rat (Edaraddpock/whiwh strain (Kuramoto et al., 2005, 2011). The EdaTa mouse and the Edaraddpock/whiwh rat have deficits in the nasopharyngeal submucosal glands, which protect the auditory (eustachian) tube, and this predisposes them to otitis media, infection and inflammation of the auditory bulla. Prenatal treatment of EdaTa mice with agonist anti-EDAR antibody rescues the submucosal glands and prevents otitis media (del-Pozo et al., 2019a).

The human tympanic membrane can perforate in recurrent acute otitis media (Folino et al., 2021) and in acute and chronic suppurrative otitis media (Mozaffari et al., 2020). Drainage of mucopurulent exudate though a tympanic membrane perforation (ototrauma) can result in the infection of the ear canal and AOE (Rosenfeld et al., 2014). Otitis externa is common in dogs, accounting for 7-10% of all canine cases seen in veterinary practice (O’Neill et al., 2014, 2021), and tympanic membrane perforation/otitis media is considered to be a common perpetuating cause of chronic otitis externa (Saridomicheakis, et al., 2007; Lorek et al., 2020). To evaluate the potential impact of otitis media on the tympanic membrane, and on the health of ear canal, we included two additional mouse models of chronic otitis media as an outgroup comparison. The aetiopathogeneses of otitis media in the Fbxo11−/− mouse is associated with a bulla cavitational defect (del-Pozo et al., 2019b), and the MecomPouieR mice has dysregulated NFkB inflammatory responses (Xu et al., 2012).

We found that EdaTa and EdarOveiOvei mice, and Edaraddpock/whiwh rats, have Zymbal’s gland hypoplasia. In addition, the EdaTa mouse has hypoplasia of ear canal pilosebaceous units (hypotrichosis) and is predisposed to bacterial otitis externa. The Zymbal’s glands and ear canal pilosebaceous units in P21 EdaTa mice are rescued by treatment with agonist anti-EDAR antibody. Furthermore, treatment with the broad-spectrum antibiotic enrofloxacin reduces the prevalence of otitis externa in P21 EdaTa mice.

**RESULTS**

**Zymbal’s gland hypoplasia in EdaTa mice**

The mouse external auditory meatus opens and extends on E12.5 (Mallo et al., 2000; Minoux et al., 2013), then closes between E15-P7 before re-opening from P7 to P12, and is fully opened at P12 (Anthwal and Thompson, 2016). We observed that the P9 Zymbal’s gland has a widely patent duct that opens into a narrowly canalised external auditory meatus (Fig. 1A,B).

The Zymbal’s gland is located between the ectotympanic ring and the annular cartilage on the rostral surface of the ear canal, and we measured EdaTa Zymbal’s tissue in this defined region. Hair follicles are reduced in number in EdaTa ear canal skin (Grüneberg, 1971) but we were careful to exclude any sebaceous glands associated with a hair follicle. Examination of serial step sections did not reveal Zymbal’s glands in 8 of 17 P21 EdaTa ear canals (Fig. 1E,F) but those detected were ∼8% the size of P21 FVB mouse controls (Fig. 1C,D,G,H,M). Depending on the plane of section, EdaTa Zymbal’s glands appeared as variably sized sebocyte lobules (Fig. 1G,H) and ducts connecting with the skin surface (Fig. S4P). Sebocyte lobules in P21 EdaTa Zymbal’s glands were significantly larger (median 10,494 µm², 95% c.i. 3584-17,786 µm², n=32) than P21 FVB hair follicle sebaceous glands (median 1273 µm², 95% c.i. 1034-1747 µm², n=33; P<0.0001, Mann–Whitney test). In contrast to P21 mice, all P79-P90 EdaTa had histologically unremarkable Zymbal’s glands, and although these were significantly larger than at P21, they were only ∼50% the size of Zymbal’s glands in P81-P84 FVB mice (Fig. 1K-M).

EdaTa, MecomPouieR, and Fbxo11−/− mutants all have middle ear pathology but its importance in the development of ear canal disease is unknown. To help differentiate the contributions of middle ear pathology and reduced Zymbal’s gland size in EdaTa mice, we compared gland size between mutant strains and found P22 MecomPouieR and P21 Fbxo11−/− mice have larger Zymbal’s glands than in P21 EdaTa mice (Fig. 1I,J,N).

**Otitis externa in P21 EdaTa mice**

Six of twelve P21 EdaTa mice had unilateral otitis externa (see Materials and Methods for diagnostic criterion) (Fig. 2A-H). These cases occurred in two litters of five and seven pups born to different parents; its prevalence of 25% (6 of 24 ears) was significantly elevated compared with the control group of P21 FVB mice in which otitis externa was absent (n=26 ears; P=0.0085, Fisher’s Exact test). In all cases of otitis externa, the neutrophil-rich suppurrative exudates contained Gram-positive cocci (Fig. 2G,H), but no fungi were detected using either PAS or Grocott stains.
Otitis externa was not observed in P79-90 EdaTa (n=52 ears) or in P81-P84 FVB (n=20 ears). The ear canal of some P79-P90 EdaTa mice can contain a thick plug of bland squamous epithelial cells that lack an inflammatory component (Fig. 2M-P).

The occurrence of otitis externa was assessed in MecomJbo/+ and Fbxo11Jf/+ mice. Neither mutants or wild-type littermates had otitis externa at weaning age or as older adults [MecomJbo/+ (P22, n=14; P84, n=12 ears); Mecom+/+ (P22, n=14 ears); Fbxo11Jf/+ (P21, n=13; P57-P223, n=41 ears); Fbxo11+/+ mice (P21, n=10 ears)].

Thirty-eight percent (6 of 16) of P21 EdaTa ears unaffected by otitis externa had a thin crust of desquamated epithelial cells, neutrophils and Gram-positive cocci on the external surface of the tympanic membrane (Fig. 2I-L); these crusts were <10% of the size of otitis externa exudate accumulations (Fig. S1A), and there was no accompanying inflammatory thickening of the tympanic membrane (Fig. 2I,J). Ears with tympanic membrane crusts had thickened ear canal soft tissue (Fig. S1B) but dermal and intra-epithelial neutrophil infiltration were also absent. Tympanic membrane crusts comprising desquamated cells and neutrophils (but not bacteria) were found in 21% (3 of 14) of P22 MecomJbo/+ ears but were absent in P22 Fbxo11Jf/+ (n=10) and wild-type control ears (P21 FVB, n=26; P22 Mecom+/+, n=14; P21 Fbxo11+/+, n=14).

**Ear canal hypotrichosis in EdaTa mice**

The skin of the osseous ear canal was haired in P21 and P81-P84 FVB mice (Fig. 3A-C,F), but was sparsely haired in P21 and P79-P90 EdaTa mice (Fig. 3D,G,P). The skin over the annular cartilage was thin and sparsely haired (Fig. 3A,J,K). No apocrine glands were found in the osseous or cartilaginous regions of the ear canal in mice P21 FVB mice (n=6).
Partial restoration of Zymbal’s gland growth and ear canal pilosebaceous unit density in EdaTa mice with agonist anti-EDAR antibody treatment

EdaTa mice treated prenatally or prenatally and postnatally with agonist anti-EDAR antibody had a higher density of ear canal pilosebaceous units at P21 (Fig. 3D,E,Q) and larger Zymbal’s glands than untreated (Fig. 3L,R) or antibiotic (enrofloxacin) mice (Fig. 3J,K). Prenatal agonist anti-EDAR antibody treatment rescued ear canal pilosebaceous units in P85 EdaTa mice (Fig. 3H,Q). However, post-weaning growth of Zymbal’s glands continued independently of EDA-EDAR signalling, and gland size in mice treated prenatally with agonist anti-EDAR antibody was no greater in P85 EdaTa mice than in untreated P79-P90 mice or P82-P84 mice treated with isotype antibody control (Fig. 3M-O). Furthermore, the density of ear canal pilosebaceous units was low in isotype antibody-treated P82-P84 EdaTa mice (Fig. 3I,Q).

Association between otitis externa susceptibility and small Zymbal’s gland size

Otitis externa and tympanic membrane crusts were not observed in agonist anti-EDAR treated P21 EdaTa mice (n=8). In contingency tests, the prevalence of otitis externa in treated mice was statistically lower than in untreated P21 controls (n=12, Table S1, Fig. S2A). Furthermore, small Zymbal’s gland size in weaning-aged mice was strongly associated with otitis externa [six of 12 affected P21 mice versus 0 of 45 P79-P90 EdaTa mice (untreated, and those administered agonist anti-EDAR or isotype antibodies); P=0.0000255, Fisher’s Exact test].

Co-existence of otitis media and otitis externa in P21 EdaTa ears

To gauge the importance of otitis media in the development of otitis externa at P21 and P22, we made an outgroup comparison and found that the prevalence of otitis media was higher in otitis media-sensitive, but otitis externa-resistant, mouse strains MecomJb/w+ and Fbxo11Jf/+ mice (13 of 14, and 9 of 10 ears, respectively) than in EdaTa mice (in which 9 of 24 ears were affected with otitis media). Nonetheless, there was a statistical association between otitis media and otitis externa in individual P21 EdaTa ears (Table S1). However, middle ear inflammation was characterised by mild serous effusion with scant leukocytes, whereas inflammation in the ear canal was severe and supplicative, with intraluminal cocci; there was no evidence of tympanic membrane perforation or cocci in the middle ear cavity.

Fig. 2. P21 EdaTa mice have otitis externa. Dorsal plane H&E- and Gram-stained sections through the bulla and ear canal of EdaTa mice. The image upper margin is rostral, and the left is lateral. A and B, C and D, E and F, I and J, M and N, and O and P are paired low and high magnification images of the same section. (A-E) Otitis externa. The tympanic membrane is intact and the ear canal contains a plug of exudate comprising neutrophils with squamous cells. (D) Ear canal dermal and epithelial inflammation (arrowheads mark the margins of epithelial erosion). (E) Thickened inflamed tympanic membrane pars tensa (arrowheads) with exudate on its external surface. Hyperplastic keratinising epithelium and inflammatory cell infiltration of the lamina propria can be observed. (F) Normal tympanic membrane in an unaffected ear. (G,H) Examples of ear canal exudate with Gram-positive cocci (Gram Twort stain). (I-L) Other P21 EdaTa mice have thin crusts overlying the tympanic membrane containing neutrophils and Gram-positive cocci (K,L). There is no marked inflammatory reaction in the skin of the ear canal. (M-P) P79 (M,N) and P90 EdaTa (O,P) mice. (M,O) The Zymbal’s gland is present and there is no otitis externa. (M,N) Minimal intrabullar exudate (otitis media; horizontal arrowheads). (O,P) The ear canal contains a plug of bland squamous cells. Severe otitis media with neutrophilic exudate in the bulla cavity and thickened inflamed mucosa can be observed (vertical arrowheads).
Otitis externa and changes in tympanic membrane and ear canal

The tympanic membrane comprises a rostral (anterior) region, the pars tensa and a posterior region, the pars flaccida, which has a thicker lamina propria. The malleolar fold forms the boundary between the pars tensa and pars flaccida, and is marked by the attachment manubrium of the malleus. The tympanic membrane has an outer squamous keratinising epithelium and a poorly characterised inner mucosal non-keratinising epithelium, which is continuous with the mucosa overlying the bulla bone (Mozaffari et al., 2020). Hereafter, we use the nomenclature relating to tympanic membrane anatomy and histology from Mozaffari et al. (2020).

In the healthy ear, the skin stratified squamous epithelium and the tympanic membrane outer epithelium (pars flaccida and pars tensa) were keratin 5 (K5, also known as KRT5)+ (Fig. S3A-C), whereas

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**Fig. 3.** See next page for legend.
the single one-layered inner mucosal epithelium had only scattered K5+ cells (Fig. S3B,C). K5 staining occurred in basal cells, supraabsal cells and external root sheath follicular epithelium (Fig. S4B,C).

K7+, K8+ and K18+ (also known as KRT7, KRT8 and KRT18, respectively) cells were present in epithelium of the inner mucosal layer of the pars flaccida and pars tensa, but absent from the outer epithelial layer; Zymbal’s gland stained with K5 and K18 (Fig. S3D-L). In situ hybridization (ISH) for RNA transcripts encoding the upper respiratory tract innate immunity protein BPIFA1 (Musa et al., 2012) showed that signals were restricted to epithelium of the inner mucosal layer of the pars flaccida and that covering the manubrium of the malleus surface, but not the pars tensa, and the Zymbal’s gland was negative (Fig. S3M-O).

P21 EdaTa ear canals with tympanic membrane crusts had minor hyperkeratosis in ear canal epithelium (Fig. S4E-G) but this was not noticeable in the tympanic membrane outer epithelium (Fig. S4H). K5-staining was present in basal and suprabasal cells but weak in superficial anucleate squames (Fig. S4F,G). Ki67 (also known as MKI67) stained basal epithelial cells in the ear canal and Zymbal’s gland, tympanic membrane and inflammatory cells in the bulla cavity (Fig. S4LJ).

In P21 EdaTa ear canals with otitis externa, the skin epithelium (Fig. S4K-M) and the outer epithelial layer of the tympanic membrane was thickened and hyperkeratotic (Fig. 2E; Fig. S4N) compared with the normal tympanic membrane (Fig. 2F; Fig. S4D), and K5 staining showed that this epithelium has a porous appearance due to transmural infiltration with polymorphonuclear neutrophils, which stain with the proliferation marker Ki67 (Fig. S4L-O). There was no evidence of full-thickness epithelial ulceration or tympanic membrane perforation in any of the P21 mice examined (Fig. 2A-C). Otitis externa in P21 EdaTa mice was also associated with inflammation of ear canal dermis (Fig. 2D). Zymbal’s gland sebocytes and duct epithelium was K5+, and the basal epithelium was Ki67+ in P21 EdaTa mice (Fig. S4P,Q). Zymbal’s gland basal cells and mature sebocytes were K5+ in P22 Mecomobs+ mice (Fig. S4R,S). The mouse tympanic membrane attains its mature size at P18 (Huangfu and Saunders, 1983), and its dorsal plane diameter was not significantly different in P21 EdaTa and P21 FVB mice [median 1.88 mm, 95% c.i. 1.80-2.02 mm (n=14), and median 1.88 mm, 95% c.i. 1.81-1.92 mm (n=26), respectively; P=0.3876, Mann–Whitney test].

Enrofloxacin treatment of EdaTa mice reduces the prevalence of otitis externa and otitis media at P21

Our EdaTa mouse colony has a high prevalence of bacterial rhinitis associated with Staphylococcus aureus (Azar et al., 2016), and we expected this bacterium would naturally colonise the skin and nasal passages of newborn pups. Enrofloxacin is a broad-spectrum antibiotic used to treat skin and soft-tissue infections in dogs. We explored the effect of enrofloxacin treatment administered via drinking water (Macy et al., 2000) on ear disease in P21 EdaTa mice.

We found S. aureus in pure culture in 11 of 12 nasal washes of untreated P21 EdaTa mice; one mouse with otitis externa had pure culture of Aerococcus viridans. S. aureus was found in 14 of 14 nasal washes in enrofloxacin-treated P21 EdaTa mice, and the titres were comparable to the untreated group (P=0.657, Mann–Whitney test). However, in the enrofloxacin treated group, there was a higher prevalence of mixed cultures with Enterococcus faecalis as a co-isolate (5 of 14) compared to the untreated group (P=0.0425, Fisher’s exact test). Furthermore within the enrofloxacin-treated group, S. aureus titres were lower when E. faecalis was a co-isolate [median 9.0×104 colony-forming units (CFUs), 95% c.i. 8.0×103-4.8×105, n=5] than when S. aureus was found in pure culture (median 1.0×106 CFUs, 95% c.i. 1.3×105-5.4×106, n=9; P=0.029, Mann–Whitney test) (Fig. S5K).

Neutrophils were the predominant leucocyte in nasal washes followed by lymphocytes and macrophages (Fig. S5A-C,F,G,J). The number of neutrophils was comparable in enrofloxacin-treated and untreated mice but lymphocytes and macrophages were significantly lower in the treated group (Fig. S5J); squamous cells, basal cells and ciliated cells (Fig. S5E,H,J) were comparable in both groups. Cocci were present in mucus and attached to squamous cells (Fig. S5D,E), and fibres and plant foreign body material were abundant in most samples.

The prevalence of otitis externa and otitis media were significantly reduced in the enrofloxacin-treated group (Table S1, Fig. S2B). There was no evidence enrofloxacin treatment affected
Zymbal’s gland size or ear canal pilosebaceous unit density in P21 EdaTa mice (Fig. 3J,K,R).

**Zymbal’s gland hypoplasia and ear canal hypotrichosis in Edar-deficient mice**

The size of the Zymbal’s gland in P82 Edar-deficient mice (EdarOFEIB/OFEIB) was not significantly different than in P82 EdaTa mice, and both were significantly smaller than in P82 wild-type mice (Fig. S6). The skin of the osseous ear canal was sparsely haired in EdarOFEIB/OFEIB and EdaTa mice compared with wild-type mice (Fig. S6).

**Zymbal’s gland hypoplasia in Edaraddswh/swh rats**

Zymbal’s glands in Edaraddswh/swh rats were significantly smaller than those in Edaraddswh/+ rats at P21, P42 and P83-P85 (Fig. 4A,D,F,H,J-L,N). There was no overt ear canal hair hypotrichosis in Edaraddswh/swh rats compared to Edaraddswh/+ rats (Fig. 4B,C,E,G,I). Pilosebaceous unit sebocytes were reduced at P21 and P42 but were comparable to littermate Edaraddswh/+ rats at P83-P85 (Fig. 4M). The tympanic membrane diameters were comparable in P21 Edaraddswh/swh (median 2.90 mm, 95% c.i. 2.53-3.19 mm, n=3) and littermate Edaraddswh/+ rats (2.93 mm, 95% c.i. 2.57-3.14 mm, n=5; P=0.7857, Mann-Whitney test). Otitis externa was not observed in Edaraddswh/swh (P21, n=6 ears; P30-P85, n=26 ears), Edaraddswh/+ (P21, n=8 ears; P30-P85, n=20 ears) or Edaradd+/+ rats (P83-P85, n=4 ears).

**Edar expression in Zymbal’s glands**

ISH of adult wild-type mouse skin showed Edar expression in basal cells and, to a lesser extent, sebocytes of follicle-associated sebaceous glands (Kowalczyk-Quintas et al., 2015). Edaradd is expressed in the follicle-associated sebaceous glands of the P13 Edaraddswh/+ rat and in the Zymbal’s gland of the P10 Edaradd+/+ rat (del-Pozo et al., 2019a). As the postnatal growth of the mouse Zymbal’s gland appears to be dependent on EDAR signalling, we investigated which cell types expressed Edar. In P10 (Fig. 5A-F) and P30 (Fig. 5G-I) FVB mice, we found the greatest density of punctate Edar ISH signals in basal cells at the periphery of the Zymbal’s gland lobules (Fig. 5F,I), whereas the positive control probe, detecting ubiquitin C (Ubc), showed signals were abundant in all gland cell types (Fig. 5D). No signal was detected by the negative control probe, which recognises the bacterial DapB gene (Fig. 5E).

Edar ISH signals were also detected in the P21 EdaTa Zymbal’s gland; however, the gland is relatively small and has less pronounced lobulation than the wild-type FVB gland (Fig. S7). Both the Edar signal and the positive control probe in P21 EdaTa (Fig. S7) were relatively lower than in P10 FVB (Fig. 5), but the interpretation of relative expression levels in these separate experiments is problematic as the samples were produced in different labs, and would likely have had differences in gland anatomy and its constituent cell populations.

**DISCUSSION**

We report that EdaTa and EdarOFEIB/OFEIB mice, and Edaraddswh/swh rats, have growth retardation of the Zymbal’s gland, resulting in gland hypoplasia, emphasizing the general importance of the EDAR signalling pathway in rodent Zymbal’s gland development. The development of Zymbal’s gland has not been investigated in detail but in EdaTa and NF-kb−/− mice, the Meibomian and preputial glands do not develop, and they are reduced in Traff−/− mice, indicating the involvement of Troy (also known as TNFRSF19) signalling, as well as EDA-EDAR signalling (Dhohaui and Offedal, 2016). Zymbal’s gland development is considered to be independent of hair follicle development (Dhohaui and Offedal, 2016), and in keeping with this, we found small sebaceous gland rudiments in P21 EdaTa located in the normal anatomical site of Zymbal’s gland. These are larger than normal ear canal pilosebaceous unit sebaceous glands, lack obvious hair follicles and develop into histologically normal Zymbal’s glands by P79-P90. We observed only a single instance of a vibrissa-sized hair follicle located in Zymbal’s gland tissue of an Edaraddswh/+ rat among the hundreds of histological sections screened in this study.

The embryonic Zymbal’s gland primordium is present at E15 (Grüneberg, 1971), and we found that prenatal (at E10.5 and E17.5) and prenatal and postnatal treatment (at E10.5, E17.5, P1, P7, and P14) of EdaTa mice with EDAR signalling agonist antibody (del-Pozo et al., 2019a) promotes Zymbal’s gland growth by P21. We found Edar gene expression in the Zymbal’s gland of P10 and P30 FVB mice, as well as in P21 EdaTa mice, suggesting the adult gland has the potential to respond to EDAR signalling. These results are consistent with the known importance of EDAR signalling in the development and function of sebaceous glands, and Meibomian glands in mice, rats, dogs and humans (Kuramoto et al., 2005, 2011; Kaercher et al., 2014; Kowalczyk-Quintas et al., 2015; Waluk et al., 2016; Schneider et al., 2018; Margolis et al., 2019).

The Zymbal’s glands continue to grow in untreated EdaTa mice, and by P79-P90 the glands attain ~50% of normal size, and prenatal agonist anti-EDAR antibody treatment did not result in additional growth in older mice. Sustained treatment of EdaTa and wild-type mice with agonist anti-EDAR antibody heightens sebum production by hair follicle sebaceous glands (Kowalczyk-Quintas et al., 2015), and we infer that continuous treatment would have the same effect on the adult Zymbal’s gland. A single treatment with agonist antibody at P21-P26 produces a long-lived but reversible effect on EdaTa hair follicle sebaceous gland size; the effect was present at 12 weeks but not after 24 weeks (Kowalczyk-Quintas et al., 2015). The apparent lack of effect on P79-90 EdaTa Zymbal’s gland size by *in utero* treatment may represent a similar time-limited response to agonist antibody treatment. Zymbal’s gland growth was also reduced (to ~46% of normal size) in P82 Edar-deficient mice compared to P82 wild-type mice.

In EdaTa and EdarOFEIB/OFEIB mice, but not Edaraddswh/swh rats, there is marked ear canal hypotrichosis. This difference between the density of ear canal hair follicles in the rat and mouse model mirror those elsewhere in the body. Edaraddswh/swh rats have sparse hair coats and hypoplasia of pilosebaceous units in skin of the head, dorsum and ventrum, but unlike EdaTa mice, Edaraddswh/swh rats have a haired tail and lack a bald patch behind the ear (Kuramoto et al., 2005, 2011). P10 EdaTa mice have few ear canal hair follicles, and these have small sebaceous glands (Grüneberg, 1971). Additionally, hair follicles in the adult pinna have small sebaceous glands (Kowalczyk-Quintas et al., 2015). The scarcity of pilosebaceous units in the ear canal may exacerbate the regional deficit of sebum caused by Zymbal’s gland hypoplasia. Although EDAR signalling deficiency in Edaraddswh/swh rats results in marked Zymbal’s gland hypoplasia, ear canal pilosebaceous units were less affected.

Six of 12 P21 EdaTa mice had unilateral otitis externa, but this was absent in P21 FVB mice with full-sized Zymbal’s glands, P21 EdaTa mice with rescued Zymbal’s glands and in P22 Mecom+/− and P21 Fbxo11+/- mice, which have unremarkable Zymbal’s
glands. Additionally, otitis externa was absent in P79-P90 Eda\textsuperscript{Ta} mice (untreated, and those administered agonist anti-EDAR or isotype antibodies).

Gram-positive cocci were found in all otitis externa lesions, and \textit{S. aureus} is a candidate pathogen (this study and Azar et al., 2016). There are potential similarities here with the nasal carriage of
S. aureus being a risk factor for the development of human skin infection (Toshkova et al., 2001). Treatment of mice with enrofloxacin via 0.25 mg/ml drinking water gives a peak plasma level of 140 ng/ml plasma (Marx et al., 2014), and this dosing regimen in EdaTa dams and their pups reduced the prevalence of otitis externa and otitis media at P21. The required minimum inhibitory concentration (MIC90) for S. aureus is 120-250 ng/ml and for Enterococcus spp is 1000-2000 ng/ml (Marx et al., 2014). The concentrations of enrofloxacin and its metabolite ciprofloxacin in skin, inflamed ear canal tissue and middle ear in dogs with end-stage otitis externa and intercurrent otitis media are significantly higher than those in plasma (4-11-fold and 2-6-fold for enrofloxacin and ciprofloxacin, respectively) (Cole et al., 2009). In addition, enrofloxacin and ciprofloxacin are lipophilic (Blokhina et al., 2016), and are likely to concentrate in sebum lipid, thereby providing antimicrobial prophylaxis that compensates for smaller Zymbal’s glands. Although enrofloxacin treatment did not reduce overall nasal bacterial load in P21 EdaTa mice, there were subtle changes, such as an increased prevalence of resistant E. faecalis as a co-isolate with S. aureus and reduced lymphocyte and macrophage populations. The change in leukocyte differentials may represent a delay in progression toward chronic inflammation and an adaptive immune response.

The microbial status of EdaTa mice also plays an important role in eye disease presentation and progression. Conventionally housed (low health status) and specific pathogen-free (SPF) (high health status) EdaTa mice have Meibomian gland deficits that result in corneal defects caused by desiccation and mechanical injury.
However, inflammation of the eyelids (blepharitis) and conjunctiva (conjunctivitis) were only observed in conventionally housed mice (Cui et al., 2005).

The pathogenesis of tympanic membrane crusts in EdaTa mice is unclear. There are some similarities with tympanic membrane inflammatory casts that are described in humans. These consist of hardened fibrin with low inflammatory cell content, and are hypothesised to be derived from serous exudate from acute otitis media with tympanic membrane perforation or otitis externa (Byun et al., 2013). The similarity in composition between EdaTa crusts and otitis externa exudates raises the possibility that crusts represent an early or resolving phase of ear canal infection and inflammation.

Pre-weaning (2-week-old) mice are more susceptible to S. aureus sepsis due to reduced neutrophil chemotaxis and macrophage phagosome maturation (Zhang et al., 2013). The absence of otitis externa in wild-type mice and in EdaTa mice with rescued Zymbal’s glands indicates that myeloid cell immaturity alone does not initiate otitis externa. Nevertheless, functional immaturity could be a contributory factor to otitis externa susceptibility, and conversely, mature myeloid cell function may aid its resolution. There are no identified humoral or cellular immune deficits in HED that might contribute to otitis externa susceptibility. Myeloid and lymphocyte cell classes in blood, spleen, bone marrow and peritoneum are broadly similar in adult EDAR-deficient downless mice (EdaTa) and unaffected heterozygous littermates (Edar+/-), and the phagocytic activity of peritoneal macrophages is comparable in EdaTa, Edar+/- and EdaTa+/- mice (Azar et al., 2016). Furthermore, dogs with XLHED have frequent respiratory tract infections but this is attributable to reduced mucociliary clearance and the absence of bronchial glands rather than any identifiable immune deficiency (Casal et al., 2005). EdaTa mice did not have congenital stenosis of the ear canal at the level of tympanic membrane attachment to the tympanic ring. However, there is minor canal wall thickening during acute inflammation.

We interpret that the pattern of ear canal infection and inflammation in EdaTa mice is primarily the result of changes in Zymbal’s gland size and function, which occurs as follows: (1) the window of susceptibility to bacterial infection and initiation of otitis externa occurs in the interval between the re-opening of the ear canal at P7 and P21, during which Zymbal’s gland growth in the absence of EDA is retarded; (2) endemic skin/nasal commensal bacteria, such as S. aureus, act as opportunistic pathogens; (3) Zymbal’s gland hypoplasia and ear canal hypotrichosis at ≤P21 results in sebum deficiency and thereby reduced innate immune protection (see discussion below); (4) agonist anti-EDAR treatment promotes Zymbal’s gland growth and sebum production by P21 to reduce susceptibility; (5) EDAR signalling-independent growth of the Zymbal’s gland between P21 and P79 restores sebum production; and (6) the absence of otitis externa in older mice indicates that acute disease is self-limiting and that tissue injury is repaired without identifiable sequelae, such as ear canal stenosis.

The numbers of Edaradd+/-msh/msh rats in this study were too low to fully assess the prevalence of otitis externa but there are reasons to suspect that rats may be less susceptible. The sebum secreted by ear canal pilosebaceous units in Edaradd+/-msh/msh rats may provide sufficient protection and/or infection by opportunistic pathogens is less common. Nasal bacteria may be a significant source in EdaTa mice, and it is noteworthy that P21-P85 Edaradd+/-msh/msh rats do not have rhinitis (del-Pozo et al., 2019a). A microbial challenge experiment might be a useful approach to explore infection of the ear canal in Edaradd+/-msh/msh rats. Otitis externa was not observed in the small number of EdaTOFE1BOFE1B mice examined in this study. Reduced Zymbal’s gland size and ear canal hypotrichosis may well predispose to ear canal infection in EdaTOFE1BOFE1B mice but spontaneous disease will also depend on the occurrence of opportunistic bacterial pathogens in colony mice.

The response of the tympanic membrane to otitis externa is not well documented but its barrier function and resilience would presumably depend on epithelial integrity and regenerative capacity. We found that the outer keratinising epithelial layer is K5+ but the inner mucosal layer epithelium has only scattered K5+ cells, which may represent a putative stem cell population. K5 is abundant in basal cells of human stratified squamous epithelium and downregulated in suprabasal cell layers (Moll et al., 2008). We observed suprabasal K5 staining in healthy, mildly hyperkeratotic and inflamed hyperkeratotic ear canal epithelium. The stratified epithelium of ear canal is thicker than that of the tympanic membrane, and number differentiating cell layers, as well as increased cell proliferation, appear to contribute to variation in suprabasal cell expression of K5 protein. The primary keratin pair K8 and K18, and the secondary keratin K7, are expressed in simple (one-layered) epithelium (Moll et al., 2008), and we found them expressed in the inner mucosal epithelial layer of the tympanic membrane. The pattern of keratin and BPIFA1 expression in the inner epithelial layer is similar to that of the non-tympanic middle ear epithelium (del-Pozo et al., 2019b); however, BPIFA1 is restricted to the epithelium of the pars flaccida and the malleus. Bacterial infection of the ear canal stimulates recruitment of neutrophils and these are observed transiting through the hyperkeratotic outer epithelium of the tympanic membrane and the ear canal.

In P21 EdaTa ears there is a statistical association between otitis media and otitis externa but tympanic membrane perforation was not observed. Ear canal infection is characterised by severe inflammation and intralesional cocci, so it is unlikely to be secondary to mild middle ear inflammation where cocci are absent. Even when otitis media is severe in >P79 EdaTa mice there is no evidence of otitis externa. We conclude that the two conditions develop independently in P21 EdaTa mice (otitis media via an auditory tube gating defect, del-Pozo et al., 2019b), and that otitis media does not predispose to otitis externa in Mecom+/- and Fbxo11+/- mice.

In contrast to EdaTa mice, ear canal pathology in human HED is characterised by cerumen impaction and stenosis (Siegel and Potsic, 1990; Daniel et al., 2002; Shin and Hartnick, 2004; Mehta et al., 2007; Yildirim et al., 2012; Callea et al., 2013). It is possible that cerumen impaction in human HED is a manifestation of impaired sebaceous and ceruminous gland secretion, and failure of keratinocyte expulsion from the ear canal.

Human cerumen comprises desquamated keratinocytes and ~50% of the dry weight is a lipid fraction, comprising long-chain fatty acids, triacylglycerols, cholesterol and cholesterol esters, wax esters, ceramides and squalene. The squalene and wax esters appear to be sebaceous gland lipids rather than squamous cell products (Bortz et al., 1990; Daniel et al., 2002; Shin and Hartnick, 2004). Although impacted cerumen is a rich medium for microbial growth (Guest et al., 2004), the action of epithelial ceramidases on ceramides and modification of sebum triglycerides by bacteria produces free fatty acids, triacylglycerols, cholesterol and cholesterol esters, wax esters, ceramides and squalene. The squalene and wax esters appear to be sebaceous gland lipids rather than squamous cell products (Bortz et al., 1990; Guest et al., 2004). Although impacted cerumen is a rich medium for microbial growth (Guest et al., 2004), the modification of sebum triglycerides by bacteria produces free fatty acids, and the action of epithelial ceramidases on ceramides produces sphingosines; both free fatty acids and sphingosines have antimicrobial activity (Wertz, 2018). In a study by Lum et al. (2009) >87% of human cerumen samples had bactericidal activity against S. aureus and Pseudomonas aeruginosa, and fungicidal activity against Candida albicans but a minority of samples showed activity. Proteomic analysis of cerumen has identified...
antimicrobial constituents such as zinc-alpha-2-glycoprotein, cathepsin D, apo-lipoprotein D, serpins, calpain, mucins and lysozyme C (Feig et al., 2013). The tympanic membrane and pilosebaceous units in the human ear canal produce beta defensins (hBD1 and hBD2) (Boe et al., 1999; Yoon et al., 2008), whereas ceruminous (apocrine) glands produce hBD1, hBD2, cathelicidin, lysozyme, lactoferrin, MUC1 and the secretory component of IgA (Stoeckelhuber et al., 2006).

Current animal models of AOE depend on disrupting the ear canal epithelial barrier. A Sprague Dawley rat model of otitis externa has had mechanical abrasion of the cartilaginous ear canal induced, and the resultant inflammation, dermal thickening and hyperkeratosis can be moderated by topical application of steroid or polymyxin B (Engmård and Hellström, 1997, 2001). This model was also infected with *P. aeruginosa* or *C. albicans* to demonstrate the efficacy of topical steroids without antibiotics (Engmård et al., 2005) and thymoquinone (Demirel et al., 2018). A guinea pig model of infectious otitis externa has been established by perturbing the ear canal environment and its commensal bacteria, which facilitates experimental *P. aeruginosa* infection (Wright and Dineen, 1972). Unlike other AOE models, gerbils were successfully infected with *Klebsiella pneumoniae* without ear canal pretreatment (Zhai et al., 2014). Topical application of tetracaine/plphorbol acetate induces acute skin inflammation, and has been used in the mouse ear canal to evaluate the efficacy of ciprofloxacin and hydrocortisone ear drops in reducing inflammation (Wright et al., 2000).

The first step toward developing an *Eda* model of otitis externa will be to establish an inoculation protocol and to gain a better understanding of the time course and its spontaneous resolution. If ear canal infection rates can be raised with microbial challenge, the numbers of mice used will be minimised. It may be necessary to establish an *Eda* mouse colony with low nasal commensal bacteria through the use of prophylactic antibiotics to avoid interference with experimental microbial inoculations. The advantages of an *Eda* mouse challenge model are that inoculation requires no pretreatment of the ear canal and is therefore minimally invasive, husbandry costs are minimised by having a short bioassay time and otitis externa appears to be a self-limiting disease. However, there are noteworthy species differences, in particular the apparent lack of apocrine ceruminous glands in rodents, which may affect cerumen composition and consistency. The issue of whether otitis externa causes otalgia needs to be addressed, but there was no obvious self-inflicted pinna injury, and ear canal inflammation was localised.

In conclusion, we report that mouse and rat models of HED have Zymbal’s gland hypoplasia, and in *Eda* mice there is also ear canal hypotrichosis. Sebum deficiency coupled with endemic nasal carriage of *S. aureus* in our SPF colony predisposes to opportunistic infection of the ear canal and AOE in *P21 Eda* mice. To our knowledge, this is the first report of naturally occurring otitis externa in inbred or genetic strains of laboratory mice. The *Eda* mouse is a model of AOE that will be useful for investigating the cellular and molecular pathology of ear canal infection.

**MATERIALS AND METHODS**

**Animals and in vivo procedures**

The animal experiments were reviewed and approved by the Roslin Institute Animal Welfare and Ethical Review Body, and were performed under the authority of an appropriate UK Home Office Licence. Tabby mice (*Eda*/*Ta*) females and *Eda*/*Y* hemizygous males; collectively termed *Eda*/*Ga* were maintained as a homozygous line. FVB mice are the background inbred genetic line for the *Eda* strain and FVB/N_Crt (Charles River) mice were bred to provide control tissues. The sparse and wavy hair (swh) *Edaradd* rat strain (Kuramoto et al., 2005, 2011) WTC-swh/Kyo [National BioResource Project (NBRP) Rat No. 0287] was supplied by the NBRP - Rat, Kyoto University, Japan. The *Edaradd* rat colony was maintained by mating heterozygous *Edaradd*/*w*/*s* rats, or mating male *Edaradd*/*w*/*s* with *Edaradd*/*w*/*s* females. The heterozygote *Edaradd*/*w*/*s* rat has a wild-type appearance but homozygous *Edaradd*/*w*/*s* animals have typical HED dental and cutaneous phenotypes.

Heterozygous *Fbox11*/*Tm* mice (Mouse Genome Informatics, MGI, 1862017; European Mouse Mutant Archive (EMMA), EM:00375) and their *Fbox11*/*T* wild-type littermates were generated by intercrossing F1 *Fbox11*/*Tm* C57BL/6 J C3H/HeJ males with C57BL/6 J (Charles River) females. Heterozygous *Mecom*/*Bc*/*m* mice (MGI, 2158381; EMMA, EM:00091) and their wild-type littermate controls, *Mecom*/*Bc*/*m* are congenic on a C3H/HeJ genetic background. These strains were obtained from the Mary Lyon Centre (Medical Research Council, Harwell). Mouse and rat husbandry, genotyping, health surveillance and SPF status are reported elsewhere (Azar et al., 2016; del-Pozo et al., 2019a,b). Male and female mice and rats were used in all analyses.

White-bellied agouti B6CBAa *A<sup>−/−/A-EdaTa/</sup>* Tabby mice (000314; Jackson Laboratory) were bred as *Eda*/*Ga*/*Eda*/*Ga* and *Eda*/*Y* mutants or as *+/−* and *−/−* wild-type controls. EDAR-deficient OVE1B mice were bred as *dlo*/*dlo*/*gNaV1.8* (described previously by Headon and Overbeek, 1999). Four to five animals per cage were housed in an SPF facility at 21°C and 50±10% humidity, with a 14 h-10 h light-night cycle. Mice were provided *ad libitum* with water at pH 2.8 and Global Rodent XP18 food (Kliba Naflag). Cages were enriched with tunnel kraft, vaginal cellulose home, sizzle pad 8G and aspen and beech brick (Serlab). These strains were bred at the University of Lausanne. All mice were handled according to the Swiss Federal Veterinary Office guidelines, under the authorization of the Office Vétérinaire Cantonal du Canton de Vaud (authorization 1370.8 to P.S.).

*Eda* mice were administered agonist anti-EDAR antibody (mAbEDAR1) (Kowalczyk et al., 2011; Schuepbach-Mallepell et al., 2021) at 2 mg/kg either prenatally (E10.5 and E17.5) (n=2) or prenatally and postnatally (E10.5, E17.5, P1, P7 and P14; n=6), and phenotyped at P21. In addition, *Eda*/*Ga* mice treated prenatally (E10.5 and E17.5) with anti-EDAR antibody (n=9) or with isotype control antibody (Aprily2) (n=10) were phenotyped at P85 or P82-P84, respectively (see details of administration routes in del-Pozo et al., 2019a).

Two *Eda*/*Ga* dams were administered enrofloxacin via drinking water (25 mg/ml injectable form of Baytrill, Bayer, Shawnee Mission, KS, USA) throughout pregnancy and lactation (Macy et al., 2000; Towne et al., 2014), and based on water intake, this is equivalent to a dosage of 40 mg/kg. Enrofloxacin, like other drugs, is secreted into milk in humans, mice and bovines via the ABCG2 efflux transporter (Jonker et al., 2005), and enrofloxacin and its metabolite ciprofloxacin are secreted in bovine milk (Idowu et al., 2010). Taken together, mouse pups are likely to acquire antibiotic through milk feeding, as well as by drinking water for themselves once they begin to eat solid food from P12 onwards.

Two litters born to enrofloxacin-dosed *Eda*/*Ga* dams (n=5 and n=9 pups) and two litters born to dams provided with normal drinking water (n=5 and n=7 pups) were euthanised at P21. Mice were sampled mortem by nasal wash microbiology and cytology as described previously (Azar et al., 2016), and the heads were prepared for histology.

**Histology and morphometric analysis**

Animals were euthanised with a rising concentration of CO₂ then decapitated, and the skin, outer pinnae and brain removed. This dissection leaves the ear canal and Zymbal’s gland intact. The fixation of tissues in neutral buffered formalin (NBF) and preparation of the decalcified and wax-embedded heads used in this study, along with details of sectioning in the dorsal plane, staining procedures for Haematoxylin and Eosin (H&E), immunohistochemistry for Ki67, and ISH for cytokeratins K5, K7, K8 and BPIFA1, have been reported previously (del-Pozo et al., 2019a,b).

K18 immunohistochemistry was performed with rabbit anti-cytokeratin 18 antibody (monoclonal EPR17347, Abcam, ab181597) diluted 1 in 1200 and applied for 30 min at room temperature. Antigen retrieval was carried out using Tris-EDTA buffer (pH 9.0) at 110°C for 5 min, and primary
antibody binding was detected using Rabbit Envision (Dako) for 40 mins and DAB chromogen. Additional special stains included Masson’s trichome, Gram Twort, PAS and Grocott’s methenamine silver stain. Histology and immunohistochemistry were performed at the Easter Bush Pathology laboratories, which are UK National External Quality Assessment Service accredited.

Bright-field images were acquired using an Olympus BX41 microscope equipped with a DP72 camera and Cell D software. Slide scans were made using a Hamamatsu NanoZoomer. Morphometrical analysis for object length and area was performed using NanoZoomer software and QuPath software (Bankhead et al., 2017), respectively.

The Zymbal’s gland is located on the anterior (rostral) ear canal between the ectotympanic ring bone and the annular cartilage. We scored the presence or absence of the Zymbal’s gland and measured its total area (sebaceous glandular lobules, connective tissue, ducts, and their lumens). P21 EdaTa mice have small sebaceous gland lobules in the normal location for Zymbal’s gland, and we interpret these to be Zymbal’s gland rudiments rather than of pilosebaceous origin because they were not associated with hair follicles. Any gland tissue microscopically associated with a hair follicle was excluded from the assessment. We also measured the size of sebaceous glands in P21 FVB ear canal pilosebaceous units to compare with the size of sebaceous gland lobules in P21 EdaTa mice.

We measured Zymbal’s glands at a standardised dorsal plane in a ~0.4-cm zone between the lateral opening of the ear canal where the Zymbal’s gland has its largest profile, upward to the level of the cochlea round window. In FVB samples (P21 and P81-P84 ears), a single Zymbal’s gland profile measurement per ear was adequately representative. In P21 EdaTa, the Zymbal’s glands are smaller and not always evident in random sections, so we assessed multiple 40-µm step sections (technical replicates) and averaged two to five measurements. For consistency, we also sampled in this way for P79-P90 EdaTa ears.

In addition, we measured Zymbal’s gland size in dissected P82 ear canals of wild-type (n=8) controls, EdaTa (n=5) and Edar-deficient mice (EdarOVETB/EdarOVETB) (n=3) from the University of Lausanne colony. Ear canals were serially sectioned in 100-µm steps, and we averaged two to six area measurements.

We also averaged Zymbal’s gland area in two to five step sections in P21, P42 and P83-P85 Edaraddswh/swh and Edaraddswh/swh rats. In P83-P85 Edaraddswh/swh rats the Zymbal’s gland can have cystic intralobular ducts and these cystic spaces were omitted from the gland area measurements.

Ear canal pilosebaceous units (the hair follicles and associated sebaceous glands) were counted on the rostral surface of ectotympanic bone between the tympanic membrane and the beginning of the annular cartilage in P21 and P79-P90 EdaTa, P21 and P81-84 FVB mice from the Roslin animal colony, and P82 wild-type, P82 EdaTa and P82 EdarOVETB/EdarOVETB mice from the University of Lausanne colony. Each data point is the median of 2-6 serial step sections.

In addition, we measured the area of sebocytes (excluding ducts and connective tissue) in the ear canal pilosebaceous units of P21, P42 and P83-P85 Edaraddswh/swh and Edaraddswh/swh rats using the same landmarks. Skin over the annular cartilage and osseous regions of the ear canal were examined in serial step sections of P21 FVB mice (n=8) for the presence or absence of acinar glands.

The thickness of ear canal soft tissue (epithelium, dermis and peristeum) overlying the rostral surface ectotympanic ring bone was measured in P21 FVB and in P21 EdaTa mice. The area of tissue was divided by the length of the underlying bone to calculate its average thickness.

The ear canal was scored for presence or absence of otitis externa or thin crusts overlying the tympanic membrane. The diagnostic criterion for otitis externa was the presence of a substantial amount of inflammatory exudate filling the ear canal base adjacent to the tympanic membrane. The exudate is composed of neutrophils mixed with eosinophilic amorphous debris, exfoliated squamous epithelial cells and Gram-positive cocci. This is accompanied by hyperkeratosis and inflammatory thickening of the tympanic membrane and ear canal dermis. Thin crusts overlying the tympanic membrane have a similar composition but the tympanic membrane is not substantially thickened, and although the ear canal epithelium can be hyperkeratotic, intraepithelial inflammatory cell infiltration and dermal inflammation are absent.

The tympanic membrane dorsal plane diameter was measured between its fibrocartilaginous insertion points. We averaged two to five 40-µm step sections for each tympanic membrane length measurement in P21 EdaTa (unaffected by otitis externa), P21 FVB mice and in P21 Edaraddswh/swh and P21 Edaraddswh/swh rats. The middle ear bullae were scored for the presence or absence of otitis media, characterised by the presence of inflammatory cells in the bulla cavity and thickening of the mucosa.

We examined the heads of P21 (n=13) and P81-84 (n=10) FVB mice; P21 EdaTa and P79-R90 (n=13) EdaTa mice; P22 Mecom4b/4b mice (n=7); P21 Fbxo11L/M−/− (n=8); and P21 (n=3), P30 (n=5), P42 (n=4), P83-P85 (n=4) Edaraddswh/swh and P21 (n=4), P30 (n=1), P42 (n=4), P83-P85 (n=5) Edaraddswh/swh rats.

Not every ear section was suitable for making all measurements and assessments, and exclusion criteria were incomplete sections through accidental overtrimming of decalcified tissue blocks, the plane of section being outside target level, or processing artefacts that obscure critical features. The number of ears (biological replicates) assessed for each feature are given in the results and figure legends. Additional mice were surveyed for the presence or absence of otitis externa: P84 Mecom4b/4b (n=6), P22 Mecom4/4+ (n=7); P57-P223 Fbxo11L/M−/− (n=22); and Fbxo11L+/+. Edar addswh/swh; rats

Edar in situ hybridization

The skulls of P10, P21 and P30 mice were skinned, and brains were removed and fixed in NBF. The ear canals were dissected within 1-2 h, then returned into fixative for a total time of 24 h then decalcified with 14% EDTA for 8 h (P10) or 12 h (P21 and P30) before processing to wax.

Edar ISH was performed manually using an RNAscope Multiplex Fluorescent V2 assay (ACD). Briefly, paraffin-embedded tissue sections (4 µm) were prepared and processed as per the manufacturer’s instructions. Sections were hybridised with a mouse Edar probe (ACD, 423011-C3), and serial sections were incubated with either a species-specific 3-plex positive control (ACD, 320881) or a 3-plex negative control (ACD, 320871). Following hybridisation and subsequent amplification steps, sections were incubated with Opal 620 dye (Akoya Biosciences, FP1495001KT), counterstained with DAPI and mounted in Prolong Gold. Samples were imaged using a Zeiss LSM 880 confocal microscope.

Statistical analysis and graphical representation

D’Agostino–Pearson normality tests showed the datasets were not normally distributed (or that the group size was too small to test for normality), so we used either Mann–Whitney tests or Kruskal–Wallis tests followed by Dunn’s multiple comparison tests. Graphs represent data as either points with the median, or Tukey’s method box-and-whisker plots; the box represents the 25% percentile, the median and the 75% percentile, the whiskers represent the minimum and the maximum, and the outliers are represented as points.

Contingency tests comparing disease frequencies were performed using Fisher’s exact tests. Ear disease frequency can be expressed by two measures, per animal and per ear. As ear disease was often unilateral, it follows that the frequency of affected animals is higher than the frequency of affected ears. For this reason, disease frequency per animal may achieve statistical significance but the same data may not show a significant difference per ear. The raw data and the contingency test results are presented in Table S1. Graphs and statistics were generated using GraphPad Prism version 8.4.3 (471). Two-tailed tests were used throughout and test values of P<0.05 were considered to be statistically significant.

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new model for hypoplasia of hair follicle and mammary glands on rat chromosome 17. J. Hered. 96, 339-345. doi:10.1093/jhered/esi053


Fig. S1. Morphometric analysis of tympanic membrane crusts and otitis externa exudate, and inflammatory thickening of ear canal soft tissue in P21 EdaTa mice.

(A) Tympanic membrane crusts have similar composition to otitis externa exudate but are <10% the size of exudate accumulations.

(B) P21 EdaTa ear canals can be categorised as normal healthy, those with tympanic membrane crusts and those with otitis externa. The thickness of ear canal soft tissues is comparable in healthy EdaTa ear canals and FVB controls, while EdaTa ear canals with tympanic membrane crusts and otitis externa exudates are thickened.

Data are represented as points and the median as the histogram bar.

Data in graph (A) was analysed with a Mann-Whitney test, and in (B) using Kruskal-Wallis tests followed by Dunn's multiple comparison tests. Two-tailed tests: not significant ns P>0.05; *P<0.05; **P<0.01; ***P<0.001; ****P< 0.0001.
Fig. S2. Otitis externa is absent in P21 *Eda Ta* mice treated with anti-EDAR antibody (mAbEDAR1) or Enrofloxacin.

(A) The Zymbal's gland is rescued by prenatal treatment with mAbEDAR1 and there is no otitis externa.
(B) The Zymbal's gland is small in an Enrofloxacin treated mouse but there is no otitis externa.
(C) An example of an untreated P21 *Eda Ta* mouse with otitis externa, the ear canal is filled with suppurative exudate mixed with squamous cells. The Zymbal's gland is not evident in its normal location between the annular cartilage and ectotympanic ring. ac, annular cartilage; be, bulla cavity; ec, ear canal; er, ectotympanic ring; ex, inflammatory exudate; tm, tympanic membrane; zg, Zymbal's gland. Scale Bars: (A-C) 1 mm.
Fig. S3. Tympanic membrane and Zymbal’s gland cytokeratin and BPIFA1 expression.

Tissue sections (A-L) P22 MecomJbo/+ , (M-O) P21 Fbxo11Jfl/+ are from mouse strains that do not have otitis externa. Note BPIFA1-positive mesenchyme associated epithelium in P21 Fbxo11Jfl/+ with a bulla cavitation defect.

(A-C) Zymbal’s gland, and ear canal stratified squamous epithelium and outer epithelium of the pars flaccida and pars tensa have high populations of KS-positive cells, whereas the epithelium of the inner mucosal layer epithelium has only scattered KS-positive cells (unlabelled arrowhead). (D-L) K7-, K8- and K18-positive cells are present in inner mucosal layer epithelium of the pars flaccida and pars tensa ; K18 stains the Zymbal’s gland. K7 stains the ear canal epithelium. (M-O) In situ hybridization signals for BPIFA 1 are restricted to epithelium of the inner mucosal layer of the pars flaccida and that covering the manubrium of the malleus but not the pars tensa. be, bulla cavity; ec, ear canal; ece, ear canal epithelium; iml, inner mucosal layer epithelium; lp, lamina propria; mae, mesenchyme associated epithelium; mm, manubrium of the malleus; oel, outer epithelial layer; pf, pars flaccida; pt, pars tensa; zg, Zymbal’s gland.

Scale Bars: (A,D,G,J,M,N) 200 µm; (B,C,E,F,H,I,K,L,O) 50 µm.
Fig. S4. Epithelial hyperkeratosis and inflammation in P21 Eda Ta mice with otitis externa.

(A-D) P22 MecomJbol+ ear canal.
(E-J) P21 Eda Ta ear canal with tympanic membrane crust.
(K-O) P21 Eda Ta ear canal with suppurative otitis externa.

Images A-D are of the same ear canal at different magnifications.

(B,C) Ear canal epithelium from the sites indicated by the unlabelled vertical arrowheads in low power image A.

(B) Non-haired skin overlying the annular cartilage and (C) haired skin of the osseous canal have KS-positive basal and suprabasal epithelial cells, but the acellular superficial squames are negative.

(D) The outer epithelial layer of the tympanic membrane is KS-positive.

(E-J) P21 Eda Ta ear canal with tympanic membrane crust. Ear canal epithelium from the sites indicated by the unlabelled vertical arrowheads in low power image E.

(F) Non-haired skin between the annular cartilage and ectotympanic ring and (G) non-haired stratified squamous epithelium of the caudal surface of osseous canal has KS-positive basal and suprabasal cells, but the acellular superficial squames are negative. (H) The outer epithelial layer stains of the tympanic membrane is KS-positive.

(I) Ki67 staining of recently proliferated basal cells in ear canal epithelium and Zymbal’s gland (arrowheads), and in the tympanic membrane outer epithelial layer and scattered inflammatory cells in the bulla cavity lumen.
(K-O) P21 Eda Ta ear canal with suppurative otitis externa, (K-L) are low and high power images of same ear canal. (L,N) Non-haired hyperkeratotic skin between the annular cartilage and ectotympanic ring is KS-positive and has a porous appearance and (M) shows intraepithelial and exudate polymorphonuclear neutrophil leukocytes (arrowheads).

(N) The outer epithelial layer of the hyperkeratotic tympanic membrane pars flaccida is KS-positive, and has a porous appearance due to infiltrating neutrophils. (O) Ki67 staining shows the neutrophils are recently proliferated (opposing arrowheads indicate margins of the tympanic membrane).

(P,Q) The hypoplastic P21 Eda Ta Zymbal's gland is uniformly KS-positive and has Ki67-positive basal cells (Q). (R,S) Normal Zymbal's glands in a P22 MecomJboll+ mouse in which basal and suprabasal sebocytes have cytoplasmic K5 staining.

ac, annular cartilage; be, bulla cavity; er, tympanic membrane crust; de, dermis; ec, ear canal; ece, ear canal epithelium; er, ectotympanic ring; ex, exudate; fe, external follicle root sheath epithelium; nl, neutrophil leukocyte; tm, tympanic membrane; tmpf, tympanic membrane pars flaccida; zg, Zymbal's gland.

Scale Bars: (A,E,K) 1 mm; (R) 500 µm; (P,Q) 250 µm; (D,H,I,J,N,O) 100 µm. (B,C,F,G,L,S) 50 µm, (M) 25 µm.
Fig. S5. Nasal cytology and microbiology of P21 Eda Ta mice treated with Enrofloxacin.

(A-I) White blood cells (WBC) and epithelial cells in nasal wash samples. 
(A) Neutrophils, solitary viable cell (vertical arrowhead) and cluster of degenerate cells (horizontal arrowhead); (B) polymorphonuclear neutrophils engaged in phagocytosis of cocci; (C) neutrophils adherent to a fibre; (D) cocci in mucus; (E) cocci adherent to a squamous cell; (F) macrophage with a reniform nucleus and cytoplasmic vacuoles; (G) lymphocytes (horizontal arrowheads) and (smaller) red blood cells (vertical arrowheads); (H) columnar ciliated epithelial cells; (I) cohesive cluster of basal epithelial cells. Giemsa stained cytospin preparations. Scale bars 20 µm.

(J) WBC numbers and (K) bacterial isolates in nasal washes of control and Enrofloxacin treated mice. Data are represented as points. In panel J, the histogram bar represents the median.

NL, neutrophils; LYMP, lymphocytes; MAC, macrophages. Note zero leukocyte counts are given a nominal value of 1 to graph on a log scale. In panel K, lines link nasal samples in which E. faecalis is co-cultured with S. aureus.

Data were analysed with Mann-Whitney tests.

Two-tailed tests: not significant ns \( P>0.05 \); * \( P<0.05 \); ** \( P<0.001 \).
Fig. S6. Edar deficient mice have Zymbal’s gland hypoplasia and ear canal hypotrichosis.  
(A-C) Examples of Zymbal’s glands from (A) P82 wild-type, (B) P82 Eda Ta and (C) P82 Edar deficient mice (EdarOVE1B/OVE1B) (H&E stain).  
(D-F) Pilosebaceous units (arrowheads) in ear canal skin of (D) P82 wild-type and (F) P82 EdarOVE1B/OVE1B mice; (E) P82 Eda Ta ear canal has no pilosebaceous units at this section level.  
(G) The size of the Zymbal’s gland at P82 is not significantly different in EdarOVE1B/OVE1B and Eda Ta mice and both are significantly smaller than same aged wild-type mice.  
(H) The ear canals of EdarOVE1B/OVE1B and Eda Ta mice have a low density of pilosebaceous units compared with wild-type mice.  
Data in graphs are represented as points and the bar is the median value. Two-tailed Kruskal-Wallis test and Dunn’s multiple comparison test; not significant ns P>0.05; *P<0.05; **P<0.01.  
ac, annular cartilage; er, ectotympanic ring; tm, tympanic membrane; zg, Zymbal’s gland. Scale Bars: (A-C) 500 µm, (D-F) 250 µm.
**Fig. S7. Edar expression in P21 EdaTa Zymbal’s gland.**

(A-C) Different magnifications of the same P21 EdaTa Zymbal’s gland section (H&E stain).

(D-F) Nearby serial sections of Zymbal’s gland are fluorescent ISH preparations with DAPI nuclear counter stain.

(D) Edar signals are punctate spots (arrowheads).

(E) ISH signals are absent with the negative control probe (DapB).

(F) The positive control probe (Ubc) shows intense ISH signals.

The ISH images (D-F) (and those of Figure 5) were all acquired using the same microscope settings. ac, annular cartilage; ec, ear canal; ece, ear canal epithelium; er, ectotympanic ring; zg, Zymbal’s gland.

Scale Bars: (A) 250 µm; (B) 100 µm; (C) 50 µm; (D-F) 20 µm.
Table S1. Reduction in otitis externa and otitis media prevalence in agonist anti-EDAR antibody (mAbEDAR1) treated and in Enrofloxacin treated P21 EdaTa mice. The prevalence of otitis externa and otitis media in treatment groups was compared to untreated controls using Fisher's exact tests. Note that this is done for assessments on a per mouse basis (where affected mice have either unilateral or bilateral disease) or on a per ear basis. Otitis externa and otitis media are significantly associated in individual ears of untreated P21 EdaTa mice. The association was analysed with a Fisher’s exact test. Statistically significant results $P<0.05$ are highlighted in bold.