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# Acquired enamel pellicle protects gastroesophageal reflux disease patients against erosive tooth wear

**Abstract:** The objective of this study was to compare the protein profile of the acquired enamel pellicle (AEP) formed in vivo in patients with or without gastroesophageal reflux disease (GERD), and with or without erosive tooth wear (ETW). Twenty-four volunteers were divided into 3 groups: 1) GERD and ETW; 2) GERD without ETW; and 3) control (without GERD). The AEP formed 120 min after prophylaxis was collected from the lingual/palatal surfaces. The samples were subjected to mass spectrometry (nLC-ESI-MS/MS) and label-free quantification by Protein Lynx Global Service software. A total of 213 proteins were identified, or 119, 92 and 106 from each group, respectively. Group 2 showed a high number of phosphorylated and calciumbinding proteins. Twenty-three proteins were found in all the groups, including 14-3-3 protein zeta/delta and 1-phosphatidylinositol. Several intracellular proteins that join saliva after the exfoliation of oral mucosa cells might have the potential to bind hydroxyapatite, or participate in forming supramolecular aggregates that bind to precursor proteins in the AEP. Proteins might play a central role in protecting the dental surface against acid dissolution.

Keywords: Tooth Erosion; Dental Pellicle; Proteomics.

## Introduction

Erosive ooth wear (ETW) is characterized by the cumulative loss of mineralized tooth substances, resulting from exposure to nonbacterial acids of intrinsic and/or extrinsic origin, where erosion is the primary causative factor.<sup>1</sup> In recent years, several studies have sought to investigate the biochemical process involving ETW, prompted by an increase in the average global prevalence from 30% to 50%.<sup>2,3</sup> To this end, it is important to identify what individuals are likely to develop ETW, so that early diagnosis and preventive measures can be established.<sup>2</sup>

An important clinical condition associated with ETW is gastroesophageal reflux disease (GERD), typical manifestations being regurgitation, dysphagia, and vomiting. It affects 15% to 25% of the high-income and 10% of the low-income population.<sup>4</sup> However, the prevalence of ETW in patients with GERD is 5-47%,<sup>3</sup> caused by direct contact of regurgitated gastric contents (pH between 1 and 2) with the



tooth surface.<sup>2,5</sup> The low pH of the gastric content seems to suggest that GERD patients would have a higher expected prevalence of ETW. This means that GERD patients without ETW might have some protective factor. The acquired enamel pellicle (AEP) is considered one of the most important factors guarding against ETW.<sup>6</sup> Previously, our team compared the proteomic profile of AEP of GERD patients with and without ETW, in an effort to find what proteins in the AEP would resist removal by intrinsic acids. Among the acidresistant proteins, hemoglobin was found to have increased more than threefold in the AEP of GERD volunteers without ETW, compared with those presenting ETW.<sup>7</sup>

In our previous study,<sup>7</sup> AEP was collected from the buccal surface of the upper and lower teeth. However, AEP composition varies depending on its location in the dental arches.<sup>8</sup> Notably, the severity of the reflux above the upper esophageal sphincter is correlated with the severity of tooth erosion, especially on the palatal/lingual surfaces,<sup>9</sup> which are directly exposed to gastric acids.

Therefore, the aim of this in vivo study was to compare the proteomic profile of the AEP of GERD patients with and without ETW, in order to investigate what proteins in the palatal/lingual AEP would resist removal by intrinsic acids, for the purpose of their use in future AEP engineering procedures.

## Methodology

#### Ethical aspects and subjects

This study was approved by the Ethics Committees of the Bauru and Ribeirão Preto School of Dentistry, (#CAAE 44.007.415.1.0000.5417 and 44.007.415.1.3001.5419, respectively). Twenty-four volunteers signed a consent form to take part in the study (n = 8 per group). These patients were the same as those who participated in our previous study, in which AEP was collected from the buccal surface of the teeth.<sup>7</sup> They were of both genders, and between 20 and 60 years of age. The criteria for inclusion and exclusion, as well as the clinical examination, have been cited elsewhere.<sup>7</sup> The patients were divided into 3 groups,<sup>10</sup> as follows:

- a. Patients with GERD-related symptoms and ETW (GE; n = 8): The inclusion criteria for ETW were BEWE (basic erosive wear examination) ≥ 9, or grade 3 in the upper anterior sextant (with all incisors affected);
- b. Patients with GERD-related symptoms without ETW (GNE; n = 8): Patients without ETW were included in this group (BEWE = 0);
- c. Control group (C; n = 8): Patients in this group did not have GERD-related symptoms or ETW (BEWE = 0).

#### In vivo experiment

The AEP collection procedures were conducted exactly as described in our previous study.<sup>7</sup> The patients were submitted to prophylaxis, and the AEP was collected from the lingual/palatal surfaces of the upper and lower teeth after 120 min, using a 5 X 10 mm electrode filter paper (Bio-Rad, Hercules, USA) pre-dipped in 3% citric acid.<sup>11</sup> One filter paper was used for each quadrant. The batches of paper were stored at -80 °C until the analysis was performed.

#### **Proteomic analysis**

Proteomic analysis of the AEP was conducted as previously reported.7 The protein was extracted by cutting strips of paper collected from all the patients in the same group into small pieces, and then grouping these pieces to form a pool in a single microtube. The processes for AEP sample preparation and shotgun proteomic analysis were performed as previously described.8 The equipment used was a nanoACQUITY UPLC-Xevo QT-MS system (Waters, Manchester, UK). ProteinLynx Global Server (PLGS) version 3.0 (Waters, Manchester, UK) was used to process and search the continuum LC-MSE data. The proteins were identified using the ion accounting algorithm embedded in the software and the Homo Sapiens database (only reviewed, UniProtKB/Swiss-Prot) downloaded in February 2020 from UniProtKB (http://www.uniprot.org/). The label-free quantitative proteomic analysis was performed by analyzing three MS raw files from each pool using PLGS software. All the identified proteins with a confidence score > 95% were included

in the quantitative analysis. Identical peptides from each triplicate by sample were grouped based on mass accuracy (< 10 ppm), and on a retention time tolerance of <0.25 min, using the clustering software embedded in the PLGS. Search results were filtered for a false discovery rate (FDR) of 1%. The difference in expression between the groups was analyzed by the t-test (p < 0.05), using PLGS software. The comparisons made were GE vs. C, GNE vs. C, and GNE vs. GE.

### Results

The characterization of the patients as to age, gender, BEWE score, and % time esophageal pH < 4 was reported elsewhere (Martini et al., 2019), and the proteins were identified (Table 1).

In all, 213 proteins were identified (Table 2), or 106, 119, and 92 from groups C, GE, and GNE, respectively. Figure 1 shows the number of proteins common to the groups, as well as the number of proteins found in just one of the groups. Twentythree proteins were found in all groups, including 14-3-3 protein zeta/delta, 1-phosphatidylinositol 4\_5-bisphosphate phosphodiesterase beta-4, actin isoforms, alpha-internexin, ankyrin-3, annexin A1, apolipoprotein A-II, Ig lambda isoforms, myeloblastin, and myosin light polypeptide 6. Some proteins typically described as existing in the AEP were also common to all groups, such as protein S100-A8, serotransferrin, and serum albumin. The number of proteins found exclusively in groups C, GE and GNE groups was 51, 41 and 40, respectively (Figure, Table 2). Regarding the proteins found exclusively in one or two of the groups, some factors should be highlighted: a) histone H3 isoforms were found only in group C,

**Table 1.** Characterization of the volunteers according to age,gender and BEWE score.

|     | Gender | Median Age $\pm$ DP | BEWE         |
|-----|--------|---------------------|--------------|
| С   | 5F 3M  | 31.37 ± 8.81        | 0            |
| GE  | 7F 1M  | 32.37 ± 8.27        | 16.1 2± 2.08 |
| GNE | 8F     | 31.25 ± 12.72       | 0            |

while histone H2B isoforms were found only in the reflux groups (GE and GNE); b) group GNE had a high number of phosphorylated and calciumbinding proteins; c) isoforms of the spectrin betachain were found exclusively in GNE.

As for the quantitative analyses (Table 3), a comparison of GE vs. C showed that the number of GE proteins increased significantly by 4, and that of C decreased significantly by 14. The proteins that increased included the zinc finger and BTB domain-containing protein 21, negative elongation factor E, serotransferrin, and protein S100-A8. On the other hand, the proteins that decreased in the GE vs. C group included myeloblastin, RNA-binding protein 25, 1-phosphatidylinositol 4 5-bisphosphate phosphodiesterase beta-4, basic salivary proline-rich protein 1, breast cancer type 1 susceptibility protein, centrosomal protein of 170 kDa, neurofilament medium polypeptide, neutrophil defensin 1, alphainternexin, and actin isoforms. As for GNE vs. C, actin\_cytoplasmic 1 increased, while protein S100-A8, alpha-internexin, and annexin A1 decreased. The most important comparison (GNE vs. GE) showed that two isoforms of actin as well as myeloblastin were higher, while protein PRR14L, annexin A1, and protein S100-A8 decreased.

## Discussion

In the previous study by our team, we compared the proteomic profile of the AEP of GERD patients with ETW vs. without ETW for the first time, in an effort to find what proteins in the AEP would help protect against ETW. However, the AEP was collected from the vestibular surface of the teeth.<sup>7</sup> In cases of intrinsic erosion, the route of the gastric acids impacts the palatal and lingual surfaces more.<sup>10-12</sup> That is why we decided to collected AEP from the palatal and lingual surfaces of the patients in the present study. Notably, the site of AEP collection is an important factor that should be taken into account in proteomic studies of this integument, because deep changes in the proteomic profile of AEPs occur according to its location in the dental arches.8 The profile of the proteins found herein was very different from that observed in our previous

| Accession number | Protein name  | С   | GE  | GNE |
|------------------|---|-----|-----|-----|
| P63104           | 14-3-3 protein zeta/delta <sup>(d, e, m, t, v)</sup>                                      | Yes | Yes | Yes |
| Q15147           | <code>l-phosphatidylinositol 4_5-bisphosphate phosphodiesterase beta-4(b, m, r, v)</code> | Yes | Yes | Yes |
| P68032           | Actin_ alpha cardiac muscle 1 <sup>(b, m, n, q, u, w)</sup>                               | Yes | Yes | Yes |
| P68133           | Actin_ alpha skeletal muscle <sup>(b, d, m, n, q, u, w)</sup>                             | Yes | Yes | Yes |
| P62736           | $Actin\_aortic \ smooth \ muscle^{(b, \ d, \ m, \ n, \ q, \ u)}$                          | Yes | Yes | Yes |
| P60709           | Actin_ cytoplasmic 1 <sup>(b, m, n, q, u, w)</sup>  | Yes | Yes | Yes |
| P63261           | Actin_ cytoplasmic 2 <sup>(a, d, g, j, n, q, u, w)</sup>                                  | Yes | Yes | Yes |
| Q8NC06           | Acyl-CoA-binding domain-containing protein $4^{(b, e, m, t, w)}$                          | -   | -   | Yes |
| Q08AH3           | Acyl-coenzyme A synthetase ACSM2A_ mitochondrial $^{(b,\ m,\ t,\ w)}$                     | -   | Yes | -   |
| Q8N6G6           | ADAMTS-like protein 1 <sup>(b, m, t, w)</sup>   | Yes | Yes | -   |
| Q9UIF7           | Adenine DNA glycosylase <sup>(b, m, t, u)</sup>   | Yes | -   | -   |
| Q8N142           | Adenylosuccinate synthetase isozyme 1 <sup>(b, m, r, w)</sup>                             | -   | -   | Yes |
| Q01518           | Adenylyl cyclase-associated protein $1^{(b, m, t, w)}$                                    | Yes | Yes | -   |
| Q9UHX3           | Adhesion G protein-coupled receptor $E2^{(b, d, m, t, u, w)}$                             | Yes | -   | -   |
| Q16352           | Alpha-internexin <sup>(b, d, m, t, u, w)</sup>  | Yes | Yes | Yes |
| P15144           | Aminopeptidase N <sup>(b, c, m, t, u, w)</sup>  | -   | -   | Yes |
| O00213           | Amyloid beta A4 precursor protein-binding family B member $1^{(b, m, t, w)}$              | Yes | -   | -   |
| Q6UB98           | Ankyrin repeat domain-containing protein 12 <sup>(b, m, r, w)</sup>                       | -   | Yes | Yes |
| Q12955           | Ankyrin-3 <sup>(a, b, m, x, w)</sup>  | Yes | Yes | Yes |
| P04083           | Annexin A1 <sup>(b, m, r, w)</sup>  | Yes | Yes | Yes |
| P03973           | Antileukoproteinase <sup>(b, f, m, t, u, w)</sup>   | -   | Yes | Yes |
| P02652           | Apolipoprotein A-II <sup>(a, m, t, w)</sup>   | Yes | Yes | Yes |
| P02656           | Apolipoprotein C-III <sup>(a, m, t, u, w)</sup>   | -   | -   | Yes |
| Q5T2S8           | Armadillo repeat-containing protein $4^{(b, m, t, u, w)}$                                 | -   | -   | Yes |
| Q7L311           | Armadillo repeat-containing X-linked protein $2^{(b,m,t,u,w)}$                            | Yes | -   | -   |
| P51689           | Arylsulfatase $D^{(b, m, t, v)}$  | Yes | -   | -   |
| Q8IUA7           | ATP-binding cassette sub-family A member 9 <sup>(b, c, m, r, u, w)</sup>                  | -   | Yes | -   |
| Q9H4G0           | Band 4.1-like protein 1 <sup>(b, m, s, w)</sup>   | -   | Yes | -   |
| O43491           | Band 4.1-like protein $2^{(b, m, s, w)}$  | -   | -   | Yes |
| Q9Y6E2           | Basic leucine zipper and W2 domain-containing protein $2^{\scriptscriptstyle (b,m,q,w)}$  | -   | Yes | -   |
| P04280           | Basic salivary proline-rich protein 1 <sup>(b, l, o, u)</sup>                             | Yes | Yes | -   |
| P02812           | Basic salivary proline-rich protein $2^{(b, \ l, \ o, \ u)}$                              | Yes | Yes | -   |
| O95342           | Bile salt export pump <sup>(c, m, r, w)</sup>   | Yes | -   | -   |
| Q96IK1           | Biorientation of chromosomes in cell division protein $1^{(b,m,t,w)}$                     | -   | -   | Yes |
| P38398           | Breast cancer type 1 susceptibility protein $^{(b, m, r, w)}$                             | Yes | Yes | -   |
| Q9ULB4           | Cadherin-9 <sup>(b, m, t, u, v)</sup>   | Yes |     | -   |
| Q5SW79           | Centrosomal protein of 170 kDa <sup>(d, m, t, x)</sup>                                    | Yes | Yes | -   |
| Q96L14           | Cep170-like protein(d, m, r, w)   | -   | -   | Yes |
| Q96G23           | Ceramide synthase 2 <sup>(e, m, t, w)</sup>   | Yes | _   | _   |

**Table 2.** Classification of proteins identified in the acquired enamel pellicle collected from volunteers with gastro-esophageal reflux disease (GERD) and erosive tooth wear (GE), GERD but no erosive tooth wear (GNE), or controls (no GERD, no erosive tooth wear; C).

| Continuation |  |     |     |     |
|--------------|--|-----|-----|-----|
| Q8TBZ0       | Coiled-coil domain-containing protein $110^{(b, m, p, x)}$                               | Yes | -   | -   |
| Q86WR0       | Coiled-coil domain-containing protein 25 <sup>(b, c, e, m, t, w)</sup>                   | -   | Yes | -   |
| Q9UBG3       | Cornulin <sup>(b, e, m, t, w)</sup>  | Yes | Yes | -   |
| Q96NY9       | Crossover junction endonuclease MUS81 $^{(b,c,d,m,t,w)}$                                 | Yes | -   | -   |
| Q13363       | C-terminal-binding protein 1 <sup>(b, m, t, w)</sup>                                     |     | Yes | -   |
| P32320       | Cytidine deaminase <sup>(b, d, m, t, w)</sup>  | Yes | Yes | -   |
| O75462       | Cytokine receptor-like factor 1 <sup>(b, m, t, w)</sup>                                  | -   | -   | Yes |
| Q5M775       | Cytospin-B <sup>(b, m, p, w)</sup>   | -   | Yes | -   |
| Q13443       | Disintegrin and metalloproteinase domain-containing protein $9^{(\text{b, m, t, u, w})}$ | Yes | -   | -   |
| Q9Y485       | DmX-like protein 1 <sup>(b, m, t, w)</sup>   | -   | Yes | -   |
| Q5T890       | DNA excision repair protein ERCC-6-like 2 <sup>(b, e, m, t, u, w)</sup>                  | Yes | -   | -   |
| Q9UBZ9       | DNA repair protein REV1 <sup>(b, m, t, u, w)</sup>                                       | -   | Yes | -   |
| P48382       | DNA-binding protein RFX5 <sup>(b, m, t, w)</sup>   | -   | -   | Yes |
| Q8N7B9       | EF-hand calcium-binding domain-containing protein $3^{\scriptscriptstyle(b,m,t,w)}$      | Yes | -   |     |
| Q12805       | EGF-containing fibulin-like extracellular matrix protein $1^{(b,f,m,t,w)}$               | -   | -   | Yes |
| Q6PCB8       | Embigin <sup>(b, m, t, w)</sup>  | Yes | -   | -   |
| Q969X5       | Endoplasmic reticulum-Golgi intermediate compartment protein $1^{(c, m, t, w)}$          | Yes | -   | -   |
| Q14152       | Eukaryotic translation initiation factor 3 subunit $A^{(b,\mbox{ m, t, w})}$             | -   | Yes | -   |
| O15360       | Fanconi anemia group A protein <sup>(b, m, t, w)</sup>                                   | Yes | -   | -   |
| O60907       | F-box-like/WD repeat-containing protein TBL1X <sup>(b, m, t, w)</sup>                    | -   | Yes | -   |
| Q9BZK7       | F-box-like/WD repeat-containing protein TBL1XR1 $^{(b,\ m,\ t,\ w)}$                     | Yes | -   | -   |
| P23142       | Fibulin-1 <sup>(b, m, t, u, w)</sup>   | Yes | -   | -   |
| P09958       | Furin <sup>(b, e, m, t, u, w)</sup>  | -   | Yes | -   |
| P09104       | Gamma-enolase <sup>(b, m, t, v, w)</sup>   | Yes | Yes | -   |
| Q8WUA4       | General transcription factor 3C polypeptide $2^{(b, m, t, w)}$                           | -   | Yes | -   |
| P04406       | Glyceraldehyde-3-phosphate dehydrogenase <sup>(a, b, c, m, t, w)</sup>                   | Yes | Yes | -   |
| Q68CQ7       | Glycosyltransferase 8 domain-containing protein $1^{(b,m,t,x)}$                          | -   | -   | Yes |
| Q08378       | Golgin subfamily A member 3 <sup>(b, d, m, t, x)</sup>                                   | -   | Yes | -   |
| Q02643       | Growth hormone-releasing hormone receptor $^{[b, m, t, x]}$                              | -   | -   | Yes |
| P61978       | Heterogeneous nuclear ribonucleoprotein $K^{(b, m, t, u, w)}$                            | Yes | -   | -   |
| P16104       | Histone H2AX <sup>(b, m, t, u, w)</sup>  | -   | Yes | Yes |
| P33778       | Histone H2B type 1-B <sup>(b, m, t, w)</sup>   | -   | Yes | Yes |
| P62807       | Histone H2B type 1-C/E/F/G/I <sup>(b, m, t, w)</sup>                                     | -   | Yes | Yes |
| P58876       | Histone H2B type 1-D <sup>(b, m, t, w)</sup>   | -   | Yes | Yes |
| Q93079       | Histone H2B type 1-H <sup>(b, m, t, w)</sup>   | -   | Yes | Yes |
| P06899       | Histone H2B type 1-J <sup>(b, m, t, w)</sup>   | -   | Yes | Yes |
| O60814       | Histone H2B type 1-K <sup>(b, m, t, w)</sup>   | -   | Yes | Yes |
| Q99880       | Histone H2B type 1-L <sup>(b, m, t, w)</sup>   | -   | Yes | Yes |
| Q99879       | Histone H2B type $1-M^{(b, m, t, w)}$  | -   | Yes | Yes |
| Q99877       | Histone H2B type $1 - N^{(b, m, t, w)}$  | -   | Yes | Yes |

| Commodiation |  |     |     |     |
|--------------|--|-----|-----|-----|
| P23527       | Histone H2B type 1-O <sup>(b, m, t, w)</sup>   | -   | Yes | Yes |
| Q16778       | Histone H2B type 2-E <sup>(b, m, t, w)</sup>   | -   | Yes | Yes |
| Q5QNW6       | Histone H2B type 2-F <sup>(b, m, t, w)</sup>   | -   | Yes | Yes |
| Q8N257       | Histone H2B type 3-B <sup>(b, m, t, w)</sup>   | -   | Yes | Yes |
| P68431       | Histone H3.1 <sup>(b, f, m, t, w)</sup>  | Yes | -   | -   |
| Q16695       | Histone H3.1t <sup>(b, f, m, t, w)</sup>   | Yes | -   | -   |
| Q71DI3       | Histone H3.2 <sup>(b, f, m, t, w)</sup>  | Yes | -   | -   |
| P84243       | Histone H3.3 <sup>(b, f, m, t, w)</sup>  | Yes | -   | -   |
| Q6NXT2       | Histone H3.3C <sup>(b, f, m, t, w)</sup>   | Yes | -   | -   |
| P62805       | Histone H4 <sup>(b, m, t, w)</sup>   | -   | Yes | Yes |
| O43719       | HIV Tat-specific factor 1 <sup>(b, m, t, w)</sup>  | -   | -   | Yes |
| Q4G0P3       | Hydrocephalus-inducing protein homolog <sup>(e, m, t, w)</sup>                           | Yes | Yes | -   |
| Q9Y4L1       | Hypoxia up-regulated protein 1 <sup>(b, m, t, u, w)</sup>                                | -   | Yes | -   |
| P01876       | lg alpha-1 chain C region <sup>(b, e, m, t, u, w)</sup>                                  | Yes | Yes | -   |
| P01834       | lg kappa chain C region <sup>(b, m, r, w)</sup>  | Yes | Yes | Yes |
| POCG04       | lg lambda-1 chain C regions <sup>(f, m, t, w)</sup>                                      | Yes | Yes | Yes |
| PODOY2       | lg lambda-2 chain C regions <sup>(b, m, o, w)</sup>                                      | Yes | Yes | Yes |
| PODOY3       | lg lambda-3 chain C regions <sup>(f, m, t, w)</sup>                                      | Yes | Yes | Yes |
| POCF74       | lg lambda-6 chain C region <sup>(b, m, t, w)</sup>                                       | Yes | Yes | Yes |
| P55899       | IgG receptor FcRn large subunit p51 <sup>(b, m, t, w)</sup>                              | -   | -   | Yes |
| POC7H9       | Inactive ubiquitin carboxyl-terminal hydrolase 17-like protein 7 <sup>(b, m, t, u)</sup> | -   | Yes | -   |
| P14735       | Insulin-degrading enzyme <sup>(b, m, t, u, w)</sup>                                      | Yes | -   | -   |
| Q86VS3       | IQ domain-containing protein $H^{(b, m, t, w)}$  | Yes | -   | -   |
| P13645       | Keratin_ type I cytoskeletal 10 <sup>(b, m, t, w)</sup>                                  | Yes | Yes | -   |
| P48668       | Keratin_ type II cytoskeletal 6C <sup>(d, m, o, u)</sup>                                 | Yes | Yes | -   |
| Q7Z4W3       | Keratin-associated protein 19-3 <sup>(b, m, t, u)</sup>                                  | Yes | -   | -   |
| C9JBD0       | KRAB domain-containing protein $1^{(b, m, t, x)}$  | Yes | -   | -   |
| Q38SD2       | Leucine-rich repeat serine/threonine-protein kinase $1^{(b, d, m, t, x)}$                | Yes | -   | Yes |
| Q8N653       | Leucine-zipper-like transcriptional regulator 1 <sup>(b, m, s, u)</sup>                  | Yes | -   |     |
| Q8WWI1       | LIM domain only protein 7 <sup>(b, d, m, t, u)</sup>                                     | -   | -   | Yes |
| Q86W92       | Liprin-beta-1 <sup>(b, d, m, t, u, v)</sup>  | Yes | Yes | -   |
| O95711       | Lymphocyte antigen 86 <sup>(b, m, t, w)</sup>  | Yes | -   | -   |
| P40925       | Malate dehydrogenase_ cytoplasmic <sup>(b, m, t, w)</sup>                                | Yes | -   | -   |
| P08493       | Matrix Gla protein <sup>(b, d, m, t, v)</sup>  | -   | Yes | Yes |
| Q9Y2H9       | Microtubule-associated serine/threonine-protein kinase 1 <sup>(b, d, m, t, u)</sup>      | Yes | Yes | -   |
| P15941       | Mucin-1 <sup>(b, i, k, o, u)</sup>   | Yes | -   | -   |
| P24158       | Myeloblastin <sup>(b, c, m, t, w)</sup>  | Yes | Yes | Yes |
| P60660       | Myosin light polypeptide 6 <sup>(b, m, r, w)</sup>                                       | Yes | Yes | Yes |
| Q86WG5       | Myotubularin-related protein 13 <sup>(b, m, t, u)</sup>                                  | -   | Yes | -   |
| P18615       | Negative elongation factor $E^{(b, m, p, w)}$  | Yes | Yes | -   |

Continue

Continuation

| CopPES2Nutracin-2-bet/s <th< th=""><th>Continuation</th><th></th><th></th><th></th><th></th></th<>  | Continuation |  |     |     |     |
|---|--------------|--|-----|-----|-----|
| PS8401Nurrain-JentrefactoreP07197Nurrainflament medium polypaptide*.cms/dVes  | Q9P2S2       | Neurexin-2 <sup>(b, m, t, w)</sup>   | -   | -   | Yes |
| PC7197Naucelianent medium polyapitalsYes <td>P58401</td> <td>Neurexin-2-beta<sup>(b, m, t, w)</sup></td> <td>-</td> <td>-</td> <td>Yes</td>   | P58401       | Neurexin-2-beta <sup>(b, m, t, w)</sup>  | -   | -   | Yes |
| Q04/21Neurogenic locus notic homolog protein 2 <sup>h, k, k, k, kYes·Q92823Neuronal cell adhesion molecule/MathemaniYesVes·Q16286NUT-3 growth factor receptor %C4 hemaniYesYesYesQ8NGQ1Olfactory receptor %C4 hemaniYesYes·Q92084Perosiredosin-5, mitochondrial/smaxiedYesYes·Q92094Phylanoyl-CoA dioxygenose, perosisonal/k-maxiedYesYes·Q92095Perosiredosin-5, mitochondrial/smaxiedYesYes·Q92096Phylanoyl-CoA dioxygenose, perosisonal/k-maxiedYesYes·Q92097Phylanoyl-CoA dioxygenose, perosisonal/k-maxiedYesYes·Q92098Perosiredosin-5, mitochondrial/smaxiedYes···Q92098Phylanoyl-CoA dioxygenose, perosisonal/k-maxiedYes····Q92098Phylanoyl-CoA dioxygenose, perosisonal/k-maxiedYes···<t< sup=""></t<></sup>  | P07197       | Neurofilament medium polypeptide <sup>(b, c, m, t, w)</sup>                          | Yes | Yes | -   |
| Q92823       Neuronal cell adhesion molecule <sup>Antimation</sup> ·       Yes       ·         PS9665       Nutrophil defensin 1 <sup>b, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1, 1,</sup>   | Q04721       | Neurogenic locus notch homolog protein $2^{(b, d, m, t, w)}$                         | Yes | -   | -   |
| PS9665Nustraphil defanin 1******YesYesYesCl /288NT-3 growth factor receptor factor for the factorNesQ8NGQ1Olfactory receptor 9G4 her toredNesNesCl /1832Phytonoyl-CoA dioxygenase_peroxisonalh first-sidYesCl /1832Phytonoyl-CoA dioxygenase_peroxisonalh first-sidYesCl /1832Phytonoyl-CoA dioxygenase_peroxisonalh first-sidYesCl /1832Phytonoyl-CoA dioxygenase_peroxisonalh first-sidYes  | Q92823       | Neuronal cell adhesion molecule $^{(b, d, e, m, e, w)}$                              | -   | Yes | -   |
| Q16288       NT-3 growth factor receptor 9C4 <sup>h,m,k,m</sup> Yes         Q8HCQ1       Otffactory receptor 9C4 <sup>h,m,k,m</sup> Nrs         P30044       Peroxiredoxin-5_mitochondrial <sup>h,m,k,m</sup> Yes       Yes          Q92508       Piezo-type mechanosensitive ion chonnel component 1 <sup>h,h,m,k,m</sup> Yes       Yes          Q92508       Piezo-type mechanosensitive ion chonnel component 1 <sup>h,h,h,m,k,m</sup> Yes       Yes          Q92508       Piezo-type mechanosensitive ion chonnel component 1 <sup>h,h,h,m,k,m</sup> Yes       Yes          Q13310       Polyadenylate-binding protein 1 <sup>h,h,h,h,m,k,m</sup> Yes       Yes          Q0583       POITE ankyrin domain family member f <sup>h,m,k,m</sup> Yes       Yes          Q65830       POITE ankyrin domain family member f <sup>h,m,k,m</sup> Yes       Yes          QCG38       POITE ankyrin domain family member f <sup>h,m,k,m</sup> Yes       Yes          Q65131       Pre-mRNA-splicing foctor XF4-dependent RNA helicase DFMIS <sup>h,h,h,m</sup> Yes        Yes         Q45143       Pre-mRNA-splicing foctor XF4-dependent RNA helicase DFMIS <sup>h,h,h,h,m</sup> .       Yes <td>P59665</td> <td>Neutrophil defensin 1<sup>(b, i, j, o, u)</sup></td> <td>Yes</td> <td>Yes</td> <td>-</td> | P59665       | Neutrophil defensin 1 <sup>(b, i, j, o, u)</sup>                                     | Yes | Yes | -   |
| QRNGQ1       Offactory receptor 9G4 <sup>k-k-1</sup> (**)       · · ·       · · ·         P30044       Peroxitedoxin-5_ mitochondrial <sup>k-k-k-k+k</sup> · · ·       · · ·         Q92508       Piezo-type mechanosensitive ion channel component 1 <sup>k-k-k-k+k</sup> · · ·       · · · ·         Q92508       Piezo-type mechanosensitive ion channel component 1 <sup>k-k-k-k+k-k-k-k-k-k-k-k-k-k-k-k-k-k-k-k</sup>  | Q16288       | NT-3 growth factor receptor <sup>(b, m, r, w)</sup>                                  | -   | -   | Yes |
| P30044Peroxiredoxin-5_mitochondrialProtectYesYesYesO14822Phytonoyl-CAA dioxygenose_peroxisomalProtectYesG92508Piezo-type mechanosensitive ion channel component like issterYesYesYesP11940Polyadenylate-binding protein like issterYesYesG13310Polyadenylate-binding protein like issterYesYesP8161Polyadenylate-binding protein like issterYesYesP8163Polymeric immonglobulin receptore issterYesYesP01833POTE ankyrin domain family member Piens isYesYesP06G38POTE ankyrin domain family member Piens isYesYesP06G38POTE ankyrin domain family member Piens isYesYesP04G38POTE ankyrin domain family member Piens isYesYesP05G38POTE ankyrin domain family member Piens isYesYesP04G38POTE ankyrin domain family member Piens isYesYesP05G38Probable RNA No-adenosine thronyCarbomoyltransferase_mitochondrialin-fieldYesYesQ43143Pre-mRNA-splicing factor XIP-dependent RNA helicase DHX156******YesQ50128Probable RNA No-adenosine thronyCarbomoyltransferase_mitochondrialin-fieldYesQ41400Probable RNA No-adenosine thronyCarbomoyltransferase_mitochondrialin-fieldYesQ50128Pr  | Q8NGQ1       | Olfactory receptor 9G4 <sup>(b, m, t, u, w)</sup>                                    | -   | -   | Yes |
| O14832       Phytonoyl-CoA dioxygenose_peroxisomal <sup>[k-m,1+4]</sup> Yes       I         Q92508       Piezo-type mechanosensitive ion channel component like.in.k.t.d       Yes       Yes         Q92508       Piezo-type mechanosensitive ion channel component like.in.k.t.d       Yes       Yes         Q911940       Polyadenylate-binding protein like.in.k.t.d       Yes       Yes         Q93101       Polyadenylate-binding protein disk.in.k.t.d       Yes       Yes         Q931310       Polyadenylate-binding protein disk.in.k.t.d       Yes       Yes         Q931310       Polyadenylate-binding protein disk.in.k.t.d       Yes       Yes         Q05381       POTE ankyrin domain family member Pie.fis.d       Yes       Yes         Q65491       ProtemRNA 3-end-processing factor PIPI.htt.d       Yes       Yes         Q641145       Pre-mRNA 3-end-processing factor PIPI.htt.d       Yes       Yes         Q642143       Pre-mRNA 3-end-processing factor PIPI.htt.d       Yes       Yes         Q641145       Pro-mRNA 3-end-processing factor PIPI.htt.d       Yes       Yes         Q641343       Pre-mRNA 3-end-processing factor PIPI.htt.d       Yes       Yes         Q64247       Proble E3 ubiquitin-protein Iggee HERC4.htt.d.f.d.                               | P30044       | Peroxiredoxin-5_ mitochondrial <sup>(b, m, s, u, w)</sup>                            | Yes | Yes | -   |
| Q92508       Piero-type mechanosensitive ion channel component 14.**.**       · <td>O14832</td> <td>Phytanoyl-CoA dioxygenase_ peroxisomal<sup>(b, m, t, w)</sup></td> <td>Yes</td> <td>-</td> <td>-</td>   | O14832       | Phytanoyl-CoA dioxygenase_ peroxisomal <sup>(b, m, t, w)</sup>                       | Yes | -   | -   |
| Q99569       Plakophilin-4 <sup>h,d,m,d,l</sup> Yes       Yes       Yes         P11940       Polyadam/late-binding protein 1 <sup>h,k,d,m,d,l</sup> Nes       -         Q13310       Polyadam/late-binding protein 1 <sup>h,k,d,m,d,l</sup> Yes       -         P98161       Polyagan-1 <sup>h,m,n,d</sup> -       Yes       -         Q65813       POIE onkyrin domain family member Fb <sup>m,n,dl</sup> Yes       -       -         Q65813       POIE onkyrin domain family member Fb <sup>m,n,dl</sup> Yes       Yes       -         Q65013       POIE onkyrin domain family member Fb <sup>m,n,dl</sup> Yes       Yes       -         Q65013       POTE onkyrin domain family member Fb <sup>m,n,dl</sup> Yes       Yes       -         Q640145       Pre-mRNA 3'-end-processing factor FIP I <sup>h,m,k,dl</sup> Yes       -       -         Q65013       Pre-mRNA-splicing factor 38 b <sup>h,m,k,dl</sup> -       Wes       -       -         Q640145       Pre-mRNA-splicing factor 38 b <sup>h,m,k,dl</sup> -       Wes       -       -         Q914480       Probable f1NA Mo-doenosine threonyclarbamolytamsførse_mintochondraffb <sup>h,d,m,dl</sup> Yes       -       -         Q914600       Protein FAMJ 4 <sup>j,h,m,dl</sup>  | Q92508       | Piezo-type mechanosensitive ion channel component $1^{(b,e,m,s,w)}$                  | -   | -   | Yes |
| P11940       Polyadenylate-binding protein 1 <sup>k,k,k,m,k,m</sup> ·       Yes       ·         Q13310       Polyadenylate-binding protein 4 <sup>k,k,k,m,k,m</sup> Yes       ·       ·         P98161       Polyaystin-1 <sup>k,m,k,m</sup> ·       Yes       ·       ·         P01833       Polymeric immunoglobulin receptor <sup>k,m,k,m</sup> ·       Yes       ·       ·         Q6583       POTE ankyrin domain family member E <sup>k,m,k,m</sup> Yes       Yes       ·       ·         Q6503       POTE ankyrin domain family member F <sup>k,m,k,m</sup> Yes       Yes       ·       ·         Q64UN15       Pre-mRNA 3'-end-processing factor FIP1 <sup>k,m,k,m</sup> ·       Yes       ·       ·       ·       ·       ·       Yes       · <td< td=""><td>Q99569</td><td>Plakophilin-4<sup>(b, d, m, t, u)</sup></td><td>Yes</td><td>Yes</td><td>-</td></td<>   | Q99569       | Plakophilin-4 <sup>(b, d, m, t, u)</sup>   | Yes | Yes | -   |
| Q13310       Polyadenylate-binding protein 4 <sup>(k-k, k-k, k-k)</sup> Yes       .         P98161       Polyagetin-1 <sup>(k-k, k-k)</sup> .       Yes       .         Q18333       Polymeric immunoglobulin receptor <sup>(k-k, k-k)</sup> Yes       Yes       .         Q65833       POTE onkyrin domain family member E <sup>[k-m, k-k)</sup> Yes       Yes       .         Q56436       POTE onkyrin domain family member I <sup>[k-m, k-k)</sup> Yes       Yes       .         Q60CG38       POTE onkyrin domain family member I <sup>[k-m, k-k]</sup> Yes       .       .         Q60UN15       Pre-mRNA-splicing factor 388 <sup>[k-m, k-k]</sup> .       Yes       .       .         Q64UN13       Pre-mRNA-splicing factor AIP-dependent RNA helicase DHX15 <sup>[k-m, k-k]</sup> .       Yes       .       .         Q54L28       Probable fRNA N6-adenosine threonylcarbamoyltransferase_mitochondriel <sup>[k-m, k-k]</sup> Yes       .       .       .         Q9H480       Probable fRNA N6-adenosine threonylcarbamoyltransferase_mitochondriel <sup>[k-m, k-k]</sup> Yes       .       .       .         Q9H480       Probable fRNA N6-adenosine fRNA M49A <sup>k, k-k, k-k</sup> Yes       .       .       .         Q9H480       Probable fRNA N6                                  | P11940       | Polyadenylate-binding protein $1^{(a, b, f, m, t, u, w)}$                            | -   | Yes | -   |
| P98161       Polycysin-1 <sup>k,m,k,d</sup> .       Yes       .         P01833       Polymeric immunoglobulin receptor <sup>k,m,k,d</sup> Yes       Yes       .         Q658J3       POTE ankyrin domain family member Fk.m,x,dl       Yes       Yes       .         A5A3E0       POTE ankyrin domain family member Fk.m,x,dl       Yes       Yes       .         Q6G38       POTE ankyrin domain family member Fk.m,x,dl       Yes       Yes       .         Q6UN15       Pre-mRNA.3'end-processing factor FIP1ens.td       .       Yes       .         Q6UN15       Pre-mRNA-splicing factor 388 <sup>km, n, m, m</sup> .       Yes       .         Q643143       Pre-mRNA-splicing factor 388 <sup>km, n, m, m</sup> .       Yes       .         Q64140       Probable RNA Ne-admonsine threonylcarbamostres_mitochondrial <sup>bit, m, t, m</sup> .       .       .         Q64247       Probable RNA Ne-admonsine threonylcarbamostres_mitochondrial <sup>bit, m, t, m       Yes       .       .         Q9H480       Probable RNA Ne-admonsine threonylcarbamostres_mitochondrial<sup>bit, m, t, m</sup>       Yes       .       .         Q9H490Q       Protein FAM161A<sup>bit, m, t, m</sup>       Yes       .       .       .   </sup>  | Q13310       | Polyadenylate-binding protein $4^{(a, b, f, m, t, u, w)}$                            | Yes | -   | -   |
| P01833       Polymeric immunoglobulin receptor <sup>1k, m, k, uff       Yes       Yes       Yes         Q658J3       POTE ankyrin domain family member P<sup>k, m, k, uff</sup>       Yes       Yes       Yes         A5A3E0       POTE ankyrin domain family member P<sup>k, m, k, uff</sup>       Yes       Yes       Yes         P0CG38       POTE ankyrin domain family member P<sup>k, m, k, uff</sup>       Yes       Yes       Yes         Q6UN15       Pre-mRNA-3Pilcing factor 388<sup>k, m, k, uff</sup>       Yes       Yes       Yes         Q6VI18       Pre-mRNA-splicing factor 388<sup>k, m, k, uff</sup>       Yes       Yes       Yes         Q5GI28       Probable E3 ubiquifin-protein ligase HERC4<sup>k, d, m, t, uff       Yes       Yes       Yes         Q9H480       Probable RNA N6-adenosine threonylcarbanoyltransforase_mitochondrial<sup>[k, d, m, t, uff</sup>}       Yes       Yes       Yes         Q9H480       Probable RNA N6-adenosine threonylcarbanoyltransforase_mitochondrial<sup>[k, d, m, t, uff</sup>}       Yes       Yes       Yes         Q9H480       Probable RNA N6-adenosine threonylcarbanostarbitochondrial Q<sup>k, d, d,</sup></sup></sup>   | P98161       | Polycystin-1 <sup>(b, m, s, u)</sup>   | -   | Yes | -   |
| QdS8J3       POTE ankyrin domain family member E <sup>(h,m,s,m)</sup> Yes       Yes       Yes       Yes         A5A3E0       POTE ankyrin domain family member f <sup>(h,m,s,m)</sup> Yes       Yes       -         POCG38       POTE ankyrin domain family member f <sup>(h,m,s,m)</sup> Yes       Yes       -         QdUN15       Pre-mRNA 3'-end-processing factor FIP16 <sup>(h,m,t,m)</sup> -       Yes       -         QdS143       Pre-mRNA-splicing factor AIP-dependent RNA helicase DLX15 <sup>(h,m,t,m)</sup> -       Yes       -         QdSU28       Probable IS ubiquitin-protein ligase HERC4 <sup>(h,d,m,t,m)</sup> -       Yes       -         QdMZM9       Probable IRNA N6-adenosine threonylcarbamoyltransferase_mitachondrial <sup>(h,d,m,t,m)</sup> Yes       -       -         QdMZM9       Protein FAM161A <sup>(h,m,t,m)</sup> Yes       -       -       -         QdMZM9       Protein FAM164A <sup>(h,m,t,m)</sup> Yes       -       -       -         QdMZM9       Protein FAM49A <sup>(h,t,t,m)</sup> Yes       -       -       -         QdMZM9       Protein FAM164A <sup>(h,t,t,m)</sup> Yes       -       -       -         QdSM200       Protein FAM164A <sup>(h,t,t,m,t)</sup> Yes       -  | P01833       | Polymeric immunoglobulin receptor <sup>(b, m, s, w)</sup>                            | Yes | Yes | -   |
| A5A3E0       POTE ankyrin domain family member Pix.m. v. vil       Yes       Yes       Yes       .         POCG38       POTE ankyrin domain family member Pix.m. v. vil       Yes       Yes       .         Q6UN15       Pre-mRNA 3'-end-processing factor FIP1b/m.t. vil       .       Yes       .         Q5VTL8       Pre-mRNA-splicing factor 38B/b.m. v. vil       .       Yes       .         Q43143       Pre-mRNA-splicing factor ATP-dependent RNA helicase DHX15/b.m.t. vil       .       Yes       .         Q5GI28       Probable E3 ubiquitin-protein ligase HERC4/b.d.m.t.vil       .       Yes       .       .         Q9H480       Probable RNA N6-adenosine threonylcarbamoyltransformes_milochondrial/b.d.m.t.vil       Yes       .   | Q658J3       | POTE ankyrin domain family member E <sup>(b, m, o, w)</sup>                          | Yes | Yes | -   |
| POCG38       POTE ankyrin domain family member (hk, m, k, w)       Yes       Yes       .         QdUN15       Pre-mRNA 3'-end-processing factor FIP16x, m, k, w)       .       Yes       .         QdSVTL8       Pre-mRNA-splicing factor ATP-dependent RNA helicase DHX15 <sup>(h, m, t, w)</sup> .       Yes       .         QdSGL28       Probable E3 ubiquitin-protein ligase HERC4 <sup>(h, d, m, t, w)</sup> .       Yes       .         QdMZM9       Probable tRNA N6-adenosine threonylcarbamoyltransferase_mitochondrial <sup>(h, d, m, t, w)</sup> Yes       .       .         QdMZM9       Probable tRNA N6-adenosine threonylcarbamoyltransferase_mitochondrial <sup>(h, d, m, t, w)</sup> Yes       .       .         QdMZM9       Protein FAM161A <sup>(h, m, t, w)</sup> Yes       .       .       .         Qd9H0Q0       Protein FAM49A <sup>(h, c, t, w)</sup> Yes       .       .       .         QSTHK1       Protein PRR14[ <sup>(h, t, t, w)</sup> ]       Yes       .       .       .         Q9Y520       Protein S100-A8 <sup>(h, t, t, w)</sup> Yes       .       .       Yes         Q96ER3       Protein unc-13 homolog Cb <sup>(m, t, w)</sup> Yes       .       .       .         Q9V5H0       Protein add GTPase-activati  | A5A3E0       | POTE ankyrin domain family member F <sup>(b, m, o, w)</sup>                          | Yes | Yes | -   |
| QdUN15Pre-mRNA 3'-end-processing factor FIP1b.ms.t.ii)·Yes·Q5VTL8Pre-mRNA-splicing factor ATP-dependent RNA helicase DHX15 <sup>b.m.t.iii</sup> ·Yes·Q5GL28Probable E3 ubiquitin-protein ligase HERC4 <sup>[b.d.m.t.iii</sup> ]·Yes·YesQ9H480Probable tRNA N6-adenosine threonylcarbamoyltransferase_mitochondrial <sup>[b.d.m.t.iii</sup> ]Yes·Yes·Q6MZM9Probable tRNA N6-adenosine threonylcarbamoyltransferase_mitochondrial <sup>[b.d.m.t.iii</sup> ]Yes··YesQ3B820Protein FAM161A <sup>b.m.t.iii</sup> Yes·····Q9H0Q0Protein FAM49A <sup>b.c.t.iii</sup> Yes······Q9H0Q0Protein FAM49A <sup>b.c.t.iii</sup> ·Yes··· </td <td>POCG38</td> <td>POTE ankyrin domain family member 1<sup>(b, m, o, w)</sup></td> <td>Yes</td> <td>Yes</td> <td>-</td>  | POCG38       | POTE ankyrin domain family member 1 <sup>(b, m, o, w)</sup>                          | Yes | Yes | -   |
| QSVTL8       Pre-mRNA-splicing factor 388 <sup>[b,m,q,m]</sup> .       Yes       .         O43143       Pre-mRNA-splicing factor ATP-dependent RNA helicase DHX15 <sup>b,m,t,m]</sup> .       Yes       .         QSGLZ8       Probable E3 ubiquitin-protein ligase HERC4 <sup>[b,d,m,t,m]</sup> .       Yes       .         Q9H480       Probable tRNA N6-adenosine threonylcarbamoyltransferase_mitochondrial <sup>[b,d,m,t,m]</sup> Yes       .       .         Q6MZM9       Probable TSA N6-adenosine threonylcarbamoyltransferase_mitochondrial <sup>[b,d,m,t,m]</sup> Yes       .       .         Q38820       Protein FAM161A <sup>[b,m,t,m]</sup> Yes       .       .       .         Q9H0Q0       Protein FAM49A <sup>[b,c,t,m]</sup> Yes       .       .       .         Q9H5Q1       Protein PRR14L <sup>[b,l,c,n]</sup> Yes       .       .       .         Q9Y520       Protein SIO0-A8 <sup>[b,c,t,m]</sup> Yes       Yes       .       Yes         Q9Y580       Protein unc-13 homolog C <sup>[b,m,t,m]</sup> .       Yes       .       .         Q9Y5910       Protein factor C subuit 1 <sup>[b,m,t,m]</sup> Yes       .       .       .         Q9UBK7       Rel GTPase-activating protein subunit alpha-2 <sup>[b,m,t,m]</sup>   | Q6UN15       | Pre-mRNA 3'-end-processing factor FIP1 <sup>(b, m, t, w)</sup>                       | -   | Yes | -   |
| O43143Pre-mRNA-splicing factor ATP-dependent RNA helicase DHX15 <sup>b,m,t,vil</sup> ·Yes·QSGLZ8Probable E3 ubiquitin-protein ligase HERC4 <sup>b,d,m,t,vil</sup> ··YesQ9H4B0Probable tRNA N6-adenosine threonylcarbamoyltransferase_mitochondrial <sup>b,d,m,t,vil</sup> Yes··Q6MZM9Probable tRNA N6-adenosine threonylcarbamoyltransferase_mitochondrial <sup>b,d,m,t,vil</sup> Yes··Q6MZM9Probable tRNA N6-adenosine threonylcarbamoyltransferase_mitochondrial <sup>b,d,m,t,vil</sup> Yes··Q3B820Protein FAM161A <sup>b,m,t,vil</sup> Yes···Q9H0Q0Protein FAM161A <sup>b,m,t,vil</sup> Yes···Q9H0Q0Protein FAM49A <sup>b,d,t,t,vil</sup> Yes···Q9H5Q1Protein PRR14L <sup>b,t,v,vil</sup> ·Yes·YesQ9Y520Protein S100-A8 <sup>(b,t,t,v,vil</sup> YesYesYesQ96ER3Protein S100-A8 <sup>(b,t,t,v,vil</sup> ·Yes·Q975H0Protein Cable in gamma-A3 <sup>(b,m,t,vil</sup> ·Yes·Q91BK7Rab ClFDase-activating protein subunit alpha-2 <sup>(b,m,t,vil</sup> Yes··Q92PJ7Reflication factor C subunit 1 <sup>(b,m,t,vil</sup> Yes···Q95100Replication factor C subunit 1 <sup>(b,m,t,vil</sup> ·Yes··Q92PJ7Reflication factor C subunit 1 <sup>(b,m,t,vil</sup> Yes···Q92Q2Rho GTPase-activating protein 18 <sup>(b,m,t,vil</sup> ··Yes·Q92Q3Rho GTPase-activating protein                                     | Q5VTL8       | Pre-mRNA-splicing factor 38B <sup>(b, m, q, w)</sup>                                 | -   | Yes | -   |
| QSGLZ8Probable E3 ubiquitin-protein ligase HERC4 <sup>(6, d, m, t, u)</sup> YesQ9H480Probable tRNA N6-adenosine threonylcarbamoyltransferase_mitochondrial <sup>(6, d, m, t, u)</sup> YesQ6MZM9Probable tRNA N6-adenosine threonylcarbamoyltransferase_mitochondrial <sup>(6, d, m, t, u)</sup> Yes-YesQ3B820Protein FAM161A <sup>(6, m, t, u)</sup> YesQ9H0Q0Protein FAM161A <sup>(6, m, t, u)</sup> YesQ9H0Q0Protein FAM49A <sup>(6, c, t, u)</sup> YesQ5GLXProtein in Gggdd-1 <sup>(6, m, t, u)</sup> -YesQ5THK1Protein PRR14L <sup>(6, t, n, t)</sup> -Yes-YesQ9Y520Protein S100-A8 <sup>(6, b, c, d, t, b, t, u, v)</sup> YesYesYesQ96ER3Protein unc-13 homolog C <sup>(6, m, t, u)</sup> -YesYesQ9Y5H0Protein unc-13 homolog C <sup>(6, m, t, u)</sup> -Yes-Q2PPJ7Rel GTPase-activating protein subunit alpha-2 <sup>(6, m, t, u)</sup> YesQ2PPJ7Rel GTPase-activating protein subunit alpha-1 <sup>(6, m, t, u)</sup> YesP35251Replication factor C subunit 1 <sup>(6, m, t, u)</sup> YesP50120Retinol-binding protein 2 <sup>(6, m, t, u)</sup> -Yes-Q8N392Rho GTPase-activating protein 18 <sup>(6, m, t, u)</sup> -Yes-Q6Q81Rho GTPase-activating protein 18 <sup>(6, m, t, u)</sup> -Yes-  | O43143       | $\label{eq:pre-mrane} Pre-mrank RNA \ helicase \ DHX15^{(b,\ m,\ t,\ w)}$            | -   | Yes | -   |
| Q9H480Probable tRNA N6-adenosine threonylcarbamoyltransferase_mitochondrial[b:d=m,t=w]Yes.Q6MZM9Probable tRNA N6-adenosine threonylcarbamoyltransferase_mitochondrial[b:d=m,t=w]YesYesYesQ3B820Protein FAM161A[b:m,t=w]YesQ9H0Q0Protein FAM161A[b:m,t=w]YesQ9H0Q0Protein FAM49A[b:c_t=w]YesP78504Protein PRR14[b:l_0,t_0,s].Yes.Q5THK1Protein PRR14[b:l_0,t_0,s].YesYesQ9Y520Protein S100-A8[a:b,c_d,t_i,t_u,v]YesYesYesP05109Protein S100-A8[a:b,c_d,t_i,t_u,v]YesYesYesQ9KER3Protein SAL1[b:m,t_u]YesYesQ9Y5H0Protein SAL1[b:m,t_u]Q2PPJ7Ral GTPase-activating protein suburi 1[b:m,t_w]YesP35251Replication factor C suburi 1[b:m,t_w]YesP50120Rtho GTPase-activating protein 18(c=m,t_w)YesQ9GQB1Rtho GTPase-activating protein 25(m,t_w)YesQ2PQGQB1Rtho GTPase-activating protein 25(m,t_w)Q2PQBARtho GTPase-activating protein 26(m,t_w)YesQ2PQBARtho GTPase-activating protein 7(b,m,t_w)  | Q5GLZ8       | Probable E3 ubiquitin-protein ligase HERC4 <sup>(b, d, m, t, u)</sup>                | -   | -   | Yes |
| Q6MZM9Proline-rich protein 27(b, 1, o, x)YesYesYesQ3B820Protein FAM161A(b, m, r, w)YesQ9H0Q0Protein FAM49A(b, c, r, w)YesP78504Protein igggdd-1(w, m, r, w)-Yes-Q5THK1Protein PRR14L(b, 1, o, x)-YesYesQ9Y520Protein PRRC2C(b, 1, o, x)YesYesYesQ9Y520Protein S100-A8(h, b, c, d, 1, 1, r, w)YesYesYesQ96ER3Protein SAL1(b, m, r, w)YesYesYesQ975H0Protein unc-13 homolog C(b, m, r, w)-Yes-Q9UBK7Rab-like protein 2A(c, m, t, w)YesQ2PJJ7Ral GTPase-activating protein subunit alpha-2(b, m, t, w)YesP50120Replication factor C subunit 1(b, m, t, w)YesQ8N392Rho GTPase-activating protein 18(c, m, t, w)-Yes-Q9CQB1Rho GTPase-activating protein 18(c, m, t, w)-Yes-   | Q9H4B0       | Probable tRNA N6-adenosine threonylcarbamoyltransferase_ mitochondrial^{(b,d,m,t,w)} | Yes | -   | -   |
| Q3B820Protein FAM161A(b:m,r.w)Yes-Q9H0Q0Protein FAM49A(b:c.f.w)Yes-P78504Protein jagged-1(w,m,r.w).Yes-Q5THK1Protein PRR14L(b,1,a,x).YesYesQ9Y520Protein PRR12C(b,1,a,x)YesYesYesQ90520Protein S100-A8(a, b,c, d, f, j, c, u, v)YesYesYesQ905109Protein S100-A8(a, b,c, d, f, j, c, u, v)YesYesYesQ96ER3Protein SAL1(b,m,r, u).YesYesQ97540Protein unc-13 homolog C(b,m,r, w).YesYesQ97540Protein unc-13 homolog C(b,m,r, w).Yes.Q918K7Rab-like protein 2A(c,m,t, u)YesQ2PPJ7Ral GTPase-activating protein 24(c,m,t, w)YesP50120Retinol-binding protein 2(c,m,t, w)YesQ8N392Rho GTPase-activating protein 18(c,m,t, w).Yes.Q96Q81Rho GTPase-activating protein 7(b,m,t,w).Yes.  | Q6MZM9       | Proline-rich protein 27 <sup>(b, l, o, x)</sup>                                      |     | Yes | Yes |
| Q9H0Q0Protein FAM49A(b, c, r, w)Yes-P78504Protein jagged -1(b, m, r, w)-Yes-Q5THK1Protein PRR14L(b, l, o, x)-YesYesQ9Y520Protein PRRC2C(b, l, o, x)Yes-YesQ9Y520Protein S100-A8(b, b, c, d, f, i, r, v, v)YesYesYesQ96ER3Protein SAL1(b, m, r, w)YesQ975H0Protein unc-13 homolog C(b, m, r, w)-YesYesQ9Y5H0Protocadherin gamma-A3(b, m, r, w)-Yes-Q2PJ7Rab-like protein 2A(c, m, t, w)YesQ2PJ71Red GTPase-activating protein subunit alpha-2(b, m, t, w)YesP50120Retinol-binding protein 2(c, m, t, w)YesQ8N392Rho GTPase-activating protein 18(c, m, t, w)-Yes-Q9CQB1Rho GTPase-activating protein 17(b, m, t, w)-Yes-   | Q3B820       | Protein FAM161A <sup>(b, m, r, w)</sup>  | Yes | -   | -   |
| P78504Protein jagged-1(e, m, r, w).Yes.Q5THK1Protein PRR14L(b, L, o, x).YesYesQ9Y520Protein PRRC2C(b, L, o, x)YesYesYesP05109Protein S100-A8(e, b, c, d, f, i, c, u, v)YesYesYesQ96ER3Protein SAAL1(b, m, r, u)YesQ8NB66Protein unc-13 homolog C(b, m, r, w).YesYesQ9Y5H0Protocadherin gamma-A3(b, m, r, w).Yes.Q9UBK7Rab - like protein 2A(c, m, t, u)YesQ2PPJ7Ral GTPase-activating protein subunit alpha-2(b, m, t, w)YesP50120Retinol-binding protein 2 <sup>(c, m, t, w)</sup> YesQ8N392Rho GTPase-activating protein 18 <sup>(c, m, t, u)</sup> .YesQ9CQB1Rho GTPase-activating protein 7 <sup>(b, m, t, u)</sup> .Yes  | Q9H0Q0       | Protein FAM49A <sup>(b, c, r, w)</sup>   | Yes | -   | -   |
| Q5THK1Protein PRR14Lb, b, a, xi.YesYesQ9Y520Protein PRRC2C(b, b, a, xi)Yes.YesP05109Protein S100-A8(a, b, c, d, f, b, r, v, v)YesYesYesQ96ER3Protein SAAL1(b, m, r, v)YesQ8NB66Protein unc-13 homolog C(b, m, r, w)YesYesQ9Y5H0Protocadherin gamma-A3(b, m, r, v)Q9UBK7Rab-like protein 2A(c, m, t, v)YesQ2PPJ7Ral GTPase-activating protein subunit alpha-2(b, m, t, w)YesP50120Retinol-binding protein 2(c, m, t, w)YesQ8N392Rho GTPase-activating protein 18(c, m, t, w)YesQ9CQBIRho GTPase-activating protein 7(b, m, t, w)   | P78504       | Protein jagged-1 <sup>(e, m, r, w)</sup>   | -   | Yes | -   |
| Q9Y520Protein PRRC2C(b, t, x, x)Yes-YesP05109Protein S100-A8(b, b, c, d, f, j, r, u, v)YesYesYesQ96ER3Protein SAAL1(b, m, r, u)YesQ8NB66Protein unc-13 homolog C(b, m, r, w)-YesYesQ975H0Protocadherin gamma-A3(b, m, r, v)-Yes-Q9UBK7Rab-like protein 2A(c, m, r, u)YesQ2PPJ7Ral GTPase-activating protein subunit alpha-2(b, m, r, w)YesP35251Replication factor C subunit 1(b, m, r, w)YesP50120Retinol-binding protein 18(c, m, r, w)-Yes-Q8N392Rho GTPase-activating protein 18(c, m, r, w)-Yes-Q9CBL1Rho GTPase-activating protein 18(c, m, r, w)-Yes-  | Q5THK1       | Protein PRR14L <sup>(b, l, o, x)</sup>   | -   | Yes | Yes |
| P05109Protein S100-A8(o, b, c, d, f, t, v, v)YesYesYesQ96ER3Protein SAAL1(b, m, r, u)YesQ8NB66Protein unc-13 homolog C(b, m, r, w)-YesYesQ9Y5H0Protocadherin gamma-A3(b, m, t, v)-Yes-Q9UBK7Rab-like protein 2A(c, m, t, u)YesQ2PPJ7Ral GTPase-activating protein subunit alpha-2(b, m, t, w)YesP50120Replication factor C subunit 1(b, m, t, w)YesQ8N392Rho GTPase-activating protein 18(c, m, t, u)-Yes-Q96QB1Rho GTPase-activating protein 7(b, m, t, w)-Yes-  | Q9Y520       | Protein PRRC2C <sup>(b, l, o, x)</sup>   | Yes | -   | Yes |
| Q96ER3Protein SAAL1(b, m, r, v)-YesQ8NB66Protein unc-13 homolog C(b, m, r, w)-YesQ9Y5H0Protocadherin gamma-A3(b, m, t, v)-YesQ9UBK7Rab-like protein 2A(c, m, t, v)Yes-Q2PPJ7Ral GTPase-activating protein subunit alpha-2(b, m, t, w)Yes-P35251Replication factor C subunit 1(b, m, t, w)Yes-P50120Retinol-binding protein 2(c, m, t, w)-YesQ8N392Rho GTPase-activating protein 18(c, m, t, w)-YesQ96QB1Rho GTPase-activating protein 7(b, m, t, w)-Yes   | P05109       | Protein \$100-A8 <sup>(a, b, c, d, f, j, r, u, v)</sup>                              | Yes | Yes | Yes |
| Q8NB66Protein unc-13 homolog C(b, m, r, w)-YesYesQ9Y5H0Protocadherin gamma-A3(b, m, t, v)-Yes-Q9UBK7Rab-like protein 2A(c, m, t, u)YesQ2PPJ7Ral GTPase-activating protein subunit alpha-2(b, m, t, w)YesP35251Replication factor C subunit 1(b, m, t, w)YesP50120Retinol-binding protein 2(c, m, t, w)-Yes-Q8N392Rho GTPase-activating protein 18(c, m, t, u)-Yes-Q96QB1Rho GTPase-activating protein 7(b, m, t, w)-Yes-  | Q96ER3       | Protein SAAL1 <sup>(b, m, r, u)</sup>  | -   | -   | Yes |
| Q9Y5H0Protocadherin gamma-A3(b, m, t, v)-Yes-Q9UBK7Rab-like protein 2A(c, m, t, v)YesQ2PPJ7Ral GTPase-activating protein subunit alpha-2(b, m, t, w)YesP35251Replication factor C subunit 1(b, m, t, w)YesP50120Retinol-binding protein 2(c, m, t, w)-Yes-Q8N392Rho GTPase-activating protein 18(c, m, t, u)-Yes-Q96QB1Rho GTPase-activating protein 7(b, m, t, u)-Yes-   | Q8NB66       | Protein unc-13 homolog C <sup>(b, m, r, w)</sup>                                     | -   | Yes | Yes |
| Q9UBK7Rab-like protein 2A <sup>(c, m, t, u)</sup> Yes-Q2PPJ7Ral GTPase-activating protein subunit alpha-2 <sup>(b, m, t, w)</sup> Yes-P35251Replication factor C subunit 1 <sup>(b, m, t, w)</sup> Yes-P50120Retinol-binding protein 2 <sup>(c, m, t, w)</sup> -YesQ8N392Rho GTPase-activating protein 18 <sup>(c, m, t, u)</sup> -YesQ96QB1Rho GTPase-activating protein 7 <sup>(b, m, t, u)</sup> -Yes  | Q9Y5H0       | Protocadherin gamma-A3 <sup>(b, m, t, v)</sup>                                       | -   | Yes | -   |
| Q2PPJ7Ral GTPase-activating protein subunit alpha-2 <sup>(b, m, t, w)</sup> Yes-P35251Replication factor C subunit 1 <sup>(b, m, t, w)</sup> Yes-P50120Retinol-binding protein 2 <sup>(c, m, t, w)</sup> -YesQ8N392Rho GTPase-activating protein 18 <sup>(c, m, t, u)</sup> -YesQ96QB1Rho GTPase-activating protein 7 <sup>(b, m, t, u)</sup> -Yes  | Q9UBK7       | Rab-like protein 2A <sup>(c, m, t, u)</sup>  | Yes | -   | -   |
| P35251Replication factor C subunit 1 <sup>(b, m, t, w)</sup> Yes-P50120Retinol-binding protein 2 <sup>(c, m, t, w)</sup> -YesQ8N392Rho GTPase-activating protein 18 <sup>(c, m, t, u)</sup> -YesQ96QB1Rho GTPase-activating protein 7 <sup>(b, m, t, u)</sup> -Yes  | Q2PPJ7       | Ral GTPase-activating protein subunit alpha-2 <sup>(b, m, t, w)</sup>                | Yes | -   | -   |
| P50120Retinol-binding protein 2 <sup>(c, m, t, w)</sup> -YesQ8N392Rho GTPase-activating protein 18 <sup>(c, m, t, u)</sup> -Yes-Q96QB1Rho GTPase-activating protein 7 <sup>(b, m, t, u)</sup> -Yes-   | P35251       | Replication factor C subunit $1^{(b, m, t, w)}$                                      | Yes | -   | -   |
| Q8N392Rho GTPase-activating protein 18(c, m, t, u)-Yes-Q96QB1Rho GTPase-activating protein 7(b, m, t, u)-Yes-   | P50120       | Retinol-binding protein 2 <sup>(c, m, t, w)</sup>                                    | -   | -   | Yes |
| Q96QB1 Rho GTPase-activating protein 7 <sup>(b, m, t, u)</sup> - Yes -  | Q8N392       | Rho GTPase-activating protein 18 <sup>(c, m, t, u)</sup>                             | -   | Yes | -   |
|   | Q96QB1       | Rho GTPase-activating protein 7 <sup>(b, m, t, u)</sup>                              | -   | Yes | -   |

| Continuation |  |     |     |     |
|--------------|--|-----|-----|-----|
| P49756       | RNA-binding protein 25 <sup>(b, m, t, u)</sup>                                       | Yes | Yes | -   |
| B4DTS2       | Serine/threonine-protein kinase <sup>(d, m, t, u)</sup>                              | Yes | Yes | Yes |
| Q6ZWH5       | Serine/threonine-protein kinase Nek10 $^{(d, m, t, w)}$                              | -   | -   | Yes |
| P02787       | Serotransferrin <sup>(c, m, t, w)</sup>  | Yes | Yes | Yes |
| P02768       | Serum albumin <sup>(a, b, c, g, o, u, w)</sup>                                       | Yes | Yes | Yes |
| O43166       | Signal-induced proliferation-associated 1-like protein $1^{(b,d,m,t,w)}$             | Yes | -   | -   |
| Q9Y5Y9       | Sodium channel protein type 10 subunit alpha $^{(b,m,t,w)}$                          | -   | Yes | -   |
| P35499       | Sodium channel protein type 4 subunit alpha $^{(b, m, t, w)}$                        | -   | Yes | -   |
| Q9Y6X4       | Soluble lamin-associated protein of 75 kDa <sup>(e, m, t, w)</sup>                   | -   |     | Yes |
| Q5M8T2       | Solute carrier family 35 member D3 <sup>(b, m, t, w)</sup>                           | -   | Yes | -   |
| Q96GZ6       | Solute carrier family 41 member 3 <sup>(c, m, r, w)</sup>                            | -   | -   | Yes |
| P11277       | Spectrin beta chain_ erythrocytic <sup>(d, f, m, q, w)</sup>                         | -   | -   | Yes |
| Q01082       | Spectrin beta chain_ non-erythrocytic 1 <sup>(d, m, t, w)</sup>                      | -   | -   | Yes |
| O15020       | Spectrin beta chain_ non-erythrocytic $2^{(d, m, t, w)}$                             | -   | -   | Yes |
| Q9H254       | Spectrin beta chain_ non-erythrocytic $4^{(d, m, t, w)}$                             | -   | -   | Yes |
| Q8WXA9       | Splicing regulatory glutamine/lysine-rich protein 1 <sup>(b, m, t, w)</sup>          | -   | Yes | Yes |
| P02808       | Statherin <sup>(b, e, i, l, o, u)</sup>  | -   | Yes | Yes |
| Q9NTJ3       | Structural maintenance of chromosomes protein $4^{(b,\ m,\ q,\ w)}$                  | -   | Yes | -   |
| Q9Y2K9       | Syntaxin-binding protein 5-like <sup>(b, c, m, t, w)</sup>                           | Yes | -   | -   |
| Q92609       | TBC1 domain family member 5 <sup>(b, c, m, t, w)</sup>                               | -   | Yes | -   |
| Q495A1       | T-cell immunoreceptor with Ig and ITIM domains $^{(b,\ m,\ r,\ w)}$                  | -   | -   | Yes |
| Q8IVF5       | T-lymphoma invasion and metastasis-inducing protein $2^{(b, m, r, w)}$               | Yes | Yes | -   |
| P19484       | Transcription factor EB <sup>(b, m, s, w)</sup>                                      | -   | Yes | Yes |
| P46100       | Transcriptional regulator ATRX <sup>(b, m, r, w)</sup>                               | Yes | Yes | -   |
| Q9Y4A5       | Transformation/transcription domain-associated protein <sup>(b, m, t, u, w)</sup>    | -   | Yes | -   |
| Q8NFB2       | Transmembrane protein 185A <sup>(b, m, t, w)</sup>                                   | Yes | -   | -   |
| Q9H7F4       | Transmembrane protein 185B <sup>(b, m, t, w)</sup>                                   | Yes | -   | -   |
| Q8NDV7       | Trinucleotide repeat-containing gene 6A protein <sup>(b, f, m, t, u, w)</sup>        | -   | Yes | -   |
| Q96LD4       | Tripartite motif-containing protein 47 <sup>(b, m, o, w)</sup>                       | -   | Yes | -   |
| Q6EMB2       | Tubulin polyglutamylase TTLL5 <sup>(b, m, t, w)</sup>                                | Yes | -   | -   |
| Q9UBB9       | Tuftelin-interacting protein 11 <sup>(b, d, m, t, w)</sup>                           | -   | -   | Yes |
| O15327       | Type II inositol 3_4-bisphosphate 4-phosphatase <sup>(b, m, t, w)</sup>              | -   | Yes | Yes |
| P17706       | Tyrosine-protein phosphatase non-receptor type $2^{(b,d,e,m,t,w)}$                   | -   | -   | Yes |
| Q9P2H5       | Ubiquitin carboxyl-terminal hydrolase $35^{(b, m, t, u)}$                            | Yes | -   | -   |
| P62256       | Ubiquitin-conjugating enzyme E2 H <sup>(b, m, t, u)</sup>                            | -   | -   | Yes |
| B2RTY4       | Unconventional myosin-IXa <sup>(c, m, t, w)</sup>                                    | -   | Yes | Yes |
| Q13459       | Unconventional myosin-IXb <sup>(c, m, t, w)</sup>                                    | -   | Yes | -   |
| Q9UID3       | Vacuolar protein sorting-associated protein 51 homolog <sup>(b, c, m, s, w)</sup>    | -   | Yes | -   |
| Q9H0V9       | VIP36-like protein <sup>(b, c, m, t, w)</sup>  | -   | -   | Yes |
| P54289       | Voltage-dependent calcium channel subunit alpha-2/delta-1 <sup>(b, f, m, t, w)</sup> | -   | Yes | -   |

Continue

Continuation

| Continuation |  |     |     |     |
|--------------|--|-----|-----|-----|
| Q9ULJ3       | Zinc finger and BTB domain-containing protein $21^{(b,m,p,\upsilon)}$                    | Yes | Yes | Yes |
| Q8IWY8       | Zinc finger and SCAN domain-containing protein $29^{\scriptscriptstyle (b,\ m,\ p,\ u)}$ | -   | Yes | -   |
| Q86VM9       | Zinc finger CCCH domain-containing protein 18 <sup>(b, m, p, u)</sup>                    | Yes | Yes | -   |
| P37275       | Zinc finger E-box-binding homeobox $1^{(b,\mbox{ m, p, u})}$                             | Yes | -   | -   |
| O60315       | Zinc finger E-box-binding homeobox $2^{(b, m, p, u)}$                                    | Yes | -   | Yes |
| O43296       | Zinc finger protein 264 <sup>(b, m, p, u)</sup>  | Yes | -   | -   |
| Q9HBT7       | Zinc finger protein 287 <sup>(b, m, p, u)</sup>  | -   | Yes | -   |
| Q8IZ26       | Zinc finger protein 34 <sup>(b, m, p, u)</sup>   | -   | -   | Yes |
| Q8N8E2       | Zinc finger protein 513 <sup>(b, m, p, u)</sup>  | -   | -   | Yes |
| Q9C0D4       | Zinc finger protein $518B^{(b, m, p, u)}$  | Yes | -   | -   |
| Q3ZCX4       | Zinc finger protein 568 <sup>(b, m, p, u)</sup>  | -   | Yes | -   |
| Q96N77       | Zinc finger protein $641^{(b, m, p, u)}$   | -   | -   | Yes |
| O75290       | Zinc finger protein 780A <sup>(b, m, p, u)</sup>   | -   | -   | Yes |
| Q6ZQV5       | Zinc finger protein 788 <sup>(b, m, p, u)</sup>  | Yes | -   | -   |

Proteins were classified according to: General Function: <sup>a)</sup> metabolism; <sup>b)</sup> biological process; <sup>c)</sup> transport; <sup>d)</sup> structure and structural organization; <sup>e)</sup> information pathways; <sup>†)</sup> miscellanea; Function in AEP: <sup>a)</sup> metabolism; <sup>h)</sup> tissue regeneration; <sup>ii)</sup> antimicrobial; <sup>iii</sup> immune response; <sup>k)</sup> lubrication; <sup>b</sup> biomineralization; <sup>m)</sup> unknown biological function; Origin: <sup>n)</sup> cytoplasm origin; <sup>o)</sup> extracellular origin; <sup>p)</sup> nucleus origin; <sup>q)</sup> cytoskeleton origin; <sup>f)</sup> intracellular origin; <sup>s)</sup> membrane origin; <sup>†)</sup> unknown protein origin; Interaction: <sup>u)</sup> protein/protein interaction; <sup>v)</sup> calcium/phosphate binding; <sup>w)</sup> other molecular interaction; <sup>x)</sup> unknown molecular interaction.

study, in which the AEP was collected from the vestibular surfaces.<sup>7</sup>

One interesting finding of the present study was the number of different types of histones. These proteins are very rich in lysine and arginine residues. This makes it easier to identify these proteins in proteomic studies, since tryptic peptides are generated upon the polypeptide chain cleavage of these residues. However, one finding was quite notable: H3 histones were found solely in the C group, while histone H2B isoforms were found only in the reflux groups (GE and GNE). The reason is not clear, but we should consider that H2B histones have a serine residue that can be phosphorylated in the N-terminus region. On the other hand, histone H3 has 2 serine residues that can be phosphorylated at the N-terminus. However, this region also contains at least 4 lysine residues that can be methylated.<sup>13,14</sup> This can make access of the phosphate ion to the tooth surface difficult, thus reducing the binding of this type of histone to the calcium in the hydroxyapatite. The increase in after-radiotherapy histones is related to the DNA methylation process, which acts concomitantly with histone acetyltransferases.



**Figure.** Venn diagram showing the number of proteins identified in the control (no GERD, no dental erosion), GE (GERD and dental erosion), and GNE (GERD but no dental erosion) groups.

This process is based on the epigenetic mechanisms associated with cellular memory and identity, which influence the cell environment and control

| °Accession | Destain a sure  | PLGS  | <sup>b</sup> Ratio: |
|------------|---|-------|---------------------|
| Number     | Profein name  | Score | GE:CL               |
| Q9ULJ3     | Zinc finger and BTB domain-containing protein 21                | 225   | 4.85                |
| P18615     | Negative elongation factor E                                    | 236   | 3.94                |
| P02787     | Serotransferrin   | 1819  | 1.40                |
| P05109     | Protein \$100-A8  | 11183 | 1.25                |
| P24158     | Myeloblastin  | 1361  | 0.84                |
| P62736     | Actin_ aortic smooth muscle                                     | 4078  | 0.73                |
| P60709     | Actin_ cytoplasmic 1  | 4078  | 0.72                |
| P49756     | RNA-binding protein 25  | 8013  | 0.70                |
| Q15147     | 1-phosphatidylinositol 45-bisphosphate phosphodiesterase beta-4 | 4085  | 0.64                |
| P63261     | Actin_ cytoplasmic 2  | 4078  | 0.63                |
| P68032     | Actin_ alpha cardiac muscle 1                                   | 4078  | 0.63                |
| P68133     | Actin_ alpha skeletal muscle                                    | 4078  | 0.63                |
| P04280     | Basic salivary proline-rich protein 1                           | 1868  | 0.46                |
| P38398     | Breast cancer type 1 susceptibility protein                     | 2136  | 0.38                |
| Q5SW79     | Centrosomal protein of 170 kDa                                  | 377   | 0.33                |
| P07197     | Neurofilament medium polypeptide                                | 7061  | 0.27                |
| P59665     | Neutrophil defensin 1   | 7061  | 0.27                |
| Q16352     | Alpha-internexin  | 327   | 0.09                |
| °Accession |   | PLGS  | <sup>b</sup> Ratio: |
| Number     | Protein name  | Score | GNE:CL              |
| P60709     | Actin_ cytoplasmic 1  | 4078  | 1.58                |
| P05109     | Protein \$100-A8  | 11183 | 0.21                |
| Q16352     | Alpha-internexin  | 327   | 0.20                |
| P04083     | Annexin A1  | 2838  | 0.19                |
| °Accession |   | PLGS  | <sup>b</sup> Ratio: |
| Number     | Protein name  | Score | GNE:GE              |
| P60709     | Actin_ cytoplasmic 1  | 4078  | 2.20                |
| P62736     | Actin_ aortic smooth muscle                                     | 4078  | 2.16                |
| P24158     | Myeloblastin  | 1361  | 1.40                |
| Q5THK1     | Protein PRR14L  | 158   | 0.21                |
| P04083     | Annexin A1  | 2838  | 0.20                |
| P05109     | Protein \$100-A8  | 11183 | 0.17                |
| Q9UIF7     | Adenine DNA glycosylase   | 338   | Cc                  |
| Q9UHX3     | Adhesion G protein-coupled receptor E2                          | 161   | С                   |
| O00213     | Amyloid beta A4 precursor protein-binding family B member 1     | 247   | С                   |
| Q7L311     | Armadillo repeat-containing X-linked protein 2                  | 230   | С                   |
| P51689     | Arylsulfatase D   | 272   | С                   |
| 095342     | Bile salt export pump   | 201   | С                   |

**Table 3.** Proteins expressed differentially in the acquired enamel pellicle collected from patients with gastroesophageal reflux disease (GERD) and erosive tooth wear (GE), GERD but no erosive tooth wear (GNE), or controls (no GERD, no erosive tooth wear; C).

| Continuation |  |     |          |
|--------------|--|-----|----------|
| Q9ULB4       | Cadherin-9   | 237 | С        |
| Q96G23       | Ceramide synthase 2  | 263 | С        |
| Q8TBZ0       | Coiled-coil domain-containing protein 110                              | 248 | С        |
| Q96NY9       | Crossover junction endonuclease MUS81                                  | 175 | С        |
| Q13443       | Disintegrin and metalloproteinase domain-containing protein 9          | 190 | С        |
| Q5T890       | DNA excision repair protein ERCC-6-like 2                              | 51  | С        |
| Q8N7B9       | EF-hand calcium-binding domain-containing protein 3                    | 212 | С        |
| Q6PCB8       | Embigin  | 208 | С        |
| Q969X5       | Endoplasmic reticulum-Golgi intermediate compartment protein 1         | 215 | С        |
| O15360       | Fanconi anemia group A protein   | 188 | С        |
| Q9BZK7       | F-box-like/WD repeat-containing protein TBL1XR1                        | 224 | С        |
| P23142       | Fibulin-1  | 318 | С        |
| P61978       | Heterogeneous nuclear ribonucleoprotein K                              | 159 | С        |
| P68431       | Histone H3.1   | 268 | С        |
| Q16695       | Histone H3.1t  | 268 | С        |
| Q71DI3       | Histone H3.2   | 668 | С        |
| P84243       | Histone H3.3   | 268 | С        |
| Q6NXT2       | Histone H3.3C  | 268 | С        |
| P14735       | Insulin-degrading enzyme   | 301 | С        |
| Q86VS3       | IQ domain-containing protein H   | 68  | С        |
| Q7Z4W3       | Keratin-associated protein 19-3  | 343 | С        |
| C9JBD0       | KRAB domain-containing protein 1                                       | 531 | С        |
| Q8N653       | Leucine-zipper-like transcriptional regulator 1                        | 304 | С        |
| O95711       | Lymphocyte antigen 86  | 222 | С        |
| P40925       | Malate dehydrogenase_ cytoplasmic                                      | 119 | С        |
| P15941       | Mucin-1  | 202 | С        |
| Q04721       | Neurogenic locus notch homolog protein 2                               | 112 | С        |
| O14832       | Phytanoyl-CoA dioxygenase_ peroxisomal                                 | 269 | С        |
| Q13310       | Polyadenylate-binding protein 4  | 43  | С        |
| Q9H4B0       | Probable tRNA N6-adenosine threonylcarbamoyltransferase_ mitochondrial | 127 | С        |
| Q3B820       | Protein FAM161A  | 316 | С        |
| Q9H0Q0       | Protein FAM49A   | 293 | С        |
| Q9UBK7       | Rab-like protein 2A  | 151 | С        |
| Q2PPJ7       | Ral GTPase-activating protein subunit alpha-2                          | 156 | С        |
| P35251       | Replication factor C subunit 1   | 167 | С        |
| O43166       | Signal-induced proliferation-associated 1-like protein 1               | 251 | С        |
| Q9Y2K9       | Syntaxin-binding protein 5-like  | 191 | С        |
| Q8NFB2       | Transmembrane protein 185A   | 213 | С        |
| Q9H7F4       | Transmembrane protein 185B   | 223 | С        |
| Q6EMB2       | Tubulin polyglutamylase TTLL5  | 172 | С        |
|              |  |     | <b>C</b> |

| Continuation |  |     |                 |
|--------------|--|-----|-----------------|
| Q9P2H5       | Ubiquitin carboxyl-terminal hydrolase 35                         | 267 | С               |
| P37275       | Zinc finger E-box-binding homeobox 1                             | 234 | С               |
| O43296       | Zinc finger protein 264  | 187 | С               |
| Q9C0D4       | Zinc finger protein 518B   | 227 | С               |
| Q6ZQV5       | Zinc finger protein 788  | 226 | С               |
| Q08AH3       | Acyl-coenzyme A synthetase ACSM2A_ mitochondrial                 | 209 | GE <sup>d</sup> |
| Q8IUA7       | ATP-binding cassette sub-family A member 9                       | 238 | GE              |
| Q9H4G0       | Band 4.1-like protein 1  | 581 | GE              |
| Q9Y6E2       | Basic leucine zipper and W2 domain-containing protein 2          | 147 | GE              |
| Q86WR0       | Coiled-coil domain-containing protein 25                         | 591 | GE              |
| Q13363       | C-terminal-binding protein 1                                     | 566 | GE              |
| Q5M775       | Cytospin-B   | 402 | GE              |
| Q9Y485       | DmX-like protein 1   | 117 | GE              |
| Q9UBZ9       | DNA repair protein REV1  | 309 | GE              |
| Q14152       | Eukaryotic translation initiation factor 3 subunit A             | 101 | GE              |
| O60907       | F-box-like/WD repeat-containing protein TBL1X                    | 415 | GE              |
| P09958       | Furin  | 121 | GE              |
| Q8WUA4       | General transcription factor 3C polypeptide 2                    | 388 | GE              |
| Q08378       | Golgin subfamily A member 3                                      | 360 | GE              |
| Q9Y4L1       | Hypoxia up-regulated protein 1                                   | 270 | GE              |
| POC7H9       | Inactive ubiquitin carboxyl-terminal hydrolase 17-like protein 7 | 154 | GE              |
| Q86WG5       | Myotubularin-related protein 13                                  | 297 | GE              |
| Q92823       | Neuronal cell adhesion molecule                                  | 275 | GE              |
| P11940       | Polyadenylate-binding protein 1                                  | 244 | GE              |
| P98161       | Polycystin-1   | 90  | GE              |
| Q6UN15       | Pre-mRNA 3'-end-processing factor FIP1                           | 175 | GE              |
| Q5VTL8       | Pre-mRNA-splicing factor 38B                                     | 286 | GE              |
| O43143       | Pre-mRNA-splicing factor ATP-dependent RNA helicase DHX15        | 126 | GE              |
| P78504       | Protein jagged-1   | 92  | GE              |
| Q9Y5H0       | Protocadherin gamma-A3   | 193 | GE              |
| Q8N392       | Rho GTPase-activating protein 18                                 | 185 | GE              |
| Q96QB1       | Rho GTPase-activating protein 7                                  | 131 | GE              |
| Q9Y5Y9       | Sodium channel protein type 10 subunit alpha                     | 165 | GE              |
| P35499       | Sodium channel protein type 4 subunit alpha                      | 129 | GE              |
| Q5M8T2       | Solute carrier family 35 member D3                               | 133 | GE              |
| Q9NTJ3       | Structural maintenance of chromosomes protein 4                  | 194 | GE              |
| Q92609       | TBC1 domain family member 5                                      | 255 | GE              |
| Q9Y4A5       | Transformation/transcription domain-associated protein           | 287 | GE              |
| Q8NDV7       | Trinucleotide repeat-containing gene 6A protein                  | 171 | GE              |
| Q96LD4       | Tripartite motif-containing protein 47                           | 211 | GE              |
|              |  |     | Continue        |

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| Continuation |  |     |                         |
|--------------|--|-----|-------------------------|
| Q13459       | Unconventional myosin-IXb                                  | 268 | GE                      |
| Q9UID3       | Vacuolar protein sorting-associated protein 51 homolog     | 369 | GE                      |
| P54289       | Voltage-dependent calcium channel subunit alpha-2/delta-1  | 147 | GE                      |
| Q8IWY8       | Zinc finger and SCAN domain-containing protein 29          | 174 | GE                      |
| Q9HBT7       | Zinc finger protein 287                                    | 283 | GE                      |
| Q3ZCX4       | Zinc finger protein 568                                    | 220 | GE                      |
| Q8NC06       | Acyl-CoA-binding domain-containing protein 4               | 263 | <b>GNE</b> <sup>e</sup> |
| Q8N142       | Adenylosuccinate synthetase isozyme 1                      | 102 | GNE                     |
| P15144       | Aminopeptidase N   | 251 | GNE                     |
| P02656       | Apolipoprotein C-III                                       | 141 | GNE                     |
| Q5T2S8       | Armadillo repeat-containing protein 4                      | 170 | GNE                     |
| O43491       | Band 4.1-like protein 2                                    | 263 | GNE                     |
| Q96IK1       | Biorientation of chromosomes in cell division protein 1    | 263 | GNE                     |
| Q96L14       | Cep170-like protein  | 263 | GNE                     |
| 075462       | Cytokine receptor-like factor 1                            | 327 | GNE                     |
| P48382       | DNA-binding protein RFX5                                   | 327 | GNE                     |
| Q12805       | EGF-containing fibulin-like extracellular matrix protein 1 | 513 | GNE                     |
| Q68CQ7       | Glycosyltransferase 8 domain-containing protein 1          | 454 | GNE                     |
| Q02643       | Growth hormone-releasing hormone receptor                  | 144 | GNE                     |
| O43719       | HIV Tat-specific factor 1                                  | 144 | GNE                     |
| P55899       | lgG receptor FcRn large subunit p51                        | 279 | GNE                     |
| Q8WWI1       | LIM domain only protein 7                                  | 215 | GNE                     |
| Q9P2S2       | Neurexin-2   | 215 | GNE                     |
| P58401       | Neurexin-2-beta  | 220 | GNE                     |
| Q16288       | NT-3 growth factor receptor                                | 174 | GNE                     |
| Q8NGQ1       | Olfactory receptor 9G4                                     | 150 | GNE                     |
| Q92508       | Piezo-type mechanosensitive ion channel component 1        | 169 | GNE                     |
| Q5GLZ8       | Probable E3 ubiquitin-protein ligase HERC4                 | 140 | GNE                     |
| Q96ER3       | Protein SAAL1  | 152 | GNE                     |
| P50120       | Retinol-binding protein 2                                  | 161 | GNE                     |
| Q6ZWH5       | Serine/threonine-protein kinase Nek10                      | 161 | GNE                     |
| Q9Y6X4       | Soluble lamin-associated protein of 75 kDa                 | 45  | GNE                     |
| Q96GZ6       | Solute carrier family 41 member 3                          | 605 | GNE                     |
| P11277       | Spectrin beta chain_ erythrocytic                          | 645 | GNE                     |
| Q01082       | Spectrin beta chain_ non-erythrocytic 1                    | 184 | GNE                     |
| O15020       | Spectrin beta chain_ non-erythrocytic 2                    | 186 | GNE                     |
| Q9H254       | Spectrin beta chain_ non-erythrocytic 4                    | 210 | GNE                     |
| Q495A1       | T-cell immunoreceptor with Ig and ITIM domains             | 193 | GNE                     |
| Q9UBB9       | Tuftelin-interacting protein 11                            | 203 | GNE                     |
| P17706       | Tyrosine-protein phosphatase non-receptor type 2           | 169 | GNE                     |
|              |  |     |                         |

| Continuation |                                   |     |     |
|--------------|-----------------------------------|-----|-----|
| P62256       | Ubiquitin-conjugating enzyme E2 H | 384 | GNE |
| Q9H0V9       | VIP36-like protein                | 410 | GNE |
| Q8IZ26       | Zinc finger protein 34            | 473 | GNE |
| Q8N8E2       | Zinc finger protein 513           | 473 | GNE |
| Q96N77       | Zinc finger protein 641           | 473 | GNE |
| 075290       | Zinc finger protein 780A          | 176 | GNE |

<sup>o</sup>Identification is based on ID proteins from Uniprot protein database, only reviewed (http://www.uniprot.org/); <sup>b</sup>Proteins with significantly altered expression are organized according to the ratio; <sup>c</sup>Indicates proteins found exclusively in the acquired enamel pellicle (AEP) of C patients (in alphabetical order); <sup>d</sup>Indicates proteins found exclusively in the acquired enamel pellicle (AEP) of GE patients (in alphabetical order); <sup>e</sup>Indicates proteins found exclusively in the acquired enamel pellicle (AEP) of GE patients (in alphabetical order); <sup>e</sup>Indicates proteins found exclusively in the acquired enamel pellicle (AEP) of GNE patients (in alphabetical order). Ratios highlighted in bold indicate proteins with higher increase or decrease in GNE compared with GE patients.

epigenetic regulation. Furthermore, non-coding RNA action and histone modification also interact in epigenetic regulation.<sup>15</sup> Acetylation is a type of histone alteration that plays an important role in modulating gene expression and cell cycles, and in neoplasm diffusion.<sup>16</sup> The acetylation process starts with the action of acetyltransferases, prompted by the binding of acetyl radicals and lysine residues of histone proteins, and results in the decompression of chromatin and transcriptional activity. Histones H2A and H2B are examples of proteins whose function is not only to act in DNA replication and repair, but also to constitute octamers with histones H3 and H4. They are also involved in the packaging of DNA in nucleosomes.<sup>17</sup> Interestingly, a study by Ventura et al.<sup>18</sup> showed that histone isoforms can be considered a strong prognostic biomarker in patients with head and neck cancer. This shows how an analysis of the AEP can benefit the patient by providing previous diagnosis and adequate treatment.

AEP proteins have long been known for their protective effect in the homeostasis of the oral cavity.<sup>11</sup> It is interesting to highlight that the focus on these proteins a few years ago was placed on those that were secreted in saliva.<sup>11</sup> However, other important sources of AEP proteins are exfoliated oral mucosa cells, and gingival cells that deliver their content to saliva. In this context, many proteins with an affinity for hydroxyapatite might be immobilized in the AEP, and play an important protective role against acid challenges. This is the case of histones, as well as other proteins recently identified in the AEP as acid-resistant, such as hemoglobin (HB).<sup>7,19-</sup> <sup>22</sup> Nevertheless, HB was evaluated in the present study because it was found to be higher in the AEP<sup>7</sup> and saliva<sup>22</sup> of GERD patients without ETW, compared with GERD patients with ETW. Moreover, bear in mind that HB has a strong affinity for hydroxyapatite, and that hydroxyapatite columns are used to purify this protein.<sup>23</sup> Interestingly, the adsorption rate of HB to hydroxyapatite increases as pH decreases.<sup>24</sup> GERD patients have an oral pH typically lower than that of healthy people.<sup>10</sup> This might increase the chance of HB adsorption onto dental surfaces.

Furthermore, a high number of phosphorylated and calcium-binding proteins were found among the proteins identified exclusively in the GNE group. Nearly 50% of a total of 40 distinct proteins of the GNE group (the vast majority being intracellular proteins) are phosphorylated or Ca-binding proteins, thus suggesting that they might interact intensively with hydroxyapatite. This finding was also observed in our previous publication<sup>7</sup>, and might also be implicated in acid resistant mechanisms.

Another important finding was that several isoforms of the spectrin beta-chain were found only in the AEP collected from the GNE group. These proteins interact with actin. Curiously, there were two actin isoforms that increased more than twofold in the GNE group, in comparison with the GE group. The spectrin beta-chain can also form protein complexes. Because it binds to actin, it could be involved in the formation of supramolecular aggregates in the second stage of AEP formation.<sup>25</sup>

As mentioned above, several intracellular proteins delivered to saliva after exfoliation of oral mucosa cells have the potential to bind to hydroxyapatite, or participate in supramolecular aggregates that bind to the precursor proteins in the AEP. This has also been observed in other recent studies,<sup>26-31</sup> and could have a central role in protecting the tooth surface against acid dissolution. Hence, it is worthwhile noting that not only do secreted salivary proteins participate in AEP formation, but oral mucosa proteins, gingival crevicular fluid and even bacteria also act strongly on the enamel pellicle. This indicates that the AEP protein could play a protective role against dental erosion caused by intrinsic acids. This topic should be investigated further in future studies.

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