Evolution of the mammalian lysozyme gene family

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Abstract

Background: Lysozyme c (chicken-type lysozyme) has an important role in host defense, and has been extensively studied as a model in molecular biology, enzymology, protein chemistry, and crystallography. Traditionally, lysozyme c has been considered to be part of a small family that includes genes for two other proteins, lactalbumin, which is found only in mammals, and calcium-binding lysozyme, which is found in only a few species of birds and mammals. More recently, additional testes-expressed members of this family have been identified in human and mouse, suggesting that the mammalian lysozyme gene family is larger than previously known.

Results: Here we characterize the extent and diversity of the lysozyme gene family in the genomes of phylogenetically diverse mammals, and show that this family contains at least eight different genes that likely duplicated prior to the diversification of extant mammals. These duplicated genes have largely been maintained, both in intron-exon structure and in genomic context, throughout mammalian evolution.

Conclusions: The mammalian lysozyme gene family is much larger than previously appreciated and consists of at least eight distinct genes scattered around the genome. Since the lysozyme c and lactalbumin proteins have acquired very different functions during evolution, it is likely that many of the other members of the lysozyme-like family will also have diverse and unexpected biological properties.

Background

The vertebrate lysozyme gene family has traditionally been considered to be composed of three genes: lysozyme c, lactalbumin, and calcium-binding lysozyme [1-4]. Lysozyme c, chicken-type (or conventional) lysozyme, is a bacteriolytic enzyme that is secreted into many body fluids of mammals (e.g., blood, tears, and milk) and is found at a high concentration in the eggs of many bird species [1,2,5]. Lysozyme c is widespread in nature; its protein and gene sequences have been characterized from numerous diverse vertebrate and non-vertebrate species [3,5,6]. Lactalbumin is related to lysozyme, with around 40% amino acid identity and nearly identical three-dimensional structure, but lacks its bacteriolytic activity [1,2,4,7]. Lactalbumin is expressed in lactating mammary glands, where it binds a calcium ion and modifies the activity of β-galactosyltransferase-1, such that the complex catalyzes the synthesis of lactose [2,4,7]. Lactalbumin has recently been shown to have a second activity in the gut, where it loses the calcium ion and binds a fatty acid; this new form of lactalbumin appears to promote apoptosis of tumor cells, and thus has been renamed HAMLET (human lactalbumin made lethal to tumors) [8]. Lactalbumin appears to be found only in mammals, and is widely distributed in this group. Calcium-binding lysozyme has bacteriolytic activity like lysozyme c, but also shares with lactalbumin the ability to bind a calcium ion. Calcium-binding lysozymes appear to be relatively rare; they have been found in the milk of only a few mammalian species (e.g., horse, dog, cat, seal, and echidna), as well as in the eggs (e.g., pigeon) and stomachs (e.g., hoatzin) of some bird species [3,9]. Indeed, calcium-binding lysozyme genes have not been reported for the human or rodent genomes.

Previous phylogenetic analyses of lysozyme c, lactalbumin, and calcium-binding lysozyme sequences had suggested that the earliest divergences within this gene family occurred between lysozyme c and the ancestor of the genes for lactalbumin and calcium-binding lysozyme,
and that this initial gene duplication may have preceded the divergence of the lineages leading to fish and mammals [10,11]. The separation of the lactalbumin and calcium-binding lysozyme genes was proposed to be more recent, with some studies [9,12] suggesting a divergence on the early mammalian lineage, which would be consistent with the restriction of the lactalbumin gene to mammals. In contrast, another study [11] suggested that the duplication generating the lactalbumin and calcium-binding lysozyme genes predated the bird-mammal divergence. Moreover, the orthology of the mammalian and avian calcium-binding lysozymes has even been questioned [3,11]. Thus, the origin of these mammalian lysozyme-like genes remains an open question.

Recently, cDNAs for several additional lysozyme-like sequences have been identified from human testis cDNA libraries [13-15]. These cDNAs were found to be encoded by genes that are now annotated by Ensembl [16] as LYZL (lysozyme-like): LYZL2, LYZL4, LYZL6 and LYZL3 (Synonym: SPACA3; SPACA, Sperm acrosome associated [15]). SPACA3 is also known as SPRSA [14] and SLLP1 [13]). The predicted protein sequences of some of these lysozyme-like sequences have amino acid substitutions at sites important for the catalytic activity of lysozyme, suggesting that these proteins would not be able to hydrolyze the glycosidic bonds of bacterial peptidoglycan [13,15]. Since these four new lysozyme-like genes (LYZL2, LYZL4, LYZL6, and SPACA3) are expressed predominantly in the testes, it has been suggested that they might have a role in reproduction [13-15,17]. Such a role has been shown for Lyl4 and Spaca3 in mice [18,19].

The identification of these LYZL genes in the human genome suggests that the mammalian lysozyme-like gene family is larger than previously appreciated, and raises the possibility that the lysozyme-like proteins encoded by these genes may have novel biological functions. Here we have used extensive similarity searches of the human and other vertebrate genomes. We thereby identified three additional intact lysozyme-like genes in the human genome; these have been annotated in the databases, but not reported in the literature. We have also identified multiple lysozyme-like genes in the genomes of diverse vertebrates. Using a combination of phylogenetic and genomic neighborhood (or synteny) analyses, wherein the relationships of the genes that flank the lysozyme-like genes in diverse species were examined, we demonstrate that homologs of the human lysozyme-like genes are found in the genomes of diverse mammalian species. Our analyses suggest that there were at least six, and perhaps as many as nine, diverse types (or subfamilies) of lysozyme-like genes in the genome of the common ancestor of all extant mammals, and that these diverse genes have been maintained on most mammalian lineages. This suggests that their protein products probably have essential biological functions that are yet to be identified.

Results and Discussion

Number of Lysozyme Genes in the Human Genome

To determine the size of the lysozyme-like gene family, we performed BLAST [20] similarity searches of the human genome for sequences that predict potential protein sequences similar to lysozyme c, and thereby identified a total of nine annotated genes (Table 1, and Additional file 1: Table S1). Of these nine annotated genes, six had previously been characterized: lysozyme c (LYZ) [21], lactalbumin (LALBA, Synonym: LYZL7) [22], LYZL2, LYZL4, LYZL6 [15], and SPACA3 (Synonyms: LYZL3, SPRSA, SLLP1) [13,14]. The three remaining genes identified in our BLAST searches – the LYZL1, SPACA5 (Synonym: LYZL5), and SPACASB genes – had been annotated as lysozyme-like in Ensembl [16], but have not been discussed in the literature. A tenth lysozyme-like sequence was later identified using our genomic neighborhood analysis (see below), but appears to be a pseudogene (ψLYSC1). These ten lysozyme-like sequences are distributed over five chromosomes in humans, with two genes on each on chromosomes 10, 17, and X, three genes on chromosome 12, and one gene on chromosome 3 (Table 1). Each of the potentially functional genes predicts a protein sequence about 140-150 amino acids long, similar to lysozyme c and lactalbumin. The mature regions of these proteins are readily aligned due to the presence of many highly conserved residues, including the eight cysteines known to be involved in disulfide bonds in lysozyme and lactalbumin (Figure 1). Each of the lysozyme-like genes has been annotated as being composed of four or five exons. The 140-150 amino acid coding regions are spread over four exons in each of the genes with the introns in exactly the same locations, including phases, as found in the lysozyme c and lactalbumin genes [5,6] (Figure 1). Moreover, the LYZL1, LYZL2, LYZL4, LYZL6, and SPACA3 genes are annotated as having an additional 5' exon, which in some cases might be translated to produce proteins that have longer N-terminal regions (especially in the case of SPACA3; not shown in Figure 1). Most of the predicted human proteins (Figure 1) show between 30% and 53% amino acid sequence identity in pairwise comparisons (Table 2), suggesting that the gene duplications that gave rise to them are fairly ancient (or, alternatively, that these proteins have evolved at extremely rapid rates). If these gene duplications were indeed ancient, then we would expect to find these genes in diverse mammalian species (as we did; see below). In contrast, the LYZL1/LYZL2 and SPACA5/SPACASB protein pairs are 97% and 100% identical, respectively
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Table 1 Chromosomal location of human lysozyme-like genes

| Gene | Chromosome | Strand | Position | Intact | Protein ID |
|------|------------|--------|----------|--------|------------|
| LYZ  | 12         | +      | 69,742,134-69,748,013 | Y       | ENSP00000261267 |
| LALB | 12         | -      | 48,961,468-48,963,829 | Y       | ENSP0000301046 |
| γLSC1| 12         | +      | 49,024,938-49,026,186 | N       | None (pseudogene) |
| LY21 | 10         | +      | 29,577,990-29,607,257 | Y       | ENSP0000364650 |
| LY22 | 10         | -      | 30,895,152-30,918,691 | Y       | ENSP0000364647 |
| LY24 | 3          | -      | 42,438,570-42,452,092 | Y       | ENSP0000287748 |
| LY26 | 17         | -      | 34,261,548-34,270,674 | Y       | ENSP0000292724 |
| SPAC3| 17         | +      | 31,318,887-31,324,895 | Y       | ENSP0000269053 |
| SPAC5| X          |        | 47,863,734-47,889,126 | Y       | ENSP0000366139 |
| SPAC5B|         |        | 47,986,603-47,991,995 | Y       | ENSP0000304762 |

* - Chromosomal localization, strand, and coordinates from release 57 of the human genome available in the Ensembl database (http://www.ensembl.org)

** - Y, full-length protein sequence; N, no intact open reading frame

* - Ensembl release 57 protein ID. None means no Ensembl Protein ID is available

(Table 2); this suggests that their genes likely duplicated fairly recently, and thus these duplicates are predicted to be more limited phylogenetically. The LYZL1 and LYZL2 genes are both located on human chromosome 10, but are separated by about 1 Mb; moreover, these genes are embedded within 60 kb long repeated sequences that are greater than 95% identical (not shown). This suggests that the LYZL1 and LYZL2 gene duplicates were generated as part of a recent segmental duplication on chromosome 10. Likewise, the SPAC5 and SPACASB genes are both on the X chromosome, separated by about 120 kb, and are within long (~100 kb) repeated DNA sequences that have high sequence identity (not shown). Thus, the LYZL1/LYZL2 and SPAC5/SPACASB gene pairs both appear to have originated from relatively recent and large genomic segmental duplications, a common form of gene duplication in mammals [23]. The high sequence identity of these gene pairs also could, at least in part, to concerted evolution [24], as discussed below.

Lysozyme Genes in Other Vertebrate Genomes

To determine whether an expanded lysozyme-like gene family is a general feature of mammalian (and other vertebrate) genomes, we conducted intensive homology searches for lysozyme-like genes in all vertebrate genomes available in the Ensembl and PreEnsembl databases (http://www.ensembl.org, http://pre.ensembl.org/index.html) [16,25]. These results are listed in Supplementary Table 1 (Additional file 1: Table S1), and summarized for mammals in Figure 2. At least one lysozyme-like gene was found in each vertebrate genome, with the exception of the lamprey. Multiple lysozyme-like genes were identified in all of the mammalian species ranging from 5 in opossum to 18 in cow. The fewest number of genes were found in bony fish, where only one gene was identified per genome, and in birds, where at most two genes were identified per genome. The anole lizard and Xenopus tropicalis, the lone representatives of reptiles and amphibians with sequenced genomes, had 8 and 16 lysozyme-like genes, respectively.

Importantly, the BLAST searches identified potential orthologs in most mammalian genomes (Figure 2) of all of the divergent lysozyme-like genes found in the
Table 2 Pairwise identity, in percent, of human lysozyme-like protein sequences

|        | LALBA | LYZL1 | LYZL2 | LYZL4 | LYZL6 | SPACA3 | SPACAS5 | SPACAS5B |
|--------|-------|-------|-------|-------|-------|--------|---------|---------|
| LYZ    | 37    | 49    | 49    | 41    | 45    | 53     | 44      | 44      |
| LALBA  | 36    | 36    | 30    | 32    | 33    | 32     | 32      | 32      |
| LYZL1  | 97    | 43    | 43    | 46    | 47    | 47     | 47      | 47      |
| LYZL2  |       | 43    | 43    | 46    | 47    | 47     | 47      | 47      |
| LYZL4  |       |       | 46    | 47    | 40    | 40     | 40      | 40      |
| LYZL6  |       |       |       | 45    | 47    | 47     | 47      | 47      |
| SPACA3 |       |       |       |       |       | 49     | 49      | 49      |
| SPACAS5|       |       |       |       |       |        | 100     |         |

Figure 2 Diversity and evolution of mammalian lysozyme-like genes. A phylogeny, adapted from recent phylogenomic analyses [71-73], of the mammalian species examined in this study is shown, branch lengths are not proportional to evolutionary time. The numbers of each type of lysozyme-like gene identified in the mammalian genomes are shown by symbols on the right of the Figure. Species with genomes of higher quality (better coverage and assembly) are labeled in bold and with asterisks. The status of the genes in each genome are indicated by symbols, listed in the key box in the figure, where solid boxes indicate genes that are predicted to encode full-length sequences, open boxes indicate incomplete genes, which may be due to either incomplete genome sequences or may be pseudogenes, the Greek letter psi identifies well-characterized pseudogenes, zero indicates that no gene was identified in our searches, and the ‘not’ sign indicates genes that appear to have been deleted from the genomes. The question mark mark beside the sloth Lysl2 gene indicates that this is a tentative assignment. Inferred evolutionary events are mapped onto their most likely lineage, based upon the data presented in this figure, coupled with our various evolutionary analyses. Duplication of genes is indicated by a gene name with an arc over it. Gene deletion is indicated by an X preceding the name of the deleted gene. X? means that the deletion is uncertain, as the missing gene may actually exist in a gap in the descendant genome (s). Pseudogene generation is indicated by the psi (ψ) that precedes the gene name; if the pseudogene is associated with a gene duplication event it also has an arc. If multiple events are shown on a lineage, the order shown does not imply historical order of occurrence.
human genome \((LYZ, LALB, LYZL1/2, LYZL4, LYZL6, SPACA3, \text{and} SPACAS5)\). Initial assignments of orthology of these mammalian genes were based upon sequence similarity, but were subsequently confirmed by performing genomic neighborhood and phylogenetic analyses (see below). Our combined findings about gene number and orthologous relationships of the mammalian lysozyme-like gene family members are outlined in Figure 2. In contrast to the mammalian genes, the lysozyme-like genes found in the other vertebrate genomes could not be readily classified into the above subfamilies based upon sequence similarities; this is because these genes and their encoded proteins displayed similar levels of sequence identity to all of the different mammalian paralogs. Furthermore, no evidence of synteny with the mammalian genes was found for any of the non-mammalian vertebrate genes, except for the lysozyme \(c\) gene. Thus, none of the non-mammalian vertebrate lysozyme-like genes could be definitively classified as orthologs of any of the lysozyme-like mammalian genes, other than \(Lyz\) itself.

Many of the genes that were identified in our searches were only partial sequences, most likely due to the incomplete nature of the genomes in question. However, all but one of these genes were consistent with a structure similar to that of the mammalian lysozyme and lactalbumin genes—i.e., their coding regions appeared to be composed of four exons having similar intron-exon structures \([5,6]\). The lone exception was a \(Lyzl1/2\)-like gene found in the treeshrew (from Genescap-fold_6044), which had a nearly full-length coding sequence that contained stop codons and frameshifts, but no introns; thus, this gene appears to be a processed pseudogene. Taken together, these observations suggest that essentially all of the vertebrate lysozyme-like genes have been generated by duplications of genomic DNA, rather than by reverse-transcription and insertion into genome.

**Phylogeny of Vertebrate Lysozymes**

The presence of multiple lysozyme-like genes in all mammalian genomes, as well as in the genomes of several other vertebrate species, raises the possibility that the lysozyme-like gene family may have amplified early in vertebrate evolution. To examine this issue, and to further establish the orthology-paralogy relationships of lysozyme-like genes, we conducted a series of phylogenetic analyses (Figure 3, and Additional files 2-4: Figures S1-S3). While many of our phylogenetic analyses, including the one shown in Figure 3 (and Additional file 3: Figure S2), suggested that the \(Lalba\) clade was the earliest diverging lineage and that most of the \(Lyzl\) and \(Spaca\) genes \((Lyzl1/2, Lyzl4, Lyzl6, Spaca3, \text{and} Spaca5, \text{but not} Lyzl8)\) were most closely related to each other, these relationships were not consistently found \(\text{e.g., see Additional file 2: Figure S1}\). Therefore, the phylogenetic analyses are inconclusive concerning the relationships of the different subfamilies of mammalian lysozyme-like genes.

The phylogenetic trees also suggested the possibility that at least some of the gene divergences occurred very early in vertebrate evolution, \(i.e.,\) prior to the mammal-fish divergence. For example, the mammalian \(Lyz\) gene sequences were found to branch with \(Lyz\) genes from fish, rather than with the other mammalian lysozyme-like genes (Figure 3); if this branching order reflects the
Figure 3 Phylogeny of vertebrate lysozyme-like genes. Bayesian phylogenetic tree of vertebrate lysozyme genes. The tree shown was generated with MrBayes [60,61] using DNA sequences of diverse vertebrate lysozyme-like sequences. (For sequences see Additional files 1 and 18: Tables S1 and S2 for sequences.) The alignment used to generate the tree is shown in Additional file 19: Figure S17. The trees were built after 2,000,000 generations with nst = 6 and rates = gamma (the TrNef+I+G model was selected as the best model by ModelTest [66-68]). Insect and amphioxus sequences were used to root the tree. Solid diamonds indicate nodes that represent the gene duplication events that generated the different types of lysozyme-like genes, or subfamilies, found in mammals. The support for each clade containing each subfamily of mammalian lysozyme-like gene is boxed, with posterior probabilities above the lineage and bootstrap support from maximum likelihood analysis below (from Additional file 3: Figure S2). The remaining lineages only display the posterior probabilities from the Bayesian analysis.
actual evolutionary history of the genes (rather than phylogenetic affinity based upon conserved lysozyme protein structure and function), then the Lyz gene lineage must have diverged from the other lysozyme-like genes prior to the mammal-fish divergence. Again, if this were true, then many species lineages must have deleted the duplicates from their genomes. Below, we discuss each subfamily of lysozyme-like gene in the mammals, and consider their potential non-mammalian orthologs.

Lysozyme c (Lyz) Genes

The Lyz gene is the best-studied lysozyme-like gene, and has been extensively characterized in many species [1-6]. Our genome searches and phylogenetic analyses identified many genes that appear to be orthologous to Lyz in diverse vertebrates (Figure 3, and Additional files 1-3: Table S1, and Figures S1 and S2). To confirm the orthology of the Lyz genes, we used a genomic neighborhood analysis, wherein we examined the orthology of the genes that flank the Lyz gene in diverse species. The avian and mammalian Lyz genes are flanked by the Cpsf6 and Yeats4 genes (Figure 4). This organization is maintained in species with tandemly duplicated Lyz genes, such as rodents and cow, where the Cpsf6 and Yeats4 genes are found flanking a cluster of Lyz genes (Figure 4). A slightly different organization is seen in the opossum, which has four Lyz genes; in this case the Cpsf6 gene is upstream of two of the Lyz genes and the Yeats4 gene is downstream of the other two, but these two clusters are separated by about 12 Mb of DNA that potentially was inserted into the opossum genome (Figure 4). Of the numerous lysozyme genes found in Xenopus tropicalis, the LyzA gene was found to be most closely related to the avian and mammalian Lyz genes in the phylogenetic trees (Figure 3); consistent with this, the LyzA gene is located adjacent to a Yeats4 ortholog (Figure 4), suggesting it is a true ortholog of the mammalian Lyz gene. Most fish Lyz sequences branch in a clade with mammalian Lyz genes (Figure 3). However, although the Cpsf6 and Yeats4 genes are neighbors in the fish genomes, the fish lysozyme genes are not adjacent to either of these genes; thus the genomic neighborhood analysis does not provide support for the orthology of fish and mammalian Lyz genes. Intriguingly, the zebrafish Lyz gene was found to be in a different genomic context from other fish Lyz genes (results not shown), which would agree with the hypothesis generated by the phylogenetic analyses (see Figure 3, and Additional files 2 and 3: Figures S1 and S2) that the zebrafish Lyz gene is not truly orthologous to the other fish Lyz genes.

The number of Lyz genes found in mammalian genomes varied from 1, in most species, to 12 (Figure 2, and Additional file 1: Table S1). Multiple Lyz genes had previously been identified in the genomes of mice and rats [26-29], rabbits [30,31], and artiodactyls [32-36]. In addition to these species, multiple Lyz genes were found in many other species, including guinea pig (9 genes), elephant (8 genes), armadillo (5 genes), sloth (6 genes), opossum (4 genes), and wallaby (8 genes) (Figure 2, and Additional file 1: Table S1). Phylogenetic analysis of the Lyz sequences, suggested that the multiple genes in diverse species are due to independent gene duplications or amplification events (Figure 5). However, previous work has shown that Lyz genes have been subjected to
or both (Additional file 5: Figure S4). Interestingly, although an ortholog of Lyz1/2 was found in the platypus genome, one could not be found in the opossum, even though the Bambi and Dnm1p17 genes are adjacent in this species; this suggests that the Lyz1/2 gene was deleted from the opossum genome (Figure 2, and Additional file 5: Figure S4). Both human and macaque have duplicated Lyz1/2 genes that reside in similar genomic neighborhoods (Additional file 5: Figure S4), which should imply that the Lyz1/2 gene duplication occurred prior to the human-macaque divergence. Phylogenetic analysis of the Lyz1/2 sequences, however, implies that independent gene duplication events occurred on the macaque, human, and marmoset lineages (Figure 6). A more likely scenario than multiple independent gene duplications in these closely-related primate species is that the original Lyz1/2 gene duplication event occurred in their common ancestor (as diagrammed in Figure 2), and concerted evolution between the Lyz1 and Lyz2 genes has obscured this original event. Although a Lyz1/2 pseudogene was found in the treeshrew (Figure 2, and Additional file 1: Table S1), it was generated independently, as discussed above. Potential Lyz1/2 orthologs in zebrafish and lizard were suggested by phylogenetic analysis (Figure 3); however, concerted evolution on the ruminant and rodent lineages [28,29,32-37]. Since the inference of independent gene duplication on sister lineages, instead of on their common ancestral lineage, is a pattern generated by concerted evolution, the distributions of duplicated Lyz genes (Figure 2, and Additional file 1: Table S1) combined with the phylogenetic analysis (Figure 5) suggests that concerted evolution might also have occurred on lineages such as Afrotheria (e.g., elephant and hyrax) and marsupials (opossum and wallaby).

Lysozyme-like 1/2 (Lyz1/2) Genes

As mentioned above, the Lyz1 and Lyz2 genes in the human genome appear to have been generated Recently via a genomic segmental duplication. Indeed, with the exception of those primate species that are close relatives of human, most other mammals have only a single gene sequence similar to Lyz1 or Lyz2 (Figure 2, and Additional file 1: Table S1). Comparison of the genomic neighborhoods surrounding the Lyz1/2 genes in diverse mammals demonstrates orthology of these genes, as the genes adjacent to them are either Bambi or Dnm1p17,

Figure 5 Phylogeny of mammalian lysozyme-like 1 (Lyz) genes. A Bayesian phylogenetic tree of mammalian lysozyme-like 1 genes was generated by MrBayes [60,61] using the DNA coding sequences of mammalian Lyz sequences. This tree was built with nst = 2 and rates = gamma as selected by ModelTest [66-68]. The tree was rooted with the platypus Lyz sequence. The posterior probability support for each node is shown.
these genes were not in genomic neighborhoods similar to those of the mammalian genes (results not shown), so their evolutionary relationships remain ambiguous.

**Lysozyme-like 4 (Lyzl4) Genes**
The Lyzl4 gene was either found as a single copy, or was missing, in all of the mammalian genomes examined (Figure 2, and Additional file 1: Table S1). Many of the missing genes likely reflect incomplete genomes rather than deletions. Genomic neighborhood analysis confirmed the orthology of the Lyzl4 genes across placental mammals. However, the flanking genes are on different chromosomes in opossum, suggesting that chromosomal recombination had occurred (Additional file 6: Figure S5). Phylogenetic analyses of Lyzl4 sequences were consistent with it being a single copy gene (Additional file 7: Figure S6). The only potential non-mammalian orthologs of Lyzl4 identified by the phylogenetic analyses were from the lizard (LyzM and LyzN in Figure 3, but no evidence for orthology in Additional files 2-4: Figures S1-S3); however, the lizard and mammalian genes were not in similar genomic neighborhoods (data not shown).

**Lysozyme-like 6 (Lyzl6) Genes**
Most mammals exhibited only one Lyzl6 gene, although the opossum had none. Yet, in contrast to most of the other paralogs, great variation in the number of Lyzl6 genes was observed across mammals, with five genes identified in the dog and four genes identified in both the alpaca and the hyrax (Figure 2, and Additional file 1: Table S1). Given the distant relationships of these three species, these gene duplication events must have occurred independently. The placental and wallaby Lyzl6 genes reside in a conserved genomic neighborhood, which again suggests that this gene was deleted on the opossum lineage (Additional file 8: Figure S7). The presence of multiple Lyzl6 genes in a genome raises the possibility of concerted evolution; however, sufficient data were not available to allow examination of this possibility for the Lyzl6 genes (Figure 2, and Additional files 1 and 9: Table S1 and Figure S8). The phylogenetic analyses did not suggest any candidates for Lyzl6 orthologs in non-mammalian species (Figure 3, and Additional files 2-4: Figures S1-S3).

**Sperm acrosomal protein 3 (Spaca3) Genes**
Spaca3, like Lyzl4, was not found to be duplicated in any of the mammalian genomes examined (Figure 2, and Additional file 1: Table S1). Spaca3 resides in a conserved genomic neighborhood in placental mammals; however, a Spaca3 gene is absent from this genomic neighborhood in the opossum (Additional file 10: Figure S9). While the wallaby and platypus Spaca3 genes could not be placed in a genomic context due to the short lengths of their genomic contigs (Additional file 10: Figure S9), phylogenetic analysis of these sequences (Additional file 11: Figure S110 was consistent with them being orthologs. These results suggest that the Spaca3 gene was deleted on the opossum lineage. Again, no non-mammalian orthologs were suggested by phylogenetic analysis (Figure 3, and Additional files 2-4: Figures S1-S3).

**Sperm acrosomal protein 5 (Spaca5) Genes**
The Spaca5 gene was found only within placental mammals, with no orthologs suggested by phylogenetic analysis or similarity searches in marsupials, platypus, or other vertebrates (Figures 2 and 3, and Additional files 1-4: Table S1 and Figures S1-S3). Thus, it is possible that this gene duplication happened in the ancestor of placental mammals. Genomic neighborhood analysis showed that the Spaca5 gene was in a similar neighborhood on the human, macaque, mouse, and dog X chromosomes (Additional file 12: Figure S11); this genomic region was not found in marsupials, platypus, or other vertebrates (results not shown). The SPACA5 gene was found to be uniquely duplicated in the human genome (Figure 2, Additional files 1 and 13: Table S1 and Figure S12). A very recent duplication of SPACA5, since human-chimpanzee divergence, could account for the perfect identity of the protein sequences (Table 2) without requiring concerted evolution; however, concerted evolution between the human SPACA5 and SPACA5B genes cannot be excluded.

**Lysozyme-like 8 (Lyzl8) Gene**
The platypus genome contained one lysozyme-like gene, named Lyzl8, which did not group with any of the other mammalian genes (Figure 3, and Additional files 2-4: Figures S1-S3). All of our phylogenetic analyses supported the designation of Lyzl8 as a unique lysozyme-like gene duplicate, as the platypus gene did not fall within any of the other monophyletic gene groups. The relationship of the platypus Lyzl8 gene to the other lysozyme-like genes was highly labile in the phylogenetic analyses (Figure 3, and Additional files 2 and 3: Figures S1 and S2). This result is in accord with the fact that the platypus Lyzl8 gene (or protein) showed little similarity to any of the other lysozyme-like genes (or proteins) in our BLAST searches. When the platypus Lyzl8 gene was used as a query to search mammalian genomes, only one genomic sequence – from the sloth (Figure 2, and Additional file 1: Table S1) – was found to have greater similarity to Lyzl8 than to any other lysozyme-like gene. When the short sloth sequence was used as a query against the platypus genome, its best match was the Lyzl8 gene. However, the sloth genomic contig was short, containing only a single exon, and
Therefore could not be used for phylogenetic or genomic neighborhood analysis; thus, the evidence supporting orthology of the sloth sequence to the platypus *Lyzl8* gene is very weak. Thus, at present, it is not clear whether this gene duplication happened on the ancestral mammal lineage, with subsequent losses on most descendant lineages, or on the monotreme lineage.

**Lactalbumin (Lalba) and Calcium-binding Lysozyme (Lysc1) Genes**

Mammalian *Lalba* genes have been well characterized, and are typically single copy in mammals [1,2,4,6,7] (Additional file 14: Figure S13). Curiously, it was previously reported that multiple *Lalba* genes exist in the bovine and ovine genomes [38,39], but here we found only a single copy of the *Lalba* gene in the cow genome (Figure 2, and Additional file 1: Table S1). Whether this reflects differences in the sources of DNA in the different studies or is due to incomplete genome assembly is unknown. The only genome that revealed a duplicate *Lalba* gene was the pika (Figure 2, and Additional file 1: Table S1). Despite hypotheses about an early origin of the *Lalba* gene [10-12], no good candidates for non-mammalian orthologs were identified by our phylogenetic (Figure 3, and Additional file 2-4: Figures S1-S3) or genomic neighborhood (Figure 7, and results not shown) analyses.

An intriguing observation from our genomic neighborhood analysis of was that the mammalian calcium-binding lysozyme gene (*Lysc1*) is located adjacent to the *Lalba* gene in the dog (Figure 7) and horse (not shown) genomes. Both previous phylogenetic analyses [9-12] and our new phylogenetic analyses (Figure 2, and Additional files 2-4: Figures S1-S3) suggested that the *Lysc1* gene originated prior to the radiation of mammals. However, our *tBLASTn* searches using either dog or horse *Lysc1* identified similar sequences in the genomes of only a few diverse mammals – dog, cat, horse, shrew, sloth, and mouse lemur (Figure 2, and Additional file 1: Table S1). It is also noteworthy that the mammalian (*Lysc1*) and avian calcium-binding lysozyme genes are not closely related in our phylogenies, a finding in agreement with some earlier analyses [3,11]. Thus, it is reasonable to speculate that calcium binding evolved independently in these bird and mammal lysozymes. The newly identified *Lysc1*-like genomic sequences all were found on short genomic contigs (Additional file 1: Table S1); nonetheless, both the cat and mouse lemur genomic contigs also encode part of the *c12orf41* gene (Additional file 15: Figure S14A), which is adjacent to the *Lysc1* gene in both the dog and horse genomes (Figure 7). This suggests that the *Lysc1* gene may be near the *c12orf41* gene in other mammalian genomes. Using a strategy that has previously worked to identify genes that could not be found through typical *BLAST* searches [40,41], we focused carefully on the sequences between the *Lalba* and *c12orf41* genes. In 17 of the 37 mammalian genomes available from *Ensembl* [16,25], the *Lalba* and *c12orf41* genes were contained in contiguous genomic sequences. In 18 of the 20 species this genomic region was fragmented into several small genomic contigs; thus, we cannot exclude the possibility that in these genomes the two genes are contiguous. In the pig and the little brown bat this genomic region was not fragmented. In the pig, the current genome assembly does not encode the *Lalba* gene and the *c12orf41* gene is embedded within a very large genomic fragment, suggesting that the *Lalba* - *c12orf41* genomic region has been reorganized in the pig genome (or that this region...
has been incorrectly assembled). In the little brown bat, the *Lalba* gene is embedded in a large genomic fragment that was not annotated to include c12orf41 (although our BLAST searches did identify a very small fragment with strong similarity). A more careful examination of the little brown bat genomic contig revealed that most of the genomic region is composed of unsequenced gaps.

For the 17 genomes that did have linked *Lalba* and c12orf41 genes, the distance between these two genes ranged from ~50 kb (mouse, rat, and rabbit) to ~250 kb (cow and opossum). For all of these genomes, except the opossum (see below), the only genes (or pseudogenes) annotated as existing between *Lalba* and c12orf41 were olfactory receptor-like genes, which are not very useful for identifying orthologous and conserved genomic neighborhoods due to their abundance. In the opossum, in addition to the olfactory receptor-like genes, three additional genes were annotated between *Lalba* and c12orf41: the genes *Mip*, *Spryd4*, and *Gls2*. Unfortunately, the wallaby genome is poorly assembled near the *Lalba* and c12orf41 genes, and thus the neighboring genes could not be identified. Although the *Mip*, *Spryd4*, and *Gls2* genes reside on the same chromosome as *Lalba* and c12orf41 in many mammals (e.g., human, rat, guinea pig, cow, horse, and elephant), they are found greater than 8 Mb away; furthermore, in some species (e.g., mouse and dog) they are on different chromosomes. These observations suggest that the organization of the *Lalba*, c12orf41, *Mip*, *Spryd4*, and *Gls2* genes, and potentially a *Lysc1* gene, has changed between the opossum (and possibly other marsupials) and placental mammals.

The genomic sequence between the *Lalba* and c12orf41 genes for the 17 genomes where these two genes were linked was aligned with *MultiPipMaker* [42,43]. Sequences with similarity to the *Lysc1* gene were not observed in 9 of the genomic sequences – those from marmoset, mouse, rat, guinea pig, rabbit, treeshrew, cow, little brown bat, and opossum (Additional file 15: Figure S14B). It should be noted, however, that for 3 of these species (cow, little brown bat, and marmoset) these genomic sequences contain large amounts of unknown sequence (i.e., sequence gaps). Thus, there are only 6 species with nearly complete genomic sequences spanning the *Lalba* and c12orf41 genes for which we have good evidence for the actual absence of a *Lysc1* gene or pseudogene – mouse, rat, guinea pig, rabbit, treeshrew, and opossum. Pairwise sequence alignments between the mouse, rat, or guinea pig genomic sequences with those from dog or horse (or primates) using *PipMaker* [42] revealed that a large genomic region, which could potentially encode a *Lysc1* gene, is missing from these rodent genomes (results not shown). This suggests that this genomic region, including the *Lysc1* gene, was deleted either early on the rodent lineage or in the common ancestor of rodents and close relatives (e.g., rabbit), but after the divergence of the rodent lineage from the primate lineage (see Figure 2).

Interestingly, some of the genomes – including those from certain haplorrhine primate species (human, chimpanzee, gorilla, orangutan, macaque, baboon, and tarsier) and the elephant – do possess sequences between the *Lalba* and c12orf41 genes that aligned with three of the four exons (exons 2 through 4) of the horse and dog *Lysc1* genes (Figures 7 and 8, and Additional files 15 and 16: Figures S14C and S15). All of these genomic sequences, except for that of the tarsier, are on large genomic segments that have only a few short unsequenced gaps. It is unlikely that all of these genomic sequences have been similarly misassembled, thus we conclude that exon 1 was deleted from all of these genes, and therefore the *Lysc1* gene is a pseudogene in all of these species (Figure 2). In addition to missing exon 1, all of these *Lysc1*-like gene sequences have both frameshift insertions and/or deletions and in-frame stop codons, strengthening the conclusion that they are pseudogenes (Figure 8, and Additional file 16: Figure S15). The loss of exon 1 from the *Lysc1* gene of haplorrhine primates and elephant must have been independent events (Figure 2) as the mouse lemur, a strepsirrhine primate, has a *Lysc1* gene that has an intact exon 1 (Figure 8, and Additional file 16: Figure S15), plus the primates and elephants are quite distant relatives.

Intact *Lysc1* genes that predict potentially functional calcium-binding lysozymes were found in only a few species (dog, horse, and shrew), whereas pseudogenes were found on several lineages (primates, elephant, and sloth). Phylogenetic and genomic analyses suggested that the pair of *Lysc1* genes found in the shrew resulted from a tandem gene duplication event on the lineage leading to this species (Additional files 1 and 17: Table S1 and Figure S16); the divergence of the predicted protein sequences of the two genes suggests that they are not undergoing concerted evolution, however. The *Lysc1* gene was deleted from the genome on the lineages leading to rodents (mouse, rat, guinea pig, and squirrel) and treeshrews. Taken together, the above observations suggest that the *Lysc1* gene likely arose from a duplication of the lactalbumin gene early in mammalian evolution, and was inactivated several times independently, as summarized in Figure 2.

**Conclusions**

Here we have shown that the mammalian lysozyme gene family is much larger than previously anticipated, and is composed of at least eight distantly-related members.
features associated with development of new mammal-specific gene families for keratin-associated proteins [44], kallikreins [45,46], and bitter taste receptors [47]. Amplification of these latter genes has been suggested to be associated with development of new mammal-specific features – e.g., hair (keratin-associated proteins), skin (kallikreins), and diet (bitter taste receptors) [44-47]. Intriguingly, lactalbumin is essential for lactose synthesis in mammary glands, a mammal-specific trait [2,4,7]. These observations raise the possibility that other members of the lysozyme-like family have also evolved mammal-specific roles. The new lysozyme-like genes have been largely conserved within mammals, suggesting that they provide important biological functions. The products of the Spaca3 and Lyz4 genes have recently been shown to be involved in fertilization in mice [18,19]. Much further study is needed to identify the enzymatic activities (if any) and biological functions of these newly identified lysozyme-like proteins.

Similar to the keratin-associated protein [44] and bitter taste receptor [47] gene families, genes for the lysozyme-like proteins are dispersed over several chromosomes (Table 1). The mechanisms by which these original gene duplications occurred are unclear, as the genes that flank the dispersed lysozyme-like genes show no homology to each other, implying that they were not generated by large segmental duplication events (as we observed for the duplications of LYZL1/LYZL2 and Spacas/Spaca5B in the human genome). The lysozyme-like gene family also shares with the keratin associated protein [44], kallikrein [45], and bitter taste receptor [47] gene families the propensity for lineage-specific gene duplications (see Figures 2 and 3). The lineage-specific expansions, in contrast to the initial tandem duplications, have frequently been tandem in nature. Such tandem organization increases the likelihood that the duplicated genes could be involved in concerted evolution [22,23], which our phylogenetic analyses suggest have occurred in the Lyz and Lyz1/2 subfamilies. The Lyz subfamily showed the greatest tendency to tandemly duplicate and evolve in concert, whereas the other lysozyme-like genes typically showed conservation in copy number. Tandem duplication or amplification of the Lyz gene has previously been observed in certain mammals, including the ruminants and rodents, where lysozyme appears to function as a digestive enzyme in the gut [3,25-34]. It is of interest to note that many of the species that we found to possess multiple Lyz genes – e.g., elephant and wallaby – are also herbivorous species, and thus may use lysozyme as a digestive enzyme upon gut bacteria. The need for higher levels of digestive lysozymes in the guts of fermenting herbivores could have driven the fixation of the tandem duplications in these lineages. Gene conversion between the tandem duplicates might provide a mechanism whereby favorable mutations in one gene copy could spread to the other copies in the cluster [33,36], as well as a mechanism for retention of sequence similarity [24] in well-adapted proteins.

| Signal peptide | Exon 1 | Exon 2 | Exon 3 | Exon 4 |
|----------------|--------|--------|--------|--------|
| Horse          | Cavalo|        |        |        |
| Dog            | ...    | ...    | ...    | ...    |
| Cat            | ...    | ...    | ...    | ...    |
| Human          | ...    | ...    | ...    | ...    |
| Chimpanzee     |        |        |        |        |
| Gorilla        |        |        |        |        |
| orangutan      |        |        |        |        |
| Babi           |        |        |        |        |
| Tarsier        |        |        |        |        |
| Mouse lemur    |        |        |        |        |
| Elephant       |        |        |        |        |
| Sloth          |        |        |        |        |

**Figure 8** Alignment of predicted calcium-binding lysozymes (Lyz1). Inferred amino acid sequences of predicted Lyz1 genes for diverse mammals are shown in the single-letter amino acid code. The DNA sequences are shown in Additional file 16: Figure S15. The diverse mammals are shown in the single-letter amino acid code.
Methods

Database Searches

All vertebrate genomes maintained in the Ensembl [16] and PreEnsembl [25] databases (release 57, see Additional file 1: Table S1 for a full list) were searched in April 2010 for lysozyme-like sequences. We initially searched the genomes using the tBLASTn algorithm [20,48] using previously-characterized human and rodent lysozyme c and lactalbumin sequences. Subsequent tBLASTn searches used all of the identified putative lysozyme-like protein sequences. Similar searches were conducted using additional databases (e.g., genome assemblies and ESTs) available at the NCBI website [49]. After identification of the dog and horse calcium-binding lysozyme gene, the other mammalian genome assemblies on the Ensembl database were searched using these sequences using tBLASTn for similar sequences. All sequences that had E-scores below 0.01 were examined. Sequences identified by BLAST searches were used in reciprocal BLASTx searches of the human, mouse and dog proteomes to ensure that their best matches were lysozyme-like sequences. Sequences that were unannotated to encode lysozyme-like sequences (see Additional file 1: Table S1) were then used in tBLASTn [20] searches of the Ensembl [16] and NCBI databases [49] for related sequences. Several insect sequences were downloaded to represent the diversity of insect lysozyme sequences.

Genomic comparisons of DNA sequences near the lysozyme-like genes were conducted using PipMaker and MultiPipMaker [42,43,55]. Genes neighboring the lysozyme-like genes were identified from the genome assemblies at Ensembl [16] and PreEnsembl [25]. The organization of genes adjacent to the lysozyme-like genes was used to determine whether the genes of interest reside in conserved genomic neighborhoods.

Phylogenetic Analysis

Phylogenies of vertebrate lysozyme-like gene coding sequences were generated with sequences from human, mouse, dog, horse, opossum, wallaby, and platypus, representing the diversity of mammals, as well as those from other vertebrate species (see Additional file 1: Table S1) and outgroups (Additional file 18: Table S2). Lysozyme-like coding sequences were aligned using MAFFT [56] and Clustal [57], as implemented at the Guidance web site [58,59], using default parameters. (A MAFFT alignment of all the full-length sequences is provided in Additional file 19: Figure S17). Protein sequences were used as guides to generate the DNA sequence alignments. The reliability of the alignments was examined using Guidance [58,59] and trimmed alignments using sites that had values above the default cut-off of 0.93 were generated. Insect and/or amphioxus lysozyme sequences were used to root the trees of vertebrate lysozyme-like sequences.

Phylogenetic trees of the sequences were generated by a variety of methods including MrBayes 3.1.2 [60,61], PhyloBayes 3.2f [62], and PhyML [63], MEGA4.0.2 [64], and PAUP* 4beta10 [65]. Bayesian trees were generated from coding sequences with MrBayes 3.1.2 using parameters selected by hierarchical likelihood ratio tests with ModelTest version 3.8, as implemented on the ModelTest server [66-68]. MrBayes was run for 2,000,000 generations with four simultaneous Metropolis-coupled Monte Carlo Markov chains sampled every 100 generations. The average standard deviation of split frequencies dropped to less than 0.02 for all analyses. The first 25% of the trees were discarded as burn-in with the remaining samples used to generate the consensus trees. Trace files generated by MrBayes were examined by Tracer [69] to verify if they had converged. Bayesian phylogenies were also generated from protein sequences using PhyloBayes, with two chains being used with the automatic stopping rule set to terminate the analysis when hpcomp and tracecomp indicated that discrepancies between the chains was equal to or below 0.2 and all effective sizes were greater than 100. The first 10% of the trees were discarded as burnin. PAUP* was used to construct parsimony trees. Bootstrapped maximum likelihood trees, 100 replications, were generated by PhyML [63] on the PhyML webserver [70] using parameters for the substitution model suggested by ModelTest. The maximum likelihood search was initiated from a tree generated by BI0NJ and the best tree was identified after heuristic searches using the nearest neighbor interchange (NNI) algorithm. MEGA4 [64] was used to construct bootstrapped (1000 replications) neighbor-joining distance trees, using either Maximum Composite Likelihood distances for the DNA sequences or JTT distances for the proteins sequences. Bootstrapped parsimony trees were also generated by PAUP [65], with 1000 replications and the same search method used for maximum likelihood.

With respect to orthology-paralogy issues, the choice of outgroup, the alignment method (MAFFT [56] or Clustal [57]), and the use of full-length or trimmed (based on Guidance scores [58,59]) alignments had little influence on the key findings of these analyses. Methods that relied on shorter sequences (i.e., trimmed
alignments or protein sequences) or simpler models of sequence evolution (i.e., neighbor-joining or parsimony) tended to yield weaker support for the earlier diverging lineages, but none of our analyses were in significant conflict with the key inferences of the phylogeny presented in Figure 3.

For phylogenies that contained just mammalian lysozyme-like sequences, Lalba sequences were arbitrarily used to root the trees. When only mammalian lysozyme-like gene sequences were used for the phylogenetic analyses, then stronger support for each of the orthologous groups was found with all of the phylogenetic methods used including Bayesian inference, maximum likelihood, distance, and parsimony (see Additional file 4: Figure S3). To generate gene-specific phylogenies, the platypus sequence was used as a root, except for Lysc1 and Spaca5 where the platypus does not have these sequences. For Lysc1, the sloth sequence was used to root the tree, whereas for Spaca5 the elephant and tenrec sequences provided the root.

**Additional material**

- **Additional file 1:** Supplementary Table 1. This file is in PDF format. Location of lysozyme genes in vertebrate genomes.
- **Additional file 2:** Supplementary Figure 1. This file is in PDF format. Phylogeny of vertebrate lysozyme-like sequences generated by PhyloBayes.
- **Additional file 3:** Supplementary Figure 2. This file is in PDF format. Phylogeny of vertebrate lysozyme-like sequences generated by PhyML.
- **Additional file 4:** Supplementary Figure 3. This file is in PDF format. Phylogeny of only mammalian lysozyme-like sequences generated by MrBayes with support for the orthologous genes by different phylogenetic methods.
- **Additional file 5:** Supplementary Figure 4. This file is in PDF format. Conservation of genomic organization near Lysl2 genes.
- **Additional file 6:** Supplementary Figure 5. This file is in PDF format. Conservation of genomic organization near Lys4H genes.
- **Additional file 7:** Supplementary Figure 6. This file is in PDF format. Phylogeny of Lys4H genes.
- **Additional file 8:** Supplementary Figure 7. This file is in PDF format. Conservation of genomic organization near Lys26 genes.
- **Additional file 9:** Supplementary Figure 8. This file is in PDF format. Phylogeny of Lys26 genes.
- **Additional file 10:** Supplementary Figure 9. This file is in PDF format. Conservation of genomic organization near Spac3 genes.
- **Additional file 11:** Supplementary Figure 10. This file is in PDF format. Phylogeny of Spac3 genes.
- **Additional file 12:** Supplementary Figure 11. This file is in PDF format. Conservation of genomic organization near Spac5 genes.
- **Additional file 13:** Supplementary Figure 12. This file is in PDF format. Phylogeny of Spac5 genes.
- **Additional file 14:** Supplementary Figure 13. This file is in PDF format. Phylogeny of Lalba genes.
- **Additional file 15:** Supplementary Figure 14. This file is in PDF format. Conservation of genomic sequences between Lalba and c12b4W1 genes.
- **Additional file 16:** Supplementary Figure 15. This file is in PDF format. DNA sequences of Lysc1 genes.

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**Authors’ contributions**

DMI and CBS together designed the research and outlined the manuscript. DMI, JMB, and CBS obtained and analyzed the data. DMI drafted the manuscript. All of the authors have read, edited, and approved the final manuscript.

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