# Comparative studies of adipose triglyceride lipase genes and proteins: an ancient gene in vertebrate evolution 

This article was published in the following Dove Press journal:<br>Open Access Bioinformatics<br>7 March 2012<br>Number of times this article has been viewed

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#### Abstract

At least eight families of mammalian patatin-like phospholipase domain-containing proteins (EC 3.1.1.3) catalyze the hydrolysis of triglycerides, including adipose triglyceride lipase (ATGL), which functions in triglyceride lipase metabolism in the body, especially in adipose tissue. Bioinformatic methods were used to predict the amino acid sequences, secondary and tertiary structures, and gene locations for $A T G L$ genes and encoded proteins using data from several vertebrate genome projects. ATGL genes usually contained nine coding exons for each of the vertebrate genomes examined, whereas the invertebrate sea squirt (Ciona intestinalis) ATGL gene contained a single exon. Vertebrate ATGL subunits contained $473-504$ residues, shared $>46 \%$ sequence identities, and exhibited sequence alignments and identities for key amino acid residues and predicted motifs: an N -terminal lipid binding region (residues $7-29$ for human ATGL); a patatin "motif" (residues $10-179$ ); a putative active site oxyanion "hole" (Cys15, Gly16, and Leu18); and catalytic dyad active site residues (Ser47 and Asp166). Predicted tertiary structures for the ATGL patatin "motif" were similar to those reported for potato patatin, suggesting that this structure is strongly conserved during animal and plant evolution. Human $A T G L$ contained a CpG131 island within the gene promoter; miR-124/506 and miR-108 binding sites within the mRNA $3^{\prime}$-noncoding region; several transcription factor binding sites, including PPARA and PPARG, which are key regulators of genes encoding enzymes of lipid metabolism; and exhibited wide tissue expression at a higher than average level ( $2.2 \times$ ). Phylogenetic analyses of vertebrate $P N P L A$-like gene families suggest that $A T G L$ is an ancient gene in vertebrate evolution which has been derived from an ancestral $A T G L$ gene (encoding adipose triglyceride lipase) and undergone successive gene duplication events, forming ancestral genes for vertebrate PNPLA1, ATGL, PNPLA3, PNPLA4, and PNPLA5 gene families.


Keywords: vertebrates, amino acid sequence, adipose triglyceride lipase, patatin-like phospholipase domain-containing proteins, gene duplication, evolution

## Introduction

At least eight mammalian patatin-like phospholipase domain-containing (PNPLAlike) proteins (EC 3.1.1.3) and genes have been reported which encode patatin motifcontaining lipases. ${ }^{1}$ Human $A T G L$ (also called PNPLA2 or PLPL2) encodes adipose triglyceride lipase (ATGL, also known as desnutrin or PNPLA2) and is localized on chromosome 11 between the genes encoding ribosomal protein P 2 ( $R P L P 2$ ) and EF-hand calcium-binding domain-containing protein 4A (EFCAB4A). ${ }^{2}$ Other human PNPLA-like genes are separately localized on the human genome, including PNPLA1 (or PLPL1, chromosome 6); ${ }^{3}$ PNPLA3 (PLPL3) and PNPLA5 (PLPL5, chromosome 22); ${ }^{4}$

PNPLA4 (PLPL4, X chromosome); ${ }^{5}$ PNPLA6 (PLPL6 or NTE) (neuropathy target esterase, chromosome 19); ${ }^{6,7}$ PNPLA7 (PLPL7, chromosome 9); ${ }^{8}$ and PNPLA8 (PLPL8 or IPLA2G, calcium-independent phospholipase A2-gamma, chromosome 7). ${ }^{9-11}$

ATGL catalyzes the first step in the hydrolysis of triglycerides in adipocytes and lipid droplets in other tissues of the body, and also acts as an acylglycerol transacylase. ${ }^{12,13}$ ATGL plays a key role in regulating the size and turnover of adipocytes and lipid droplets in nonadipose tissues in association with hormone-sensitive lipase. ${ }^{13}$ Hormone-sensitive lipase serves as the major lipase for catecholamine-stimulated and natriuretic peptide-stimulated lipolysis, whereas ATGL mediates the hydrolysis of triglycerides during basal lipolysis. ${ }^{14-16} A T G L$ gene expression has been shown to be upregulated during fasting, ${ }^{17}$ downregulated by insulin, ${ }^{18}$ and is associated with type 2 diabetes ${ }^{2}$ and obesity. ${ }^{19}$ Human ATGL also serves as a likely target for transcriptional activation by peroxisome proliferator-activated receptor gamma, a key adipogenic transcription factor, ${ }^{18}$ and is upregulated in skeletal muscle by exercise training. ${ }^{20}$ In addition, human $A T G L$ gene mutations have been shown to cause a neutral lipid storage condition and severe myopathy. ${ }^{21-23}$ Notari et al ${ }^{24}$ have also identified ATGL in retinal and nerve cell plasma membranes as an extracellular pigment epithelium-derived factor which may contribute to ligand/receptor signaling function in these cells.

ATGL and other members of the PNPLA-like enzymes belong to the patatin family of acyl hydrolases, the proteins of which are characterized by conserved amino acid sequences for their active sites (human ATGL residues identified) involved in catalysis (Gly45-X46-Ser47-X48-Gly49) at their active sites ${ }^{25}$ and a Ser-Asp catalytic dyad (Ser47/Asp166 for human ATGL) instead of the Ser-His-Asp/Glu triad reported for other lipases. ${ }^{26}$ Although three-dimensional structural analyses have not been reported for mammalian ATGL, the crystal structure for human PNPLA8 (also IPLA2G or cytosolic phospholipase A2) has been described ${ }^{27}$ which shows structural similarities to potato patatin. ${ }^{25}$

This paper reports the predicted gene structures and amino acid sequences for $A T G L$ vertebrate genes and proteins, including primate (human [Homo sapiens] and chimpanzee [Pan troglodytes]), other eutherian mammals (mouse [Mus musculus], rat [Rattus norvegicus], guinea pig [Cavia porcellus], pig [Sus scrofa], cow [Bos taurus], dog [Canis familiaris]), panda (Ailuropoda melanoleuca), a marsupial mammal (opossum, Monodelphis domestica) and other vertebrates, including chicken (Gallus gallus),
lizard (Anolis carolensis), frog (Xenopus tropicalis), and zebrafish (Danio rerio), and an invertebrate sea squirt (Ciona intestinalis). Predicted secondary and tertiary structures for vertebrate ATGL protein subunits are also described, as well as the structural, phylogenetic, and evolutionary relationships of these genes and enzymes with other PNPLAlike gene families.

## Materials and methods

## Vertebrate ATGL and other PNPLA-like gene and protein identification

BLAST (Basic Local Alignment Search Tool) studies were undertaken using web tools from the National Center for Biotechnology Information (http://blast.ncbi.nlm.nih.gov/ Blast.cgi). ${ }^{28}$ Protein BLAST analyses used human ${ }^{29}$ and mouse ${ }^{12}$ ATGL and PNPLA-like amino acid sequences deduced from reported sequences for these genes. ${ }^{3-10}$ Nonredundant protein sequence databases for several mammalian and other vertebrate genomes were examined using the BLASTP algorithm with the default scoring parameters. The following genomes were examined: human (H. sapiens), ${ }^{30}$ chimpanzee ( $P$. troglodytes), ${ }^{31}$ horse (Equus caballus), ${ }^{32}$ cow (B. Taurus), ${ }^{33}$ mouse (M. musculus), ${ }^{34}$ rat (R. norvegicus), ${ }^{35} \operatorname{dog}$ (C. familiaris), ${ }^{36}$ opossum (M. domestica), ${ }^{37}$ chicken (G. gallus), ${ }^{38}$ lizard (A. carolensis), ${ }^{39}$ frog (X. tropicalis), ${ }^{40}$ zebrafish (D. rerio), ${ }^{41}$ and sea squirt (C. intestinalis). ${ }^{42}$ This procedure produced multiple BLAST "hits" for each of the protein databases which were individually examined and retained in FASTA format, and a record kept of the sequences for predicted encoded PNPLA-like proteins. These records were derived from annotated genomic sequences using the gene prediction method: GNOMON and predicted sequences with high similarity scores were generated (usually $>800$ with full sequence coverage). Patatin "motif" amino acid sequences were obtained from the following sources: potato POT1, ${ }^{25}$ mouse ATGL, ${ }^{12}$ mouse PNPLA1, ${ }^{34}$ PNPLA3, ${ }^{43}$ rat PNPLA4, ${ }^{35}$ and mouse PNPLA5 ${ }^{34}$ using web tools from the National Center for Biotechnology Information (http://blast. ncbi.nlm.nih.gov/Blast.cgi).

BLAST analyses were subsequently undertaken for each of the predicted ATGL and other PNPLA-like amino acid sequences using the UC Santa Cruz web browser (http:// genome.ucsc.edu/cgi-bin/hgBlat) ${ }^{44}$ with default settings to obtain the predicted locations for each of the vertebrate PNPLA-like genes, including predicted exon boundary locations and gene sizes (see Table 1). Structures for human $A T G L$ isoforms were obtained using the AceView website to
Table I Vertebrate ATGL and other PNPLA-like lipase genes and proteins

| Vertebrate | Species | PNPLA gene (other name) | Chromosome coordinates | Gene size | Exons strand | Subunit MW | Amino acids | pl | GenBank ID | UNIPROT ID | 'NCBI reference ID ${ }^{2}$ NCBI predicted ID |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Human | Homo sapiens | ATGL (PNPLA2) | 11:819,719-824,859 | 5,141 | $9+\mathrm{ve}$ | 55,316 | 504 | 6.7 | BCOII958 | Q96AD5 | 'NM_023376 |
| Chimpanzee | Pan troglodytes | ATGL (PNPLA2) | \| 1:879,389-884,735 | 5,347 | $9+\mathrm{ve}$ | 55,429 | 504 | 6.9 | na | na | ${ }^{2} \mathrm{XP}$ _003339482.1 |
| Mouse | Mus musculus | Atgl (Pnpla2) | 7:148,641,186-148,645,564 | 4,379 | $9+\mathrm{ve}$ | 53,657 | 486 | 6.1 | BC064747 | Q8BJ56 | 'NR_028142 |
| Rat | Rattus norvegicus | Atgl (Pnpla2) | I:201,642,058-201,646,343 | 4,286 | $9+\mathrm{ve}$ | 52,567 | 478 | 6.2 | ACI09542 | P0C548 | 'NM_001108509.2. |
| Guinea pig | Cavia porcellus | ATGL (PNPLA2) | ${ }^{3} 218: 129,068-132,994$ | 3,927 | $9-\mathrm{ve}$ | 53,964 | 486 | 6.3 | ${ }^{5}$ ENSCPOT5922 | na | na |
| Pig | Sus scrofa | ATGL (PNPLA2) | 2:149,082-153,354 | 4,273 | $9-\mathrm{ve}$ | 53,147 | 486 | 7.5 | ABW06598 | A5JTI5 | 'NM_001098605.1 |
| Cow | Bos taurus | ATGL (PNPLA2) | ${ }^{4}$ Un.004.171:137,584-141,740 | 4,157 | $9-\mathrm{ve}$ | 53,418 | 486 | 6.8 | BCII2803 | Q2KII 8 | 'NM_001046005.1 |
| Panda | Ailuropoda melanoleuca | ATGL (PNPLA2) | ${ }^{3} \mathrm{GLI} 95194.1: 5,234-9,734$ | 4,501 | $9-\mathrm{ve}$ | 52,834 | 482 | 8.1 | ${ }^{6}$ ENSAMET7602 | na | na |
| Opossum | Monodel/phis domestica | ATGL (PNPLA2) | ${ }^{4}$ Un:45,368,040-45,372,831 | 4,792 | $9+\mathrm{ve}$ | 53,547 | 490 | 6.8 | na | na | ${ }^{2} \mathrm{XP}$ _001380646.2 |
| Chick | Gallus gallus | ATGL (PNPLA2) | 5:16,838,493-16,868,610 | 30,118 | $9-\mathrm{ve}$ | 53,610 | 483 | 6.7 | EU419874 | A8WEN5 | 'NM_001113291.1 |
| Lizard | Anolis carolensis | ATGL (PNPLA2) | 1:73,687,462-73,733,079 | 45,618 | $9-\mathrm{ve}$ | 53,795 | 483 | 6.4 | na | na | ${ }^{2} \mathrm{XP}$-003214822.1 |
| Frog | Xenopus tropicalis | ATGL (PNPLA2) | ${ }^{39} 90: 125,824-140,107$ | 14,284 | $9+\mathrm{ve}$ | 55,162 | 498 | 5.9 | BC089233 | Q5FWR9 | 'NM_001015693.1 |
| Zebrafish | Danio rerio | ATGL (PNPLA2) | ${ }^{3} 3512: 30,775-45,374$ | 14,600 | $10+\mathrm{ve}$ | 52,253 | 473 | 6.8 | BC075928 | na | 'NM_001002338.1 |
| Human | Homo sapiens | PNPLAI (PLPLI) | 6:36,238,237-36,275,490 | 37,254 | $8+\mathrm{ve}$ | 57,876 | 532 | 8.2 | BCI03906 | Q8N8W4 | 'NM_001145716.1 |
| Mouse | Mus musculus | Pnplal (P\|plI) | 17:28,995,812-29,023,893 | 28,082 | $9+\mathrm{ve}$ | 65,171 | 592 | 8.6 | AK132521 | Q3VID5 | 'NM_00I034885.3. |
| Rat | Rattus norvegicus | Pnpla3 (P/p/3) | 7:122,152,145-\|22,171,991 | 19,847 | $9+\mathrm{ve}$ | 45,908 | 414 | 6.8 | EDMI5609 | na | na |
| Mouse | Mus musculus | Pnpla3 (Plp/3) | 15:83,998,304-84,016,512 | 18,209 | $9+\mathrm{ve}$ | 45,772 | 413 | 6.6 | BC028792 | Q91WW7 | 'NM_054088.3 |
| Human | Homo sapiens | PNPLA4 (PLPL4) | X:7,866,804-7,895,475 | 29,493 | 6 -ve | 27,980 | 253 | 9.0 | BC020746 | P41247 | 'NM_001142389.1 |
| Rat | Rattus norvegicus | Pnpla4 (P\|p|4) | X:64,019,691-64,022,515 | 2,825 | 6 -ve | 27,439 | 252 | 9.1 | FQ216301 | na | ${ }^{2}$ XP_343791.1 |
| Cow | Bos taurus | PNPLA4 (PLPL4) | Un.004.9:49,923-92,708 | 42,786 | 6 -ve | 28,117 | 253 | 9.7 | BT021623 | na | ${ }^{2}$ XP_590366.2 |
| Human | Homo sapiens | PNPLA5 (PLPL5) | 22:44,276,678-44,287,760 | 12,299 | $9-\mathrm{ve}$ | 47,912 | 429 | 6.3 | BC031820 | Q7Z6Z6 | 'NM_138814 |
| Rat | Rattus norvegicus | Pnpla5 (P\|p|5) | 7:122,105,840-122,115,990 | 10,151 | $9-\mathrm{ve}$ | 50,408 | 453 | 8.5 | na | D3ZXU1 | 'NM_001130497.1 |
| Mouse | Mus musculus | Pnpla5 (P/p/5) | 15:83,943,618-83,953,543 | 9,926 | $9-\mathrm{ve}$ | 48,480 | 432 | 9.0 | BCIO9360 | Q32LZ8 | 'NM_029427.1 |
| Human | Homo sapiens | PNPLA6 (NTE) | 19:7,600,675-7,626,445 | 25,771 | $32+\mathrm{ve}$ | 149,996 | 1,366 | 7.7 | BC050553 | Q8IYI7 | 'NM_006702 |
| Rat | Rattus norvegicus | Pnpla6 (Nte) | 12:2,610,798-2,638,622 | 27,825 | $34-\mathrm{ve}$ | 148,558 | 1,326 | 8.2 | na | D3ZRF6 | ${ }^{2} \mathrm{XP}$ _001057249.2 |
| Mouse | Mus musculus | Pnpla6 (Nte) | 8:3,516,954-3,544,025 | 27,072 | $33+\mathrm{ve}$ | 149,537 | 1,355 | 8.0 | AFI73829 | Q3TRM4 | 'NM_001122818.1 |
| Human | Homo sapiens | PNPLA7 (PLPL7) | 9:140,354,848-\|40,441,85| | 87,007 | $34-\mathrm{ve}$ | 145,733 | 1,317 | 7.8 | AK697623 | Q6ZV29 | 'NM_001098537.1 |
| Rat | Rattus norvegicus | Pnpla7 (P\|p|7) | 3:5,125,219-5,200,872 | 75,654 | $35+\mathrm{ve}$ | 150,156 | 1,349 | 6.7 | BC083547 | Q5BK26 | 'NM_144738.2 |
| Mouse | Mus musculus | Pnpla7 (P\|p|7) | 2:24,835,554-24,909,310 | 73,757 | $35+\mathrm{ve}$ | 150,494 | 1,352 | 6.5 | BC025621 | A2AJ88 | 'NM_146251.4 |
| Human | Homo sapiens | PNPLA8 (IPLA2G) | 7:108,112,848-108,155,935 | 43,088 | $9-\mathrm{ve}$ | 88,477 | 782 | 9.3 | BC032999 | Q9NP80 | 'NM_015723.2 |
| Rat | Rattus norvegicus | Pnpla8 (Ipla2g) | 6:63,671,683-63,704,337 | 32,655 | $9+\mathrm{ve}$ | 87,961 | 776 | 9.2 | FQ2II866 | na | ${ }^{2}$ EDM03350.1 |
| Mouse | Mus musculus | Pnpla8 (Ipla2g) | 12:45,383,654-45,412,583 | 28,930 | $9+\mathrm{ve}$ | 87,294 | 775 | 9.3 | BC019364 | Q8KINI | 'NM_026164.2 |
| Opossum | Monodelphis domestica | PNPLAI (PLPLI) | 2:275,536,646-275,586,990 | 50,345 | $9+\mathrm{ve}$ | 60,027 | 540 | 6.3 | na | na | ${ }^{2} \mathrm{XP}$ _001378816.2 |
| Sea squirt | Ciona intestinalis | PNPLA2 (ATGL) | ${ }^{3} 127: 40,380-41,927$ | 1,548 | $1+\mathrm{ve}$ | 57,387 | 516 | 8.1 | AKII2234 | na | na |
| zebrafish | Danio rerio | PNPLA7 (PLPL7) | 5:26,715,402-26,738,943 | 23,542 | $34-\mathrm{ve}$ | 147,166 | 1,293 | 6.5 | BC092427 | B0S710 | ${ }^{4}$ BX842571.5 |
| Frog | Xenopus tropicalis | PNPLA7 (PLPL7) | ${ }^{3} 886: 206,377-247,043$ | 40,667 | $32+\mathrm{ve}$ | 148,479 | 1,323 | 7.9 | BCI29023 | F7ASR4 | 'NM_001097275.1 |
| Sea squirt | Ciona intestinalis | PNPLA7 (PLPL7) | 02q:1, 109,384-1,121,040 | 11,657 | $26+\mathrm{ve}$ | 123,881 | 1,097 | 6.4 | AKII6835 | na | ${ }^{2} \mathrm{XP}$-002127361.1 |

Table I (Continued)

| Vertebrate | Species | PNPLA gene <br> (other name) | Chromosome <br> coordinates | Gene <br> size | Exons <br> strand | Subunit <br> MW | Amino <br> acids | pl | GenBank ID | UNIPROT <br> ID |
| :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- | :--- |
| Zebrafish | Danio rerio | PNPLA8 (PLPL8) | $4: 12,692,490-12,707,208$ | 14,719 | $9+\mathrm{ve}$ | 76,937 | 696 | 9.0 | na |  |
| 2NCBI peference ID |  |  |  |  |  |  |  |  |  |  |

 sequences.
Abbreviations: pl, theoretical isoelectric points (the number of coding exons are listed); na, data not available.
examine predicted gene and protein structures to interrogate this database of human mRNA sequences ${ }^{45}$ (http://www.ncbi. nlm.nih.gov/IEB/Research/Acembly/).

## Predicted structures and properties of vertebrate ATGL subunits

Alignments of predicted ATGL amino acid sequences were undertaken using a ClustalW method (http://www.ebi.ac.uk/ Tools/msa/clustalw2/). ${ }^{46}$ Predicted secondary and tertiary structures for vertebrate ATGL subunits were obtained using PSIPRED ${ }^{47}$ and SWISS MODEL web tools, respectively. ${ }^{48,49}$ The reported tertiary structure for potato patatin ${ }^{25}$ served as the reference for the predicted ATGL tertiary structures, with a modeling range of residues 3-173 (pig ATGL), 4-176 (opossum ATGL), 9-176 (zebrafish ATGL), and 13-183 (sea squirt ATGL). Theoretical isoelectric points and molecular weights for vertebrate ATGL and PNPLA-like subunits were obtained using Expasy web tools (http://web.expasy. org/compute_pi/)..$^{50}$ Predicted membrane binding helices for PNPLA-like sequences were obtained using web tools from the Center for Biological Sequence Analysis, Technical University of Denmark (http://www.cbs.dtu.dk/services/ THMMH/). ${ }^{51}$

## Human ATGL gene expression and predicted gene regulation sites

The human genome browser (http://genome.ucsc.edu) ${ }^{44}$ was used to examine GNF Expression Atlas 2 data using various expression chips for the human and mouse $A T G L$ genes (http://biogps.gnf.org). ${ }^{52}$ Predicted CpG islands, microRNA (miRNA) binding sites, and transcription factor binding sites for human $A T G L$ were obtained using the UC Santa Cruz genome browser (http://genome.ucsc.edu). ${ }^{44}$

## Phylogenetic studies and sequence divergence

Alignments of ATGL and other PNPLA-like protein sequences were assembled using BioEdit v.5.0.1 and the default settings. ${ }^{53}$ Alignment-ambiguous regions were excluded prior to phylogenetic analysis, yielding alignments of 241 residues for comparisons of vertebrate PNPLAlike sequences with the sea squirt PNPLA-like sequences (Table 1). Evolutionary distances were calculated using the Kimura option ${ }^{54}$ in TREECON. ${ }^{55}$ Phylogenetic trees were constructed from evolutionary distances using the neighbor-joining method ${ }^{56}$ and "rooted" using the internode at the PNPLA6-PNPLA7-PNPLA8 branch. Tree topology was re-examined by the boot-strap method (100 bootstraps
were applied) of resampling and only values that were highly significant $(\geq 95)$ are shown. ${ }^{57}$

## Results and discussion

## Alignments and biochemical features of vertebrate ATGL amino acid sequences

Amino acid sequence alignments for 13 previously unreported vertebrate and invertebrate ATGL amino acid sequences and an invertebrate (sea squirt, C. intestinalis) ATGL sequence are shown in Figure 1, together with the reported sequence for human ${ }^{29}$ and mouse ${ }^{12}$ ATGL. The vertebrate ATGL sequences exhibited $>46 \%$ identities, suggesting that these protein subunits are products of the same gene family, whereas the sequences for the predicted vertebrate PNPLA1, PNPLA3, PNPLA4, and PNPLA5 subunits were $23 \%-42 \%$ identical with the ATGL sequences, indicating that these are members of distinct but related PNPLA-like gene families (Table 2). The sequences for the vertebrate PNPLA6, PNPLA7, and PNPLA8 subunits examined were even more distantly related, having vertebrate ATGL sequences with identities of $<16 \%$ (Table 2). However, two of these sequences (PNPLA6 and PNPLA7) showed comparatively high sequence identities ( $58 \%-61 \%$ ), suggesting that these are closely related gene families. Amino acid sequences for the eight human PNPLA-like proteins examined contained 473-504 (ATGL), 253 (PNPLA4), 429-532 (PNPLA1, PNPLA3, and PNPLA5), 782 (PNPLA8), and 1317-1366 (PNPLA6 or NTE and PNPLA7) residues (Table 1). Consequently, vertebrate PNPLA4 is the smallest among these PNPLA-like proteins with an average molecular weight of about 28,000, while others exhibited molecular weights which are about two times (ATGL, PNPLA1, PNPLA3, and PNPLA5), three times (PNPLA8), or five times larger (PNPLA6 and PNPLA7) than PNPLA4.

Structural studies for potato patatin ${ }^{25,57,58}$ and sitedirected mutagenesis studies of human PNPLA4 $4^{59,60}$ have enabled identification of key catalytic residues among those aligned for the vertebrate and invertebrate ATGL (Figure 1) and PNPLA-like (mouse PNPLA1, ATGL, PNPLA3, and PNPLA5; and rat PNPLA4) sequences examined (Figure 2). These included two proposed active site motifs for vertebrate ATGL. Active site motif 1 was identified as Cys-Gly-Xaa-Leu (mouse ATGL residues 15,16 , and 18) which may function in ATGL catalysis because it is aligned with the potato patatin motif (residues 37-40 Gly-Gly-Xaa-Lys) which functions as an oxyanion hole and stabilizes the oxyanion formed during triglyceride cleavage. ${ }^{25}$ A second active

ATGL motif was identified as Gly-Xaa-Ser-Yaa-Gly (mouse ATGL residues 45-49), together with active site residues Ser47 and Asp166 which serve as the catalytic dyad during catalysis. ${ }^{25}$ ATGL Ser 428 has been identified as a site subject to site-specific phosphorylation ${ }^{61}$ which is conserved among all vertebrate ATGL sequences examined, with the exception of guinea pig, panda, and zebrafish ATGL. Neutral isoelectric points ( pI ) were observed for each of the vertebrate ATGL subunits examined ( pI values 5.9-8.1), as well as the human, mouse, and rat PNPLA1, PNPLA3, PNPLA5, PNPLA6, and PNPLA7 sequences. This is in contrast with the human, rat, and cow PNPLA4 sequences and the human, mouse, and rat PNPLA8 subunits which exhibited high predicted pI values ( $>9.0$, Table 1).

## Predicted secondary and tertiary structures for vertebrate ATGL subunits

Analyses of predicted secondary structures for ATGL sequences revealed similar $\alpha$-helix and $\beta$-sheet structures for all of the vertebrate subunits examined, particularly near key residues or functional domains (Figure 1). Predicted secondary (Figure 1) and tertiary structures (Figure 3) were very similar to those reported for potato patatin ${ }^{25}$ which have been retained for all of the vertebrate ATGL sequences examined. The predicted vertebrate and invertebrate ATGL tertiary structures (pig, opossum, zebrafish, and sea squirt, Figure 2) are based on partial sequences for this enzyme (residues 6-173, 4-176, 9-176, and 13-183, respectively) which reveal the relative positioning and predicted structures for each of $5 \alpha$-helices and $6 \beta$-sheets. These included the N -terminus $\alpha$-helix (designated as $\alpha 1$ ) which contained a predicted membrane bound helix (residues 7-29, see Supplementary Figure 1) and the proposed first active site motif Cys-Gly-Xaa-Leu (human ATGL residues 15,16 , and 18 ); a second $\alpha$-helix ( $\alpha 2$ ) and $\beta$-sheet ( $\beta 2$ ) which contains the second active site motif Gly-Xaa-Ser-Yaa-Gly (residues $45-49$ for human ATGL); and a $\beta$-sheet ( $\beta 6$ ) which contains Asp166, the second member of the active site dyad of catalytic residues. These structures are proximally located within a putative active site cleft which supports the predicted three-dimensional structure for this enzyme. However, any firm conclusions must await further structural studies. Several conserved serine residues were also observed for the vertebrate and invertebrate ATGL sequences examined (human ATGL Ser117, 130, 145, and 170 residues, Figure 1) which may correspond to residues previously proposed for performing structural roles in potato papatin phospholipase A. ${ }^{25-57}$

--------PHGPEDK--DQAVESAQAEDYSQL--------PGEDHILEHLPARLNEALLEACVEPTDLLTTLSNMLPVR---LATAMMVPYTLPLESALSFTIRLLEWLPDVPEDIRWMKEQTGSICQYLVMRAKRKLGRH 388
(
---------PDGPEDEEEDEAVESARAEDHSQP---------PEEGHILEHLPARLNEALLEACVEPTDLLTTLSNMLPVR---LATAMMVPYTLPLESALSFTIRLLEWLPDVPEDIRWMKEQTGSICQYLVMRAKRKLGRH 390
-------PQEEDAEEAAVTEERTGGED--------------------RILEHLPARLNE $\bar{A} L L E A C V E P K D L M T T L S N M L P V R-$

P-----APEDADAEEGAVAMERTGGKDHLPP---------PREDHILEHLPSRLNE A $\bar{A} L L E A C M E P T D L L T T L S N M L P V R-~$
-------PNPNLSGGPHPGLLHRGHLQP--------PLEDPILEHLPSRLNEALLEACIEPKDLLTTFSNMLPVR-
APAPAPAPEEEEEEEEAAAAAAAGGRGRAAQGELALGRGPVEDHILEHLPARLNEALLEACVEPKDLMTTLSNMLPVR-
P------PGEK--KEEEKEAEDQMEDNTALA---------VVEDHIFEHLPPKLNQALLEACAERRGLLSGISNTLPIR-
G------AGDRNPEEEAEEEEDQLGENPALV--------EAEEHILEHLPPKLNQALLEACAERRGLFTGITNLLPIR---LASTLMLPYTLPLESAVSFTVRLLEWLPDVPEDIRWMKEQTRKVCHYLMRKAKNKLGSH 392
P------ENIANENEASRNSEEPEDDIAAVQ--------AAEDHIFEHLPQRLNQALLEACKEKRSMYLSISNLLPVR---MASALMLPYTLPLESALSFTVRLLEWLPDVPEDIEWIKDQTRKVCRYVLRNAKKQLGNH 391
SN----VKGRRNLTRRLSLTAQRISDDDVEGSDVSYTSSLSDYDIEECFDENCNDYDVIKPLPSPSVFTVVHVYPPDKSPTLQLNFQLPPPVLAALETAFQSSATPWKIPLSHFIGNCKKSLIV RVPLDRTYSIATFLLR 414
LPSRLPEQVELRRVQSLPSVPLSCAAYREALPGWMRNNLSLG-DALAKWEECQRQLLLGLFCTNVAFPPEALRMRAP----ADPAPAPADPASPQHQLAGPAPLLSTPAPEARPVIGALGL 504
LPSRLPEQVELRRVQSLPSVPLSCAAYREALPGWMRNNLSLG-DALAKWEECQRQLLLGLFCTNVAFPPEALRMRAP----ADPAPAPADPASPQHQLARPAPLLSTPAPEARPVIGALGL 504
LPSRMSEQVELRRTQSLPSVPLSCATYSEALPGWMRNNLSLG-GALAKWEACQRQLLLGLFCTNVAFPPDALRMHAP----SGPAPAPADPAP------AGLPPC----------------- 484
LPSRLSEQVELRRAQSLPSVPLSCATYSEALPNWVRNNLSLG-DALAKWEECQRQLLLGLFCTNVAFPPDALRMR------APASPTAADPATPQDP-PGLPPC---------------------- 486
LPSRLSEQVELRRAQSLPSVPLSCATYSEALPNWVRNNLSLG-DALAKWEECQRQLLLGLFCTNVAFPPDALRMR------APASPTATDPATPQDP-SGLPPC------------------ 478
LPSRLSEPEELRRTRSLPSVPLSCASYREALPSWVRSNLLLG-DALAKWEECQRQLLLGLFCTNVAFPPDALRMRTPVDHPTPVDPMAMDPTPAQHP-PGLPPR------------------- 485
LPSRLTEQVELRRSQSLPSVPLSCAAYGGVLPSWMRNSLSLG-DVLAKWEERQRQLLLGLFCTNTAFPPDALRML----------APAGPAPAPSQHP-PGSPPC------------------ 486

LPSRLSEQMELRRAQSLPSVPLSCAAYSDALPSWMRNSLLLG-DALAKWEECQRQLLLGLFCTNVAFPPDTLRMH----------APASPASRLPQHP-ATLPPC------------------- 482

LSARLYYHLELGGPQSLP-ISSA-SPCGEAMPMWMRSNRSLS-DVMLKWEEYQRQLVMGLLCINVDMQASLFPWEGF-----QMKLPPLD-------------------------- 483
LSARLYYHLELRKTQSLP-LPLS-VSYGEALPKWLRSNLSLT-DVMTKWEEYQRQLMLGLFCINVDLQASIFPSEGL-----RMRHPSSD------------------------ 483
ITARLYYHLELRRAQSVT-VPLT-FSSSDCLPRWVRTSTSLM-DMMQKWEEYQRQLMPGLFCINVDMQATLYHHQHP-----KQPSLSMDPDSLIMVDVSKESSNETF-------------- 491

HTASFPDDITWLGDHLRMLVDKLQSYFRCRGSKLVGKIRNMGSDAAQDLTHLTQNMIITVQLLTKALWDRGYPYYTT-----WLAPLWCRSFSGGIPQEDKNILQLV--------------- 516
-I

Figure I Amino acid sequence alignments for vertebrate ATGL sequences.
Notes: See Table I for sources of ATGL sequences. *Identical residues; I or 2 conservative substitutions; I or 2 nonconservative substitutions; patatin refers to predicted motif residues (6-I73); predicted oxyanion hole motif (Cys-Gly-Xaa-Leu); active site motif (Gly-Xaa-Ser-Xaa-Gly); catalytic dyad residues Ser and Asp; predicted helix $\alpha$ I, $\alpha 2$; predicted sheet $\beta 1 \beta 2$; conserved serine residues; conserved Ser428 ( $\overline{\mathrm{P}})$ for reported phosphorylation site; and bold underlined font shows predicted exon junctions. Abbreviations: Hu, human; Ch, chimpanzee; Ma, marmoset; Mo, mouse; Ra, rat; Gp, guinea pig; Pi, pig; Bo, bovine; Pa, panda; Op, opossum; Ck, chicken; Li, lizard; Fr, frog; Zf, zebrafish; Ss, sea squirt.

## Predicted gene locations, exonic structures, and expression for vertebrate ATGL genes

Table 1 summarizes the predicted locations and exonic structures for vertebrate $A T G L$ genes based upon BLAST interrogations of several vertebrate genomes using the sequence for human ${ }^{29}$ and mouse ${ }^{12}$ ATGL and the predicted sequences for other vertebrate and invertebrate (sea squirt) ATGL subunits (Table 1) and the UC Santa Cruz
web browser. ${ }^{44}$ Vertebrate $A T G L$ genes predominantly contained nine coding exons with the predicted exonic start sites in identical or similar positions (Figure 1). An exception was the zebrafish $A T G L$ gene, which contained 10 coding exons, with the second exon for most of the other vertebrate ATGL genes being split into two coding exons. $A T G L$ genes varied in size with all of the mammalian ATGL genes examined containing about 4-5 kilobase pairs, whereas other vertebrate $A T G L$ genes were three
(frog [X. tropicalis] and zebrafish [D. rerio]) to nine (lizard [A. carolensis]) times larger (Table 1). This is mainly due to a major increase in the size of the first intron, particularly for chicken and lizard $A T G L$ genes, for which intron 1 is more than 10 times larger than for the other vertebrate ATGL genes (Supplementary Table 1).

Figure 4 illustrates the comparative predicted structures of premessenger RNA human $A T G L$ gene transcripts (http://www.ncbi.nlm.nih.gov/IEB/Research/Acembly/). ${ }^{45}$ There were nine introns and nine coding exons present for the premessenger mRNA reference (RefSeq) sequence (designated as transcript b [NM] in Figure 4) which contained a CpG131 island in the $5^{\prime}$-noncoding segment corresponding to the gene promoter and a second CpG108 island within the coding exon 6 region. Several transcription factor binding sites were also observed in this region: the aryl hydrocarbon receptor; the estrogen receptor, which regulates cellular responses to estrogens; early growth response 4 receptor, a transcription activator for genes involved in mitogenesis and differentiation; forkhead fox protein 01 (FOX01), a transcription factor which regulates response to oxidative stress; peroxisome proliferator-activated receptors alpha and gamma (PPARG/PPARA) which activate genes encoding enzymes of the peroxisomal $\beta$-oxidation pathway; chicken ovalbumin upstream promoter (COUP); homeobox protein 3 (HMX3), involved in the differentiation of nerve cells; and transcription regulator protein, BACH1 (Supplementary Table 2). These predicted gene regulation sites may contribute to the high level of gene expression ( $2.2 \times$ the expression of the average human gene) and wide tissue expression observed for $A T G L .{ }^{45}$ Elango and $\mathrm{Yi}^{62}$ have recently reported that larger CpG islands are associated with gene promoters showing a broad range of gene expressions which contain more RNA polymerase II binding sites than other promoters. Consequently, the presence of CpG131 and the transcription factor binding sites observed within the $A T G L$ gene may contribute significantly to the broad tissue expression observed for $A T G L$ transcripts, as shown in Figure 5, which presents "heat maps" showing the comparative gene expression for various human and mouse tissues obtained from GNF Expression Atlas data using U133A, GNF1H, and U74A ATGL chips (http://genome.ucsc.edu; http://biogps. gnf.org). ${ }^{52}$ Of particular interest are the very high levels of $A T G L$ gene expression observed in human adipocytes, and in mouse brown fat, adipose tissue, mammary gland, trachea, and epidermis, which are consistent with the primary role for this enzyme in degrading triglycerides within fat cells and other tissues of the body. ${ }^{12,13}$ The reference human $A T G L$
transcript also contained an extended $3^{\prime}$-noncoding segment with two predicted miRNA binding sites (miR-124/506 and miR-377). miRNAs have been reported to function as post-transcriptional regulators that bind to complementary sequences on target messenger RNA transcripts (mRNAs), which result in translational repression or target degradation and gene silencing. ${ }^{63}$

## Sequence identities and phylogeny of vertebrate ATGL and other PNPLA-like lipases

Supplementary Figure 2 shows a UCSC genome browser comparative genomics track depicting evolutionary conservation and alignment of nucleotide sequences for the human $A T G L$ gene, including the intronic, exonic, and untranslated regions of this gene, with the corresponding sequences for eight vertebrate genomes. Extensive conservation was observed among these $A T G L$ genomic sequences for the eutherian mammalian genomes, particularly for the primate species but also for the exonic regions for all eutherian genomes examined. In contrast with the mammalian genomes examined, the other vertebrate genomes examined lacked conserved sequences for exon 1 and the intron regions, which may reflect increased evolutionary divergence for these sequences.

A phylogenetic tree (Figure 6) was constructed from alignments of vertebrate PNPLA-like amino acid sequences with the predicted sea squirt (C. intestinalis) ATGL sequence, as well as sea squirt PNPLA7 and PNPLA8 sequences, serving to "root" the tree. The dendrogram showed clustering into three major groups of vertebrate PNPLA-like sequences: group 1, ATGL (the "apparent" ancestral sequence together with PNPLA1, PNPLA4, PNPLA3, and PNPLA5 sequences; group 2, PNPLA6 and PNPLA7; and group 3, PNPLA8. Group 1 is further divided according to the designation of PNPLA-like gene families, including the vertebrate ATGL sequences, which cluster with the sea squirt ATGL-like sequence. This suggests an ancestral relationship between invertebrate ATGL and vertebrate ATGL, as well as with other members of PNPLA-like group 1 sequences, ie, PNPLA1, PNPLA4, and PNPLA3/PNPLA5. Moreover, this is consistent with $A T G L$ being an ancient gene present throughout vertebrate evolution, as well as within the invertebrate sea squirt genome.

Figure 6 also shows the number of times a clade (sequences common to a node or branch) occurred in the bootstrap analyses with replicate values of 90 or more (which are highly significant) for the 100 replicates
Table 2 Percentage identities for vertebrate ATGL and other PNPLA-like amino acid sequences

| PNPLA-like Gene | Human | Mouse | Opossum | Chicken Zebrafish |  | Sea squirt | Human | Mouse | Human | Mouse | Human | Rat | Human | Mouse | Human | Mouse | Human | Mouse | Human | Mouse |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
|  | ATGL | ATGL | ATGL | ATGL | ATGL |  |  |  |  |  |  |  | 4 PNPLA5 |  |  |  |  |  |  |  |
| Human ATGL | 100 | 86 | 81 | 67 | 46 | 30 | 26 | 27 | 37 | 41 | 35 | 37 | 34 | 34 | 9 | 14 | 14 | 13 | 9 | 7 |
| Mouse ATGL | 86 | 100 | 85 | 68 | 46 | 33 | 25 | 27 | 36 | 40 | 35 | 37 | 34 | 33 | 7 | 9 | 10 | 10 | 9 | 10 |
| Opossum ATGL | 81 | 85 | 100 | 70 | 47 | 30 | 25 | 26 | 37 | 41 | 36 | 37 | 37 | 33 | 13 | 16 | 6 | 13 | 10 | 6 |
| Chicken ATGL | 67 | 68 | 70 | 100 | 46 | 30 | 25 | 26 | 35 | 42 | 34 | 34 | 33 | 34 | 11 | 12 | 7 | 12 | 8 | 8 |
| Zebrafish ATGL | 46 | 46 | 47 | 46 | 100 | 30 | 26 | 23 | 38 | 40 | 39 | 39 | 33 | 33 | 8 | 10 | 8 | 9 | 11 | 8 |
| Sea squirt ATGL | 30 | 33 | 30 | 30 | 30 | 100 | 23 | 21 | 23 | 28 | 35 | 33 | 27 | 26 | 8 | 8 | 8 | 10 | 5 | 4 |
| Human PNPLAI | 26 | 25 | 25 | 25 | 26 | 23 | 100 | 63 | 26 | 25 | 30 | 33 | 25 | 25 | 10 | 10 | 7 | 12 | 3 | 3 |
| Mouse PNPLAI | 27 | 27 | 26 | 26 | 23 | 21 | 63 | 100 | 25 | 26 | 30 | 32 | 26 | 24 | 8 | 6 | 5 | 6 | 5 | 5 |
| Human PNPLA3 | 37 | 36 | 37 | 35 | 38 | 23 | 26 | 25 | 100 | 65 | 32 | 32 | 34 | 32 | 10 | 9 | 6 | 9 | 5 | 6 |
| Mouse PNPLA3 | 41 | 40 | 41 | 42 | 40 | 28 | 25 | 26 | 65 | 100 | 28 | 28 | 27 | 12 | 7 | 11 | 12 | 12 | 8 | 5 |
| Human PNPLA4 | 35 | 35 | 36 | 34 | 39 | 35 | 30 | 30 | 32 | 28 | 100 | 61 | 28 | 27 | 12 | 7 | 11 | 12 | 8 | 5 |
| Rat PNPLA4 | 37 | 37 | 37 | 34 | 39 | 33 | 33 | 32 | 32 | 28 | 61 | 100 | 30 | 30 | 19 | 16 | 13 | 11 | 13 | 14 |
| Human PNPLA5 | 34 | 34 | 37 | 33 | 33 | 27 | 25 | 26 | 34 | 27 | 28 | 30 | 100 | 65 | 9 | 9 | 9 | 7 | 8 | 6 |
| Mouse PNPLA5 | 34 | 33 | 33 | 34 | 33 | 26 | 25 | 24 | 32 | 12 | 27 | 30 | 65 | 100 | 7 | 7 | 5 | 11 | 9 | 11 |
| Human PNPLA6 | 9 | 7 | 13 | 11 | 8 | 8 | 10 | 8 | 10 | 7 | 12 | 19 | 9 | 7 | 100 | 94 | 59 | 60 | 9 | 8 |
| Mouse PNPLA6 | 14 | 9 | 16 | 12 | 10 | 8 | 10 | 6 | 9 | 11 | 7 | 16 | 9 | 7 | 94 | 100 | 61 | 58 | 6 | 7 |
| Human PNPLA7 | 14 | 10 | 6 | 7 | 8 | 8 | 7 | 5 | 6 | 12 | 11 | 13 | 9 | 5 | 59 | 61 | 100 | 82 | 11 | 7 |
| Mouse PNPLA7 | 13 | 10 | 13 | 12 | 9 | 10 | 12 | 6 | 9 | 12 | 12 | 11 | 7 | 11 | 60 | 58 | 82 | 100 | 9 | 8 |
| Human PNPLA8 | 9 | 9 | 10 | 8 | 11 | 5 | 3 | 5 | 5 | 8 | 8 | 13 | 8 | 9 | 9 | 6 | 11 | 9 | 100 | 83 |
| Mouse PNPLA8 | 7 | 10 | 6 | 8 | 8 | 4 | 3 | 5 | 6 | 5 | 5 | 14 | 6 | 11 | 8 | 7 | 7 | 8 | 83 | 100 |

[^1] Abbreviations: ATGL, adipose triglyceride lipase; PNPLA, patatin phospholipase.
undertaken in each case. Of particular interest are the nodes demonstrating highly significant separations for each of the vertebrate PNPLA-like group 1 gene family sequences (ATGL, PNPLA4, PNPLA1, PNPLA3, and PNPLA5) sequences during vertebrate evolution, which supports the distinct status for each of these genes. However, there are differences in the distribution of some gene families, with PNPLA3 and PNPLA5 being observed only among eutherian mammalian genomes, while PNPLA1 was present in both eutherian and marsupial genomes, but apparently absent in other vertebrate genomes (Table 1, Figure 6). In contrast, both $A T G L$ and PNPLA4 genes were found among all vertebrate genomes examined, with $A T G L$ also found in an invertebrate genome (sea squirt, C. intestinalis). This suggests that invertebrate $A T G L$ may have served as a primordial gene for subsequent gene duplication events generating the five families of vertebrate PNPLA-like group 1 lipase genes. The likely sequence of these proposed group 1 PNPLA-like gene duplication events is summarized in Figure 6: an invertebrate $A T G L$ gene undergoing duplication to form ancestral vertebrate $A T G L$ and PNPLA4 genes; vertebrate PNPLA4 and $A T G L$ genes are subsequently retained into mammalian genomes; vertebrate $A T G L$ undergoes successive gene duplications to form primordial mammalian PNPLA1, PNPLA3, and PNPLA5 genes, which are retained throughout subsequent eutherian mammalian evolution. The PNPLA-like gene phylogram for group 2 sequences is also consistent with a primordial invertebrate PNPLA7 gene serving as an ancestor for vertebrate $P N P L A 7$, followed by a gene duplication event generating the PNPLA6 and PNPLA7 sequences in eutherian mammalian genomes. The phylogram for vertebrate PNPLA8 sequences suggests that this is an ancient gene in invertebrate and vertebrate evolution.

Previous phylogenetic studies of PNPLA-like proteins are predominantly consistent with these results, although these reports were restricted to vertebrate PNPLA1-8, ${ }^{64}$ chicken PNPLA1-8, ${ }^{65}$ and pig PNPLA2-5 ${ }^{66}$ genes and proteins, with each group inclusive of $A T G L$ genes and proteins. In each case, evidence of separation into distinct PNPLAlike family groups was reported. However, this current study has also examined PNPLA-like genes and proteins from an invertebrate genome (sea squirt, C. intestinalis), and provides evidence for at least three ancestral genes (Figure 5): $A T G L$ (proposed ancestral gene for vertebrate PNPLA1, ATGL, PNPLA3, PNPLA4, and PNPLA5 genes and proteins); PNPLA7 (proposed ancestral gene for vertebrate PNPLA6 and PNPLA7 genes and proteins); and PNPLA8 (proposed ancestral gene for the vertebrate PNPLA8 gene and protein).

## Summary

The results of this study support those of previous studies for at least eight vertebrate $P N P L A$-like genes and encoded lipases, including five group 1 genes, namely $A T G L$ (encoding adipose triglyceride lipase), PNPLA1, PNPLA3, and PNPLA5, and PNPLA4 genes; two group 2 genes, PNPLA6 (encoding neuropathy target esterase) and PNPLA7; and the group 3 gene, PNPLA8 (encoding cytosolic phospholipase A2). Vertebrate ATGL sequences shared putative key conserved sequences, including two proposed active site motifs: Cys15-Gly16-Xaa-Leu18; and Gly45-Xaa-Ser47-Yaa-Gly49; active site residues (Ser47 and Asp166 for human ATGL); a phosphorylated Ser308 site; and several conserved serine residues. A predicted N -terminal membrane binding site proposed as a membrane anchor involved in the localization of the enzyme near adipocyte stores was also conserved within vertebrate ATGL sequences.


[^2]

Figure 3 Predicted tertiary structures for vertebrate ATGL.
Notes: The predicted structures for pig, opossum, zebrafish, and sea squirt ATGL subunits are based on the reported structure for potato patatin ${ }^{25}$ and obtained using the SWISS MODEL web site (http://swissmodel.expasy.org/workspace/). The rainbow color code describes the three-dimensional structures from the N -terminus (blue) to C-terminus (red color); predicted $\alpha$-helices, $\beta$-sheets, proposed active site "motifs" I and 2; and active site residues (Ser47 and Aspl66) are shown.
Abbreviation: ADPL, adipose triglyceride lipase.


Figure 4 Gene structure and major isoforms for human ATGL.
Notes: From AceView website ${ }^{46}$ http://www.ncbi.nlm.nih.gov/IEB/Research/Acembly/Mature isoform variants (designated as, eg, "a","b") are shown for each ATGL transcript; capped $5^{\prime}$-ends and $3^{\prime}$-ends for the predicted mRNA sequences are identified; predicted $\mathrm{CpG}^{\prime}$ islands ( CpGI 31 and CpGI 08 ), miRNA binding sites ( miR - $124 / 506$ and miR 377); transcription factor binding sites (AHRARNT, ${ }^{68}$ estrogen receptor, ${ }^{69}$ early growth response 4 receptor, ${ }^{70}$ FOXOI, ${ }^{71}$ PPARG,,${ }^{72}$ COUP, ${ }^{73}$ PPARA, $,{ }^{74} \mathrm{HMXI},{ }^{75}$ BACHI ${ }^{76}$ ); and a scale of base pairs of nucleotide sequences are shown.
Abbreviation: TFBS, transcription factor binding sites.

GNF expression atlas 2 data from U133A and GNF1 human chips for ATGL


GNF expression atlas data from U74a mouse chip for ATGL

based on Su et al., 2004
Figure 5 Human and mouse tissue gene expression "heat maps" for ATGL.
Notes: Taken from the human and mouse genome browsers (http://genome.ucsc.edu); ${ }^{44}$ GNF Expression Atlas 2 data using expression chips for human and mouse ATGL (http://biogps.gnf.org); ${ }^{52}$ comparative gene expression levels among human tissues: red (high); black, (intermediate); and green (low) expression levels.


Figure 6 Phylogenetic tree of vertebrate ATGL and other PNPLA-like amino acid sequences with sea squirt (Ciona intestinalis) ATGL, PNPLA7, and PNPLA8 sequences.
Notes: The tree is labeled with the ATGL or other PNPLA-like gene name and the name of the vertebrate or invertebrate; note the major clusters into three groups of PNPLA-like gene families: Group I (ATGL, PNPLAI, PNPLA3, PNPLA4, and PNPLA5); Group 2 (PNPLA6 and PNPLA7); and Group 3 (PNPLA8); proposed ancestral invertebrate ATGL, PNPLA7, and PNPLA8 sequences which were used to "root" the tree are shown; gene duplication events generating these gene families are proposed to have occurred at different times: prior to the vertebrate common ancestor (ATGL and PNPLA4); prior to the appearance of marsupial mammals (PNPLAI and PNPLA3); and prior to the appearance of eutherian mammals (PNPLA6). A genetic distance scale is shown. The number of times a clade (sequences common to a node or branch) occurred in the bootstrap replicates is shown. Only replicate values of 90 or more which are highly significant are shown. 100 bootstrap replicates were performed in each case.

Gene expression data ${ }^{52}$ showed that the human $A T G L$ gene is broadly expressed at higher levels than those for the average gene. ${ }^{45} \mathrm{~A} \mathrm{CpG}$ island in the ATGL promoter region and several transcription factor binding sites of the $A T G L$ gene may contribute to this high level of gene expression. Two miRNA binding sites localized in the extended $3^{\prime}$ 'noncoding region of the $A T G L$ mRNA may contribute to the regulation of this gene during development. Phylogeny studies using several vertebrate and invertebrate ATGL subunits demonstrated that this is an ancient gene in vertebrate and invertebrate evolution. In addition, these studies of PNPLA-like subunits have suggested that there are at least three major groups based on their comparative structures and sequence phylogeny: group 1, ATGL (as the proposed ancestral gene), PNPLA1, PNPLA3, PNPLA4, and PNPLA5; group 2, PNPLA6 and PNPLA7; and group 3, PNPLA8. Ancestral invertebrate group 1, group 2, and group 3 PNPLA-like genes and proteins were observed in the sea squirt ( $C$. intestinalis) genome and used to "root" the phylogenetic tree for vertebrate $A T G L$ and other PNPLA-like genes.

## Acknowledgment

The expert assistance of Bharat Patel of Griffith University, Brisbane, Australia, with the phylogenetic analysis is gratefully acknowledged.

## Disclosure

The author reports no conflict of interest in this work.

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## Supplementary materials



Figure SI Predicted transmembrane structures (THMMH) for vertebrate ATGL sequences.
Notes: Derived from predictions using CBS web tools (Center for Biological Sequence Analysis, Technical University of Denmark (http://www.cbs.dtu.dk/services/THMMH/). Red peaks indicated positioning of predicted transmembrane structures (THMMH amino acid sequences are identified); pink-sequence outside membrane; blue sequence inside membrane; $y$ axes show probabilities of forming THMMH ( $0 \rightarrow \mathrm{I}$ ); x axes show amino acids; individual ATGL sequences examined include human (Homo sapiens), cow (Bos taurus), opossum (Monodelphis domestica), and zebrafish (Danio rerio).


Figure S2 Comparative sequences for vertebrate ATGL sequences.
Notes: Derived from the UCSC genome browser using the comparative genomics track to examine alignments and evolutionary conservation of ATGL gene sequences; genomic sequences aligned for this study included primate (human and rhesus), other mammal (mouse, dog, elephant, and opossum), and other vertebrate species (chicken, frog, and zebrafish); conservation measures were based on conserved sequences across all of these species in the alignments; regions of sequence identity are shaded.

Table SI Comparative intron sizes for vertebrate ATGL genes

| Vertebrate | ATGL gene size (bps) | Intron I bps | Intron 2 bps | Intron 3 bps | Intron 4 bps | Intron 5 bps | Intron 6 bps | Intron 7 bps | Intron 8 bps |
| :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: | :---: |
| Human | 5,141 | 1,722 | 97 | 373 | 920 | 106 | 142 | 183 | 86 |
| Mouse | 4,379 | 1,799 | 86 | 347 | 215 | 101 | 85 | 189 | 99 |
| Rat | 4,286 | 1,773 | 83 | 353 | 189 | 100 | 82 | 188 | 84 |
| Pig | 4,273 | 1,770 | 89 | 320 | 191 | 109 | 79 | 170 | 87 |
| Cow | 4,157 | 1,564 | 79 | 355 | 221 | 101 | 96 | 191 | 92 |
| Opossum | 4,792 | 1,543 | 146 | 184 | 783 | 106 | 314 | 177 | 69 |
| Chicken | 30,118 | 21,146 | 171 | 4,655 | 848 | 886 | 423 | 730 | 710 |
| Lizard | 45,618 | 29,925 | 239 | 6,135 | 1,526 | 2,008 | 652 | 1,576 | 2,048 |
| Frog | 14,284 | 6,276 | I,52 I | 1,610 | 278 | 672 | 116 | I,165 | I,055 |

Note: Numbers show the comparative sizes for vertebrate ATGL introns.
Abbreviation: bps, base pairs of nucleotides.

Table S2 Identification, location and comparative functions for human ATGL transcription factor binding sites

| TFBS | Name | Chromosome coordinates | Chromosome location (strand) | Role |
| :---: | :---: | :---: | :---: | :---: |
| AHRARNT | Aryl hydrocarbon receptor ${ }^{1}$ | chrll:819809-819824 | Exon 2 (-) | Activates multiple phase I and II xenobiotic chemical metabolizing enzyme gene expression |
| ER | Estrogen receptor ${ }^{2}$ | chrll:819883-81990\| | Exon 2 (-) | Involved in the hormonal regulation of eukaryotic gene expression |
| NGFIC | Early growth response $4^{3}$ | chrll:820189-820200 | Intron 2 (+) | Transcription activator of genes required for mitogenesis and differentiation. |
| FOXOI | Forkhead fox protein $\mathrm{Ol}^{4}$ | chrll:820601-820614 | Intron $2(+)$ | Transcription factor which acts as a regulator of cell responses to oxidative stress |
| PPARG | Peroxisome proliferatoractivated receptor gamma ${ }^{5}$ | chrll:820704-820724 | Intron 2 (-) | Activates genes encoding enzymes of the peroxisomal beta-oxidation pathway of fatty acids |
| COUP | Chicken ovalbumin upstream promoter ${ }^{6}$ | chrll:820708-82072 I | Intron 2 (+) | Stimulates transcription initiation |
| PPARA | Peroxisome proliferatoractivated receptor alpha ${ }^{7}$ | chrll:820708-820727 | Intron 2 (-) | Activates genes encoding enzymes of the peroxisomal beta-oxidation pathway of fatty acids |
| HMXI | Homeobox protein HMXI ${ }^{8}$ | chrll:821301-821310 | Intron 2 (+) | Transcription factor involved in the differentiation of neuronal cell types |
| PPARA | Peroxisome proliferatoractivated receptor alpha ${ }^{7}$ | chrll:82\|315-82|334 | Intron 2 (+) | Activates genes encoding enzymes of the peroxisomal beta-oxidation pathway of fatty acids |
| BACHI | Transcription regulator protein BACHI ${ }^{9}$ | chrll:82\|379-821393 | Intron 2 (-) | Coordinates gene transcription activation and repression by MAFK ${ }^{10}$ |
| MYB | Transcriptional activator Myb ${ }^{\text {I }}$ | chrll:82\|397-82|406 | Intron 3 (-) | Control of proliferation and differentiation of hematopoietic progenitor cells |

Notes: Predicted transcription factor binding sites identified: 'AHRARNT;;6,2 estrogen receptor; ${ }^{67,3}$ early growth response 4 receptor; ${ }^{68,4}$ FOXOI; ${ }^{69,5}$ PPARG; ${ }^{70,6}$ COUP;; ${ }^{7,7}$ PPARA; ${ }^{72,8} \mathrm{HMXI} ; ;^{73,9} \mathrm{BACHI} ;{ }^{74,10}$ MAFK; ${ }^{75,11} \mathrm{MYB}^{76}$.
Abbreviations: TFBS, transcription factor binding sites; UCSC, University of California Santa Cruz.

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[^1]:    Notes: Numbers show the percentage of amino acid sequence identities. Numbers in bold show higher sequence identities for more closely related PNPLA-like family members.

[^2]:    Figure 2 Amino acid sequence alignments for potato patatin, mouse ATGL, PNPLAI, PNPLA3, and PNPLA5, and rat PNPLA4 patatin "motif" sequences. Notes: See Table I for sources of mouse and rat PNPLA-like sequences; the potato patatin "motif" sequence was obtained from. ${ }^{25}$ *Identical residues; I or 2 conservative substitutions; I or 2 nonconservative substitutions; oxyanion hole motif (Gly-Gly-Xaa-Lys for potato patatin); active site motif (Gly-Xaa-Ser-Xaa-Gly); catalytic dyad residues Ser and Asp; predicted helix $\alpha I \alpha 2$; and predicted sheet $\beta$ I $\beta 2$. Patatin "motif" sequences examined included Po PATI (potato patatin); Mo ATGL (mouse ATGL); Mo PLAI (mouse PNPLAI); Mo PLA3 (mouse PNPLA3); Ra PLA4 (rat PNPLA4); and Mo PLA5 (mouse PNPLA5).

