

Product datasheet for **RG237715**

AIPL1 (NM_001285401) Human Tagged ORF Clone

Product data:

Product Type:	Expression Plasmids
Product Name:	AIPL1 (NM_001285401) Human Tagged ORF Clone
Tag:	TurboGFP
Symbol:	AIPL1
Synonyms:	AIPL2; LCA4
Mammalian Cell Selection:	Neomycin
Vector:	pCMV6-AC-GFP (PS100010)
E. coli Selection:	Ampicillin (100 ug/mL)
ORF Nucleotide Sequence:	>RG237715 representing NM_001285401. Blue=ORF Red=Cloning site Green=Tag(s)

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GCTCGTTTGTAGTAACCGTCAGAATTTGTAAACGACTCACTATAGGGCGCCGGGAATTCGTCGACTG
GATCCGGTACCGAGGAGATCTGCCGCCGCGATCGCC
ATGGATGCCGCTCTGCTCCTGAACGTGGAAGGGGTCAAGAAAACCATCTGCACGGGGCACGGGCGAG
CTCCCAAACCTTCATCACCGGATCCCGAGTGATCTTTCAATTCGCACCATGAAATGTGATGAGGAGCGG
ACAGTCATTGACGACAGTCGGCAGGTGGGCCAGCCCATGCACATCATCATCGAAACATGTTCAAGCTC
GAGGTCTGGGAGATCCTGCTTACCTCCATGCGGGTGCACGAGGTGGCCGAGTTCTGGTGCGACACCATC
CACACGGGGTCTACCCATCCTATCCCGAGCCTGAGGCAGATGGCCAGGGCAAGGACCCACAGAG
TGGCACGTGCACACGTGCGGGCTGGCCAACATGTTGCGCTACCACACGCTGGGCTACGAGGACCTGGAC
GAGCTGCAAGAAGGAGCCTCAGCCTCTGGTCTTTGTGATCGAGCTGCTGCAGGTTGATGCCCGAGTGAT
TACCAGAGGGAGACCTGGAACCTGAGCAATCATGAGAAGATGAAGGCGGTGCCCGTCTCCACGGAGAG
GGAAATCGGCTCTTCAAGCTGGGCGCTACGAGGAGGCCTCTTCCAAGTACCAGGAGGCCATCATCTGC
CTAAGGAACCTGCAGACCAAGTGCTGTGAAGAAGGAGGAGTACTATGAGGTGCTGGAGCACACCAAGT
GATATTCTCCGGCACCAACCGGATCGTGAAGGCCTACTACGTGCGTGCCCGGGCTCACGCAGAGGTG
TGAATGAGGCCGAGGCCAAGGCGGACCTCCAGAAAGTGTGGAGCTGGAGCCGTCATGCAAGAAGCGG
GTGCGCAGGGAGCTGAGGCTGCTGGAGAACCAGCATGGCGGAGAAGCAGGAGGAGGAGCGGCTGCGCTGC
CGGAACATGCTGAGCCAGGGTCCACAGCAGCCTCCCGCAGAGCCACCCACAGAGCCACCCGACAGTCA
TCCACAGAGCCACCTGCAGAGCCACCCACAGCACCATCTGCAGAGCTGTCCGAGGGCCCCCTGCAGAG
CCAGCCACAGAGCCACCCCGTCCCGAGGGCACTCGCTGCAGCAC
ACGCGTACGCGGCCGCTCGAG - GFP Tag - GTTTAAAC
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Protein Sequence: >Peptide sequence encoded by RG237715
 Blue=ORF Red=Cloning site Green=Tag(s)

MDAALLLNVEGVKKTILHGGTGELPNFITGSRVIFHFRTMKCDEERTVIDDSRQVGQPMHIIIGNMFKL
 EVWEILLTSMRVHEVAEFWCDTIHTGVYPILSRSLRQMAQKDPTEWHVHTCGLANMFAYHTLGYEDLD
 ELQKEPQPLVFVIELLQVDAPSDYQRETWNLSNHEKMKAVPVLHGEGNRLFKLGRYEEASSKYQEAIIC
 LRNLQTKCLLKKEEYVELEHTSDILRHHPGIVKAYVVRARAHAEVWNEAEAKADLQKVLELEPSMQKA
 VRRELRLLENRMAEKQEEERLRCRNMLSQGATQPPAEPPTPEPPAQSSTEPPAEPPTAPSAELSAGPPAE
 PATEPPPPSPGHSLQH
 TRTRPLEMESDESGLPAMEIECRITGTLNGVEFELVGGGEGTPEQGRMTNKMSTKGALTFSPYLLSHV
 MGYGFYHFGTYPSTYENPFLHAINNGGYNTRIEKYEDGGVLHVSFSYRYEAGRVIGDFKVMGTGFPED
 SVIFTDKIIIRSNATVEHLHPMGDNDLDGSFTRTFSLRDGGYSSVVDSHMHFKSAIHPSILQNGGPMFA
 FRRVEEDHSNTELGIVEYQHAFKTPDADAGEERV

Restriction Sites: SgfI-MluI

Cloning Scheme:



ACCN: NM_001285401

ORF Size: 1080 bp

OTI Disclaimer: The molecular sequence of this clone aligns with the gene accession number as a point of reference only. However, individual transcript sequences of the same gene can differ through naturally occurring variations (e.g. polymorphisms), each with its own valid existence. This clone is substantially in agreement with the reference, but a complete review of all prevailing variants is recommended prior to use. [More info](#)

OTI Annotation: This clone was engineered to express the complete ORF with an expression tag. Expression varies depending on the nature of the gene.

Components: The ORF clone is ion-exchange column purified and shipped in a 2D barcoded Matrix tube containing 10ug of transfection-ready, dried plasmid DNA (reconstitute with 100 ul of water).

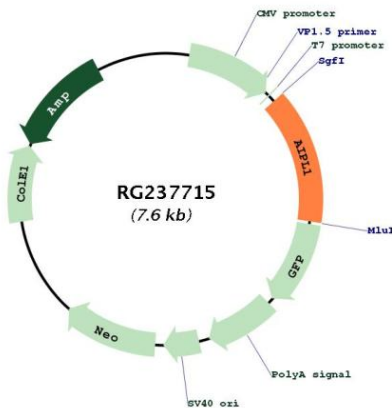
RefSeq: [NM_001285401.3](#)

RefSeq Size: 2918 bp

RefSeq ORF: 1083 bp
Locus ID: 23746
UniProt ID: [Q9NZN9](#)
Cytogenetics: 17p13.2
Protein Families: Druggable Genome
MW: 41.4 kDa

Gene Summary: Leber congenital amaurosis (LCA) is the most severe inherited retinopathy with the earliest age of onset and accounts for at least 5% of all inherited retinal diseases. Affected individuals are diagnosed at birth or in the first few months of life with nystagmus, severely impaired vision or blindness and an abnormal or flat electroretinogram. The photoreceptor/pineal-expressed gene, AIPL1, encoding aryl-hydrocarbon interacting protein-like 1, is located within the LCA4 candidate region. The encoded protein contains three tetratricopeptide motifs, consistent with chaperone or nuclear transport activity. Mutations in this gene may cause approximately 20% of recessive LCA. Alternative splicing results in multiple transcript variants. [provided by RefSeq, Jan 2014]

Product images:



Circular map for RG237715