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Identification of Immune Relevant Genes Using Expressed Sequence Tags (ESTs) in Common carp (*Cyprinus carpio*) Gills and Intestine

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Identification of immune related molecules that function in fish organs is important for better understanding of the host defense mechanisms in fish. Expressed sequence tags (ESTs) represent the expressed portion of a genome, so they have proven to be useful tool for gene identification and confirmation of gene predictions. In the present study, a transcriptome analysis of carp gill and intestine EST libraries have been done as an attempt to identify the immune relevant genes expressed in those organs. A total of 2148 EST clones were generated from the two libraries: 1099 clones from gill library in which 632 clones were matched with functional proteins and 1049 clones from intestine library in which 559 clones were matched with functional proteins. The results of gill EST library has more frequency of innate immune–molecules containing MHC class I and MHC class II, which showed 18.3% of the immune molecules than those in intestine library which represents 10.8% only. The gill library also showed higher frequency of the cytokines and chemokine molecules and/or their receptors compared to the intestine library.

Key words: Common carp, expressed sequence tags (EST), gills, intestine, immune relevant genes.

INTRODUCTION

Expressed sequence tags (ESTs) are single-pass reads of approximately 200-800 base pairs (bp) generated from randomly selected cDNA clones. They represent the expressed portion of a genome, so they have proven to be useful for gene identification and confirmation of gene predictions. They therefore represent a lowcost substitute to the full genome sequencing (Parkinson and Blaxter, 2009). Currently, the number of fish-related ESTs in the public databases is still small compared with mammalian sequences and there are relatively few tissuespecific cDNA libraries (Ton et al., 2000). Large-scale EST studies have been conducted on zebrafish (Gong et al., 1997; Gong, 1999; Kelly et al., 2000; Borchardt et al., 2010). In addition to other commercially important species as fugu (Aparicio et al., 2002); Japanese flounder (Nam et al., 2000; Kono and Sakai, 2001; Nam et al., 2003); channel catfish (Ju et al., 2000); rainbow trout (Kono et al., 2000); Atlantic salmon (Davey et al., 2001) and common carp (Savan and Sakai, 2002; Kono et al., 2004; Ji et al., 2012; Liao et al., 2013). Still, comprehensive information on mRNA levels at steady state is not known for the most known fish transcripts (Virlon et al., 1999). Actually, EST-based approaches have been especially effective in fish for identification of cytokines, a wide range of small proteins that are secreted from various cells of the immune system to act on immune-related cells, functioning as a signal messenger for immune responses (Bird *et al.*, 2006; Savan and Sakai, 2006; Wang *et al.*, 2009). This was largely because many of the cytokines including interleukins, interferons, and chemokines of fish have poorly conserved primary structure compared with mammalian homologues, inhibiting application of conventional homology cloning techniques dependent on high degree of sequence conservation.

The common carp, Cyprinus carpio, belonging to the family Cyprinidae is one of the most widely cultured fish species in aquaculture and considered to be an important source of animal protein (FAO, 2009). Despite the importance in global aquaculture, genomic information in the cyprinid is still very limited, except for zebrafish (Danio rerio) (Christoffels et al., 2006), partly due to the polyploidy that represent a characteristic feature of several members of this family (Larhammar and Risinger, 1994; David et al., 2003). Since the species, Cyprinus carpio, is believed to have been established as a result of allotetraploidization about 50 million years ago, the pseudotetraploid nature of the carp genome has been implicated to have an impact to allow diversification of various innate immune-related genes in this species (David et al., 2003, 2007; Christoffels et al., 2006).

To date, identification of immune-related genes have gained considerable successes in the common carp, resulting from EST analysis of immune-related organs such as kidney (Savan and Sakai 2002; Kono *et al.*, 2004), testes (Christoffels *et al.*, 2006), thymus (Huang *et al.*, 2009), peritoneal cell exudate (Fujiki *et al.*, 1999, 2000), in addition to skin, which is an immediate interface between fish body and environmental water (Gonzalez *et al.*, 2007). The present study was started as further attempts to unveil novel immune-related genes by applying the EST approaches to the gill and intestine, which

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are considered to be important interfaces for outer environment and potential sites of microbial infections.

In fish gill, intestine and skin are organs exposed to the outer environment. They are considered the first line of defense against any invading pathogen or infection. Therefore identification of immune related molecules that function in those organs is important for better understanding of the host defense mechanisms in fish. In the present study, a recent transcriptome analysis of carp gill and intestine EST libraries have been done as an attempt to identify the immune relevant genes expressed in those organs. The obtained data will be a good tool for establishment of prevention of infectious diseases, which damage the cultured fish stock.

MATERIALS AND METHODS

Reagents

Scleroglucan (a fungal β -1,3-glucan from Sclerotium glucanicum) and Edwardsiella tarda NG8104 were obtained as described elsewhere (Yano et al., 1991). E. tarda was cultured using the heart infusion agar and broth (Eiken Chemical Co., Tokyo, Japan) and harvested during the logarithmic phase. ISOGEN reagent was purchased from Nippon Gene (Tokyo, Japan). Quick Prep Micro mRNA kit was from GE Healthcare (Tokyo, Japan). cDNA Synthesis Kit, Uni–ZAP XR vector and Gigapack II Gold were obtained from Stratagene (USA), PCR Super Mix HF (Invitrogen), Exo SAP–IT kit was from GE Healthcare (Tokyo, Japan).

Fish

Mature common carp (*Cyprinus carpio*) weighing about 1 kg purchased from a local fish farm were maintained in a recirculation tank system kept at 25°C with daily feeding of commercial pellets for a month before experiments.

Construction of cDNA libraries for EST sequence analysis of carp gill and intestine

The carp was stimulated by oral administration of scleroglucan for 3 days at a dose of 5 mg/kg–body weight/ day, and by an immersion in a live *E. tarda* suspension $(3 \times 10^7 \text{ CFU/ml})$ for five minutes in the third day. Then the fish was kept without feeding for two more days until sacrificed for RNA isolation.

The fish was anesthetized in 25 mg/L quinaldine and subjected to cannulation to the bulbus arteriosus in the thoracic cavity by surgical operation. Then the circulating blood was removed as completely as possible by perfusion with freshwater fish physiological saline (0.75% NaCl, 0.02% KCl, 0.02% CaCl₂ and 0.002% NaHCO₃; Kawamoto, 1970) supplied from the cannula. After the color of the gill became almost white, the gill lamellae and the intestine (the mid gut part or the second segment) were excised and immediately homogenized in ISOGEN reagent for total RNA purification according to the manufacturer's instructions. Poly (A)⁺ RNA was purified using Quick Prep Micro mRNA kit. Double stranded cDNA was synthesized with the cDNA Synthesis Kit, and after ligation with EcoRI adaptor and digestion with XhoI, unidirectional cDNA libraries for gill and intestine were constructed in Uni–ZAP XR vector. In vitro packaging was performed using Gigapack III Gold. Titers of 3.6×10^7 pfu and 2×10^7 pfu were obtained for the gill and intestine cDNA libraries, respectively. The libraries were amplified once, aliquoted, and stored at -80° C.

Nucleotide sequencing

Phages of an aliquot of the once amplified ZAP library were converted to pBluescript SK (–) plasmids by *mass excision* with ExAssist helper phage and XLOLR bacterial host strain, according to the manufacturer's instructions. Resultant recombinant colonies were randomly selected for colony–direct PCR of the insert using PCR Super Mix HF with a set of M13 forward and reverse primers under the following conditions: 30 cycles of 94°C for 30 sec, 55°C for 30 sec, and 68°C for 7 min. After analysis of the amplicon size by agarose gel electrophoresis, templates for sequencing reactions were purified from the amplified product using Exo SAP–IT kit.

Nucleotide sequences were determined from the 5'-end of each cDNA insert with the PCR-amplified template and the sequencing primer was ACAAAAGCTGGAGCTCCACCG. ABI Prism Big Dye™ Terminator Cycle Sequencing Ready Reaction Kit version 3.0 was used for ABI310 and ABI377 sequencers (Applied Biosystems), and GenomeLab[™] DTCS- Quick Start Kit was for analysis on CEQ8800 sequencer (Beckman). In addition, mass sequencing was also conducted in Takara Bio Co. (Shiga, Japan) using ABI3700 sequencer with the same primer.

Sequence analysis

Nucleotide sequences were clustered and cleared from vector-derived sequences by using ATGC software (GENETYX), and the non-redundant insert sequences were subjected to a BlastX homology search against non-redundant set (nr) of amino acid sequence on the NCBI database (<u>http://www.ncbi.nlm.nih.gov/blast/</u>). The homology was determined to be significant when the E-value was less than 10⁻⁸. The sequences that showed homology to known proteins were classified based on KEGG orthology (KO) System (<u>http://www.genome.jp/</u> <u>kegg/ko.html</u>). The EST frequency (percentage) was calculated from the number of clone in the number of total clones.

RESULTS

Annotation analysis of gill and intestine EST clones

As shown in Table 1, the database analysis of a total number of gill EST 1099 random clones yields 632 clones that were matched with functional proteins and 134 clones with unknown function while 333 clones showed non–significant similarity. In case of intestine cDNA library, the database analysis of EST 1049 random clones yields 559 clones that were matched with functional proteins and 223 clones were matched with unknown func-

| | Gills ES | T clones | Intestine EST clones | | |
|---------------------------|----------------------------------------|----------|----------------------|----------------|--|
| Group | Clones number Percentage (%) Clones nu | | Clones number | Percentage (%) | |
| Functional | 632 | 57.5 | 559 | 53.3 | |
| Unknown function | 134 | 12.2 | 223 | 21.3 | |
| No significant similarity | 333 | 30.3 | 267 | 25.4 | |
| Total | 1099 | 100 | 1049 | 100 | |

Table 1. Annotation analysis of the carp gill and intestine EST clones

tion while 267 clones showed non-significant similarity.

Clustering and functional classification of the EST clones

The sequenced clones that showed homology to known proteins were classified based on KEGG Orthology (KO) System. The summary of the results is shown in Table 2. For gill library, out of 143 clones (13.0%) classified into metabolism category, 185 clones (17.0%) classified into genetic information processing category, 80 clones (7.3%) classified into environmental information processing category, 212 clones (19.3%) classified into cellular processing category, 12 clones (1.1%) classified into diseases category while unknown function represents 134 clones (12.0%) and the non-significant similarity represents 333 clones (30.3%). For intestine library, out of 163 clones (15.5%) classified into metabolism category, 144 clones (13.7%) classified into genetic information processing category, 87 clones (8.3%) classified into environmental information processing category, 161 clones (15.4%) classified into cellular processing category, 4 clones (0.4%) classified into disease category while unknown function represents 223 clones (21.3%) and the non-significant similarity represents 267 clones (25.4%).

Clusters of immune related genes of the gill and intestine EST clones

All the EST sequences have been deposited in DDBJ/ EMBL/GenBank databases under the accession numbers from DC995888 to DC997897. Of the many transcripts that had homology to known genes in the gill and intestine libraries, a number appeared to be related to the host defense mechanisms were occupied 104 clones (9.5%) and 93 clones (8.9%) from gill and intestine EST clones respectively. In addition, the gill library has more frequency of innate immune–molecules containing MHC class I, MHC class II which showed 18.3% of the immune molecules than those in intestine library which represents 10.8% only. Gill library also showed higher frequency of the cytokines and chemokine molecules and/ or their receptors compared to cytokines expressed from intestine library (Tables 3 and 4).

DISCUSSION

In fish, gills, intestine and skin are important immune organs (Gonzalnez et al., 2007). In order to identify the different immune molecules inside carp gills and intestine, cDNA libraries have been constructed after stimulation with Edwardsiella tarda. In this paper, stimulation of carp gill and intestine leads to expression of newly discovered molecules in carp such as XCR1, CysLTR2 receptors, and melanin-inhibiting protein. Others showed higher similarity to mammalian homologues rather than to fish homologues such as IL-8, an interleukin-8 (IL-8like CXC-chemokine) encoded by EST clone accession number (DC996606) obtained from gill library shows significantly higher sequence similarity to sheep (Ovis aries) than to chemokines from other fish species. This clone was designated CaIL-8 and now considered the second legend of IL-8 family in carp fish (IL-8L2) (Abdelkhalek et al., 2009). Many other immune related molecules have been identified either related to innate immunity such as complement factors, cathepsins, lectins and fucolectin or acquired immunity such as CD2, CD3, CD83, T-cells receptor, MHC complex and immunoglobulin heavy chain and immunoglobulin light chain. In addition, chemokine CK-1, chemokine CXCL2, CC chemokine SCYA117, monocyte chemo-attractant protein, NF-kappa

Table 2. Functional classification of the carp gill and intestine EST clones

| | Gills EST clones | | | Intestine EST clones | | |
|--------------------------------------|--------------------|---------------------------------------|-------------------|----------------------|---------------------------------------|-------------------|
| Group | Clone frequency | immune– related clone frequency | Percentage (%) | Clone frequency | immune– related clone frequency | Percentage (%) |
| Metabolism | 143 | 0 | 13.0 | 163 | 0 | 15.5 |
| Genetic information processing | 185 | 0 | 17.0 | 144 | 4 | 13.7 |
| Environmental information processing | 80 | 21 | 7.2 | 87 | 14 | 8.3 |
| Cellular processing | 212 | 83 | 19.2 | 161 | 75 | 15.4 |
| Diseases | 12 | 0 | 1.1 | 4 | 0 | 0.4 |
| Total | 632 | 104 | 57.5 | 559 | 93 | 53.3 |

| Table 3. | Common carp gill EST | clones encoding for possib | ble immunologically-relevant proteins |
|----------|----------------------|----------------------------|---------------------------------------|
| | | | |

| Hit name | Species | Identified genes accession numbers | AA region/ AA–length (%) | E-value | Clone accession numbers |
|------------------------------------------------------|----------------------------|---------------------------------------|-------------------------------------------------------------|------------------------------------|------------------------------------|
| Adenosine A1 receptor | Xenopus laevis | CAB62281 | 434-667/326 (65%) | 4.00E-21 | DC996583 |
| Art2a protein | Mus musculus | AAI20754 | 25–174/287 (34%) | 2.00E-13 | DC996184 |
| B–cell translocation gene 2 | Danio rerio | NP_570997 | 182–670/165 (89%) | 3.00E-80 | DC996623 |
| Beta–2 microglobulin | Danio rerio | NP_998291 | 17-116/116 (77%) | 3.00E-42 | DC996032 |
| Beta–2–microglobulin precursor | Cyprinus carpio | Q03422 | 25-116/116 (97%) | 7.00E–50 | DC995975 |
| Bone morphogenetic protein 4 | Danio rerio | NP_571417 | 381-638/400 (96%) | 1.00E-39 | DC996563 |
| C–type lectin | Cyprinus carpio | BAA95671 | 1-151/163 (70%) | 1.00E-61 | DC996096 |
| Carcinoembryonic antigen–cell adhesion molecule 1 | Rattus norvegicus | NP_001029032 | 155-329/519 (30%) | 6.00E-18 | DC996273 |
| Cathepsin B preproprotein | Cyprinus carpio | BAE44111 | 1-195/330 (92%) | 2.00E-112 | DC996115 |
| Cathepsin L precursor | Sus scrofa | Q28944 | 139–666/334 (39%) | 2.00E-26 | DC996691 |
| CC chemokine SCYA117 | Ictalurus punctatus | ABA54964 | 67-261/96 (48%) | 6.00E-09 | DC996712 |
| CD2–associated protein | Xenopus tropicalis | CAJ83986 | 119-406/482 (80%) | 5.00E-23 | DC996668 |
| CD3 gamma delta–A | Salmo salar | ABO10198 | 90-452/181 (35%) | 2.00E-10 | DC996356 |
| CD83 | Ginglymostoma cirratum | AAO62993 | 26–127/194 (33%) | 3.00E-10 | DC996062 |
| Chemokine CK–1 | Oncorhynchus kisutch | AAF23861 | 154–378/100 (45%) | 4.00E–15, 2.00E–11 | DC996582, DC996601 |
| Chemokine receptor | Oncorhynchus mykiss | CAA05917 | 17-94/368 (66%) | 4.00E-20 | DC995899 |
| Chemokine C–X–C motif receptor 3 | Ctenopharyngodon idella | AAW69766 | 10-136/341 (76%) | 2.00E–27 | DC995896 |
| Claudin e | Danio rerio | NP_571840 | 87–656/209 (84%), 102–653/209 (80%), 100–654209 (85%) | 5.00E–70, 2.00E–67, 7.00E–55 | DC996343, DC996511, DC996304 |
| Claudin h | Danio rerio | NP_571842 | 189-656/214 (88%) | $7.00E{-}71$ | DC996449 |
| Connective tissue growth factor | Danio rerio | NP_00101501 | 69-662/345 (85%) | 2.00E-102 | DC996521 |
| Coagulation factor IX | Gallus gallus | NP_989674 | 1-107/471 (47%) | 4.00E-23 | DC996171 |
| CXC chemokine receptor–1 | Cyprinus carpio | AB010468 | 318-414/2852 (81%) | 1.00E-14 | DC996593 |
| CXCL2–like chemokine | Ictalurus punctatus | AAX40735 | 15-112/131 (42%) | 2.00E-09 | DC996211 |
| Cysteinyl leukotriene receptor 2(CysLTR2) | Sus scrofa | Q95N03 | 254-646/345 (31%) | 5.00E-11 | DC996646 |
| Fc fragment of IgG binding protein | Homo sapiens | EAW56926 | 62–658/3004 (41%) | 3.00E-30 | DC996806 |
| Fish-egg lectin | Cyprinus carpio | P68512 | 186–662/238(62%), 186–669/238(79%) | 2.00E–46, 4.00E–72 | DC996679, DC996706 |
| Fucolectin | Fundulus heteroclitus | AAU21486 | 98-670/227 (40%) | 7.00E–27, 2.00E–26 | DC996431, DC996804 |
| G–protein signaling modulator 2 | Danio rerio | NP_956732 | 299-661/649 (93%) | 1.00E-60 | DC996665 |
| Glucocorticoid receptor | Oncorhynchus mykiss | P49843 | 58-648/758 (82%) | 2.00E-94 | DC996525 |
| Granulin | Carassius auratus | ABD03952 | 114-509/158 (67%) | 4.00E-54 | DC996408 |
| Growth hormone protein gene | Catla catla | AY053360 | 557-650/11576 (84%) | 2.00E-18 | DC996592 |
| Immunoglobulin mu heavy chain | Oncorhynchus mykiss | ABH09025 | 142-669/198 (62%) | 2.00E-56 | DC996512 |
| Immunoglobulin heavy chain | Cyprinus carpio | BAA3471 | 252-445/445 (100%) | 8.00E-104 | DC996158 |
| Immunoglobulin light chain | Cyprinus carpio | BAB91004 | 133-666/204 (92%) | 5.00E-69 | DC996338 |
| Inhibitor of DNA binding 2 | Danio rerio | AAH56303 | 123–473/ 137 (97%) | 6.00E–57 | DC996526 |
| Interferon induced protein 2 | Ictalurus punctatus | AAN04880 | 359-57792 (47%) | 2.00E-10 | DC996736 |
| Interferon regulatory factor 5 isoform b | Bos taurus | AAX46454 | 300-426/499 (70%) | 5.00E-44 | DC995900 |
| Interleukin 8 | Ovis aries | NP_001009401 | 116-409/101 (36%) | 4.00E-08 | DC996606 |

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| Invariant chain like protein 2 | Cyprinus carpio | BAC53768 | 57–350/198 (100%), 121–654/198 (85%), 173–655198 (99%), 155–673/198 (86%) | 5.00E–54, 1.00E–74, 6.00E–89, 4.00E–74 | DC996412, DC996658, DC996688, DC996801 |
|--------------------------------------------------------------------------------|----------------------------|-------------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------|---------------------------------------------------------------------------|---------------------------------------------------------------------------|
| Invariant chain like protein 1 | Cyprinus carpio | BAC53767 | 53-664/234 (97%) | 3.00E-101 | DC996826 |
| Legumain (lgmn), mRNA | Danio rerio | NM_214759 | 57–295/2340 (77%) | 2.00E-43 | DC996814 |
| LY6D protein | Homo sapiens | AAH22806 | 56-352/129 (26%) | 2.00E-05 | DC996282 |
| Lymphocyte cytosolic plastin 1 | Danio rerio | NP_571395 | 117-656/624 (93%) | 3.00E-92 | DC996365 |
| Macrophage inflammatory protein–1– alpha(XCR1) | Homo sapiens | AAA36543 | 70–657/355 (36%) | 2.00E-26 | DC996427 |
| MHC class I antigen | Barbus intermedius | CAD12381 | 108-641/418 (85%) | 1.00E-87 | DC996562 |
| MHC class I antigen | Ctenopharyngodon idella | BAD01529, BAD01666 | 168 –635/254 (61%), 1–200 / 254 (65%), 146–295 /329 (77%) | 3.00E–50, 1.00E–72, 2.00E–64 | DC996370, DC996006, DC996193 |
| MHC class I UFA gene | Danio rerio | AAH66754 | 1-239/350(64%) | 4.00E-87 | DC995993 |
| MHC class II–associated invariant chain | Danio rerio | AAD24542 | $\begin{array}{l} 123-608/234\ (33\%),\\ 83-676/\ 234\ (72\%),\\ 86-646/\ 234\ (65\%),\\ 360-451/\ 234\ (66\%),\\ 4-145/234\ (44\%),\\ 7-224/234\ (33\%)\end{array}$ | 5.00E-24, 1.00E-80, 2.00E-64, 1.00E-29, 1.00E-53, 7.00E-24 | DC996452, DC996614, DC996673, DC996345, DC996157, DC996025 |
| MHC class II beta chain | Cyprinus carpio | CAA64709 | 71–664/249 (81%), 19–435/246 (89%) | 2.00E–89, 2.00E–53 | DC996478, DC996581 |
| MHC class II antigen | Barbus intermedius | CAD44962 | 86–652/224 (87%), 91–675/224 (61%), 1–183/224 (87%) | 6.00E–94, 1.00E–67, 6.00E–91 | DC996700, DC996813, DC996239 |
| MHC class I protein Zr2 | Cyprinus carpio | CAB96993 | 104–661/244 (93%) | 8.00E-100 | DC996744 |
| Metastasis suppressor homolog | Rattus norvegicus | AAC05159 | 123-647/266 (54%) | 7.00E-34 | DC996628 |
| Mucolipin 3 | Xenopus tropicalis | NP_001016804 | 162-410/553 (67%) | 2.00E-43 | DC996570 |
| Melanin-inhibiting protein | Drosophila melanogaster | AB205184 | 9–54/ 1208 (100%), 8–52/ 1208 (97%), 8–64/ 1208 (93%) | 6.00E–13, 3.00E–10, 6.00E–13 | DC996707, DC996600, DC996508 |
| Nuclear factor (erythroid–derived 2)–like 2 | Danio rerio | AAH45852 | 165-659/586 (78%) | 1.00E–50 | DC996663 |
| Nuclear NF-kappaB activating protein | Siniperca chuatsi | AAY79251 | 506-670/147 (75%) | $7.00E{-}16$ | DC996664 |
| Protein kinase C binding protein 1 | Danio rerio | NP_955935 | 50–532/1111 (69%) | 1.00E-39 | DC996751 |
| RAN, member RAS oncogene family, isoform CRA_a | Homo sapiens | EAW98515 | 34-669/236 (88%) | 2E-106 | DC996595 |
| RAR–related orphan receptor A isoform a | Homo sapiens | NP_599023 | 272–658/523 (87%) | 6.00E-62 | DC996349 |
| Ras-related protein Rab-7A | Bos taurus | Q3T0F5 | 318-656/207 (100%) | 9/E-59 | DC996754 |
| Regulator of G–protein signalling 3, isoform CRA_c | Homo sapiens | EAW87389 | 216-602/1088 (57%) | 1.00E-38 | DC996711 |
| Serine/threonine_protein phosphatase 2A (PP2A_alpha) | Sus scrofa | P67776 | 228-656/309 (97%) | 2.00E-77 | DC996578 |
| Serine/threonine kinase A | Danio rerio | NP_997731 XP_700037 | 57-656/320 (94%) | 4.00E-110 | DC996735 |
| Suppressor of typt1 | Danio rerio | NP_878281 | 46-675/632 (97%) | 4.00E-112 | DC996803 |
| Tumor–associated calcium signal transducer | Danio rerio | NP_99834, AAH66716 | 213–662/ 302 (74%), 223–660/ 302 (73%), 175–302 /302 (76%), | 2.00E–62, 3.00E–60, 4.00E–38, | DC996591, DC996533, DC996185 |
| Tumor suppressor candidate 3 | Mus musculus | NP_084530 | 110-640/347 (70%) | 4.00E-73 | DC996441 |
| Transmembrane 9 superfamily member 2 | Danio rerio | NP_997893 | 568–658 / 658 (100%) | 3.00E-39 | DC996261 |
| Tyrosine 3–monooxygenase / tryptophan 5–monooxygenase activation protein | Danio rerio | NP_958892, NP_998310 | 55–657/245 (96%), 127–654/ 242 (92%) | 9.00E–106, 1.00E–85 | DC996321, DC996501 |

| 1 | |
|---|--|
| | |

$\textbf{Table 4.} \ \text{Common carp intestine EST clones encoding for possible immunologically-relevant proteins}$

| | | | | | Clone |
|-------------------------------------------------------------------|---------------------------|---------------------------------------|-------------------------------------|-----------------|------------------------------------|
| Hit name | Species | Identified genes accession numbers | AA region/ AA–length (%) | E-value | Clone accession number |
| Angiopoietin–like 2 | Danio rerio | NP_001012502 | 12-213/510 (97%) | 1E-105 | DC997336 |
| Attractin | Mus musculus | NP_033860 | 1037–1228/1428 (84%) | 1E-103 | DC997227, DC997212 |
| Attractin–like 1 (Atrnl1) protein | Mus musculus | AAH50020 | 655-787/787 (73%) | 3E-43 | DC997188 |
| Beta-catenin | Carassius auratus | AAP94282 | 670–780/780 (97%) | 4E-61 | DC996928 |
| Beta–2–microglobulin precursor | Cyprinus carpio | Q03422 | 1-116/116(98%) | 9E-64 | DC997091 |
| Beta-2-microglobulin | Barbus intermedius | CAD44965 | 1-116/116 (91%) | 1E–59, 4E–60 | DC997439, DC997085 |
| C–type lectin domain family 3. member B | Homo sapiens | NP_003269 | 12-199/202 (53%) | 1E–55 | DC997874 |
| Cathepsin L | Sus scrofa | NP_999057 | 26-204/334 (53%) | 2E-44 | DC997269, DC997391, DC997140 |
| CD2 antigen (cytoplasmic tail) binding protein 2 | Danio rerio | NP_957255 XP_001341245 | 172–364/378 (65%) | 5E-66 | DC997518 |
| CD80–like protein | Oncorhynchus mykiss | CAG25516 | 201-349/372 (35%) | 2E-16 | DC997516 |
| Cell division cycle 42 (GTP binding protein 25 kDa) | Xenopustropicalis | NP_001017070 | 1-190/191 (99%) | 1E-106 | DC997410 |
| Class I helical cytokine receptor number 1 | Tetraodon nigroviridis | AAR25664 | 110-313/394 (78%) | 4E-92 | DC997764 |
| Claudin 14b | Takifugu rubripes | AAT64039 | 2-131/246 (49%) | 6E-29 | DC996974 |
| Cytokine inducible SH2–containing protein | Gallus gallus | NP_989957 | 1-168/249 (46%) | 2E-32 | DC997371 |
| Dipeptidyl peptidase 4 | Rattus norvegicus | NP_036921 | 411-590/767 (57%) | 1E-56 | DC996958 |
| Leukocyte immune–type receptor TS32.15 L1.2a1 | Ictalurus punctatus | ABI16039 | 128–296/483 (32%) | 1E-15 | DC997165 |
| Lymphocyte antigen 75 | Mus musculus | NP_038853 | 959–1130/ 1723 (39%) | 7E–23 | DC997698 |
| Lymphocyte antigen 75 variant | Homo sapiens | BAD92152 | 918–1115/1340 (36%) | 1E-26 | DC997636 |
| Glutamyl amino–peptidase (EAP) CD249 | Homo sapiens | Q07075 | 95–288/957 (65%) | 1E-68 | DC997775 |
| GTPase activating protein (SH3 domain) binding protein 1 | Danio rerio | AAH65323 | 1-169/477 (89%) | 6E-81 | DC997689 |
| Heat shock 70 kDa protein 5 (glucose- regulated protein) | Danio rerio | AAH52971 | 1-141/650(97%) | 1E-74 | DC997149 |
| HGF–regulated tyrosine kinase substrate | Mus musculus | CAM27049 | 615–741/767 (47%) | 3E-19 | DC996991 |
| Human immunodeficiency virus type I enhancer binding protein 1 | Homosapiens | NP_002105 | 2087–2227/2718 (56%) | 6E-40 | DC997885 |
| Inositol polyphosphate–5– phosphatase, isoform CRA_d | Homo sapiens | EAX07310 | 195–375/913 (60%) | 1E-56 | DC996986 |
| Insulin receptor substrate 2–B (IRS– 2–B) | Xenopus laevis | Q5RJW5 | 688–898/1077 (30%) | 2E-17 | DC997048 |
| Interferon gamma inducible protein 30 | Danio rerio | AAH83267 | 1-205/255(75%) | 8E-91 | DC997060 |
| Interferon regulatory factor 1 | Danio rerio | NP_991310 | 1-176/287 (70%) | 3E-62 | DC99768, DC997685 |
| Interferon regulatory factor 2 | Channa argus | ABK63484 | 228-328/328(45%) | $6E{-}10$ | DC996873 |
| Invariant chain like protein 2 | Cyprinus carpio | BAC53768 | 1-197/198 (84%) | 4E-85 | DC99764, DC997504 |
| Invariant chain like protein 2 | Cyprinus carpio | BAC53768 | 1-181/198 (99%) | 1E-100 | DC997791 |
| Legumain | Danio rerio | NP_999924 | 295-437/438 (84%) | 7E-62 | DC997010 |
| Legumain | Bos taurus | NP_776526 | 19–199/433(78%) | 2E-75 | DC997151 |
| Peptide transporter PEPT2 | Danio rerio | NP_001034917 | 1–186/719(90%), 521–670/719(83%) | 3E–94, 1E–65 | DC99717, DC997095 |

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| Platelet–derived growth factor alpha polypeptide | Danio rerio | NP_919407 | 1-106/195 (90%) | 4E-48 | DC996918 |
|-------------------------------------------------------------|-------------------------------|------------------------|-----------------------------------------|-----------------|-----------------------|
| Plexin C1 | Mus musculus | NP_061267 XP_290070 | 1364-1405/1574 (76%) | 1E-15 | DC997384 |
| Protein tyrosine kinase 2 beta | Danio rerio | NP_997735 | 701–906/1004 (80%), 1–133/1004 (88%) | 1E–89, 2E–64 | DC997593, DC997534 |
| RAS–related C3 botulinum substrate 1 | Danio rerio | NP_956065 | 1-129/192 (100%) | 1E-71 | DC997846 |
| Similar to Contactin 3 precursor | Danio rerio | XP_695624 | 62-263/697(37%) | 5E-23 | DC997089 |
| Similar to membrane–Toll–like receptor 5 | Danio rerio | XP_001343149 | 357-519/881 (40%) | 2E-18 | DC996968 |
| Small inducible cytokine SCYA105 | Paralabidochromis chilotes | AAO21207 | 1-87/105 (45%) | 3E-19 | DC996970 |
| Solute carrier family 3, member 2 | Danio rerio | AAH53236 | 1–180/180 (89%), 1–173/485 (90%) | 1E–90, 2E–87 | DC996849, DC997378 |
| Syndecan 4 | Danio rerio | NP_001041614 | 16-50/201 (80%) | 7E-10 | DC997001 |
| Transforming growth factor beta–1 precursor (TGF–beta–1) | Cyprinus carpio | Q9PTQ2 | 1-59/376 (81%) | 3E-17 | DC997847 |
| Transmembrane 4 superfamily member protein | Homo sapiens | EAW78874 | 1-163/202 (41%) | 4E-24 | DC996876 |
| Transmembrane and immunoglobulin domain 1 | Homo sapiens | NP_996663 XP_371037 | 60-184/262 (37%) | 5E-18 | DC997621 |
| TRPC4–associated protein A | Danio rerio | NP_571644 | 271-465/774(98%) | 2E-93 | DC997129 |
| Tumor necrosis factor receptor superfamily member8 | Gallus gallus | NP_989775 | 1-201/467(33%) | 3E-09 | DC997192 |
| Tumor necrosis factor, alpha–induced protein 8–like 3 | Homo sapiens | NP_997264 | 89-223/292 (70%) | 2E-47 | DC996858 |
| X–box binding protein 1 | Danio rerio | AAH66493 | 1-158/263 (94%) | 3E-65 | DC997144 |
| | | | | | |

B activating protein, interferon induced protein and cytokine receptors such as CXCR3, CXCR1, CRFB5 and TNFR member19 that were more frequent in gill library than in intestine library, which suggests the important role of gills as a first line of defense relying on innate and acquired immunity. This may be because the gill is very fragile and weak organ acting as a physical barrier easily to enter the forging substance from the surrounding environment. In addition, the outer environment surrounding gills is full of many kinds of pathogen, so gills become an important place for the antigen presenting molecules compared to intestine or other studies in carp ESTs libraries (Savan and Sakai 2002; Kono et al., 2004; Christoffels et al., 2006; Gonzalez et al., 2007). The results of gill and intestine cDNA libraries indicating the useful role of EST for identifying immune-related genes of fish as well as identifying novel genes.

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