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First Identification of a Chemotactic Receptor in an Invertebrate Species: Structural and Functional Characterization of Ciona intestinalis C3a Receptor

Daniela Melillo,* Georgia Sfyroera,‡ Rosaria De Santis,* Rita Graziano,† Rita Marino,† John D. Lambris,1‡ and Maria Rosaria Pinto1**

In mammals, the bioactive fragment C3a, released from C3 during complement activation, is a potent mediator of inflammatory reactions and exerts its functional activity through the specific binding to cell surface G protein-coupled seven-transmembrane receptors. Recently, we demonstrated a Ciona intestinalis C3a (CiC3a)-mediated chemotaxis of hemocytes in the deuterostome invertebrate Ciona intestinalis and suggested an important role for this molecule in inflammatory processes. In the present work, we have cloned and characterized the receptor molecule involved in the CiC3a-mediated chemotaxis and studied its expression profile. The sequence, encoding a 95,394 Da seven-transmembrane domain protein, shows the highest sequence homology with mammalian C3aRs. Northern blot analysis revealed that the CiC3aR is expressed abundantly in the heart and neural complex and to a lesser extent in the ovaries, hemocytes, and larvae. Three polyclonal Abs raised in rabbits against peptides corresponding to CiC3aR regions of the first and second extracellular loop and of the third intracellular loop react specifically in Western blotting with a single band of 98–102 kDa in hemocyte protein extracts. Immunostaining performed on circulating hemocytes with the three specific Abs revealed that CiC3aR is constitutively expressed only in hyaline and granular amoebocytes. In chemotaxis experiments, the Abs against the first and second extracellular loop inhibited directional migration of hemocytes toward the synthetic peptide reproducing the CiC3a C-terminal sequence, thus providing the compelling evidence that C. intestinalis expresses a functional C3aR homologous to the mammalian receptor. These findings further elucidate the evolutionary origin of the vertebrate complement-mediated proinflammatory process.

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In all vertebrates, local injuries to living tissues result in an inflammatory response, which represents an important first line of defense, operating in tight association with both the innate and adaptive immune responses. In this context, the complement system plays a pivotal role. In fact, its activation results in the generation of the bioactive fragments, C3a and C5a, which are potent mediators of inflammation (1, 2).

In particular, the anaphylatoxin C3a is a 74- to 78-aa peptide generated by proteolytic cleavage of the N-terminal end of the C3 α-chain during the activation of either the classical, lectin, or alternative pathways of the complement system (3, 4). Our knowledge of the functional and structural features of this anaphylatoxin comes mainly from studies of the mammalian C3a molecules, which possess various immunomodulatory and proinflammatory activities (2, 5, 6). Most of the biological effects of C3a, including the recruitment and activation of leukocytes and other effector cells at an inflammatory site, are elicited via specific binding to cell surface G protein-coupled seven-transmembrane receptor(s) (2, 7) belonging to the rhodopsin family. The C3a receptor (C3aR) has been cloned from different mammalian species (8–14), and its functional characterization has been conducted mainly in humans and mice (2, 7). A feature common to all the mammalian C3aRs is their broad pattern of tissue distribution, which includes lymphoid organs (8–14).

The most remarkable feature of mammalian C3aRs is the presence of an unusually large second extracellular loop (EL) of ~ 175 aa between the fourth and fifth transmembrane domains, exhibiting low sequence homology among species (7). The existence of this unique structural feature among the G protein-coupled receptors has suggested a special role for this loop in ligand binding. Indeed, the analysis of chimeric and deletion mutants of the human C3aR has established that as much as 65% of the second EL sequence (aa 198–308) can be removed without affecting C3a binding or the calcium response (15). This analysis has also suggested a two-site C3a-C3aR interaction model similar to that established for C5a-C5aR. A primary effector site on the receptor interacts with the C3a C-terminal sequence GLAR, whereas the anionic residues near the N and C termini of the C3aR second EL constitute a secondary interaction site with the cationic residues in the C-terminal helical region of C3a (15). More recently, it has been demonstrated that sulfation of Tyr174 of human C3aR is essential for C3a binding (16). Following ligand binding, C3aR is internalized within minutes, and the phosphorylation of cytosolic C-terminal Ser and Thr residues seems to play a crucial role in this process (17, 18).

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2 Abbreviations used in this paper: EL, extracellular loop; CiC3a, C. intestinalis C3a; JTT, Jones-Taylor-Thornton; IL, intracellular loop.
Whereas the mammalian inflammatory pathway of the complement system is well characterized at both the molecular and functional levels, little information is available about its structure, function, and evolution in nonmammalian vertebrates and invertebrates (19). Very recently, a trout cDNA has been cloned that encodes a 364-aa molecule, which shows a high degree of sequence homology and a strong phylogenetic relationship with mammalian C3αRs. However this C3α-like receptor was found to lack a significant portion of the large second EL characterizing the mammalian C3αRs (20). C3-like genes have been also identified in different species of invertebrates (21), and in limited instances, C3 mediated functions have been reported in these organisms (22–25).

Using the ascidian *Ciona intestinalis* as a model, we have previously shown the presence of an inflammatory effector pathway of the complement system in deuterostome invertebrates (26). In fact, we have demonstrated a specific chemotactic activity exerted on *Ciona* hemocytes by the cell-free hemolymph or the recombinant *Ciona* C3α. As is true for mammalian C3α, the chemotactic activity of *C. intestinalis* C3α (CiC3α) was localized to the C terminus, but the terminal Arg was not critical for the activity. The C3α-mediated chemotaxis was inhibited by pretreatment of the cells with pertussis toxin, suggesting that the receptor molecule mediating the chemotactic effect is G protein coupled (26).

To clarify the molecular mechanisms underlying the complement-mediated proinflammatory processes in invertebrates, we have now identified and sequenced a *CiC3αR* gene that exhibits a wide pattern of tissue expression. Three polyclonal Abs raised against peptides reproducing different portions of the deduced amino acid sequence of this receptor have been used in chemotaxis experiments, Western blot analysis, and immunostaining of circulating hemocytes. The results of these experiments demonstrate for the first time the presence and the functional activity of a C3αR in an invertebrate species.

**Materials and Methods**

**Hemocyte preparation**

Specimens of *C. intestinalis* were collected in the Gulf of Napoli and maintained in circulating seawater until use. The hemolymph was collected from the perivisceral cavity with a syringe.

To prepare hemocytes, hemolymph was immediately diluted 1/2 with ice-cold artificial seawater, Ca²⁺ and Mg²⁺-free, containing 10 mM EDTA (pH 7.4). Following centrifugation at 500 × g for 5 min at 4°C, hemocytes were resuspended in marine solution: 0.45 M NaCl, 26 mM MgCl₂, and 1 mM complex included the adult neural gland. The results of these experiments demonstrate for the first time the presence and the functional activity of a C3αR in an invertebrate species.

**RNA isolation**

Total RNA was prepared from hemocyte pellet, ovary, heart, neural complex, and larval stage of *Ciona* using the kit SV Total RNA isolation system (Promega), according to the manufacturer’s instruction. The neural complex included the adult *Ciona* single cerebral ganglion and the associated neural gland.

**RT-PCR amplification**

Oligo(dT)primed single-stranded cDNA was synthesized from hemocyte RNA with a RT-PCR kit (Invitrogen Life Technologies), according to the manufacturer’s instruction. The cDNA was PCR amplified using two specific primers (sense primer, 5'-GATTCAAAATCGCCGAACAGA-3', and antisense primer, 5'-ATGAAGCTCGGCAATGCGCATGA-3'), designed on the basis of the 381-nt gene model sequence cII010014062, Scaffold:455:55844–558678, of the *C. intestinalis* genome, version 1.0, provided by JGI (www.jgi.doe.gov/). The conditions used were as follows: 94°C for 15 s, 56°C for 30 s, and 72°C for 1 min. A single band of the expected size (333 bp) was detected. This product was cloned into the pCRII-TOPO vector (Invitrogen Life Technologies) and verified by sequencing.

**5'- and 3'-RACEs**

To determine sequences at the 5'- and 3'-ends of the *CiC3α* mRNA, RACE assays were conducted using the 5'- and 3'-RACE system (Invitrogen Life Technologies). To amplify the 3'-end, an oligo(T) containing adapter primer was used for the first-strand cDNA synthesis of hemocyte total RNA. The cDNA-adapter DNA was UV cross-linked at 120 mJ/cm², prehybridized in a buffer composed of 5× SSC, 5× Denhardt’s solution, 5 mM EDTA, 0.5% SDS, 200 μg/ml salmon sperm, and 50 mM sodium phosphate (pH 7.2), then hybridized for 16 h at 60°C in the same buffer containing the 843-bp 32P-labeled random-primed probe (28), corresponding to the 597–1439 region of the coding sequence. PCR analysis of genomic DNA indicated that this fragment, out of the several amplified, did not contain introns. At the end of the hybridization time, the membranes were briefly rinsed in 2× SSC with 0.1% SDS at room temperature and then washed in 2× SSC with 0.1% SDS for 15 min at room temperature and 15 min at 60°C. The blot was exposed in an autoradiography cassette to the x-ray BioMax MS film (Eastman Kodak) at −80°C.

**Southern blot analysis**

Genomic DNA was isolated from *Ciona* sperm according to standard protocols (27). The DNA (4 μg) was digested to completion with the restriction enzymes, EcoRI and HindIII, separated on a 0.8% agarose gel, and transferred to a Hybond-N⁺ nylon membrane (Amershams Biosciences) in 20× SSC. The transferred DNA was UV cross-linked at 120 mJ/cm², prehybridized in a buffer composed of 5× SSC, 5× Denhardt’s solution, 5 mM EDTA, 0.5% SDS, 200 μg/ml salmon sperm, and 50 mM sodium phosphate (pH 7.2), then hybridized for 16 h at 60°C in the same buffer containing the 843-bp 32P-labeled random-primed probe (28), corresponding to the 597–1439 region of the coding sequence. PCR analysis of genomic DNA indicated that this fragment, out of the several amplified, did not contain introns. At the end of the hybridization time, the membranes were briefly rinsed in 2× SSC with 0.1% SDS at room temperature and then washed in 2× SSC with 0.1% SDS for 15 min at room temperature and 15 min at 60°C. The blot was exposed in an autoradiography cassette to the x-ray BioMax MS film (Eastman Kodak) at −80°C.

**Determination and analysis of DNA sequences**

Sequences were determined for both strands using vector- or gene-specific primers. Sequence reactions were obtained with the BigDye Terminator Cycle Sequencing technology (Applied Biosystems) and purified by automatic separation using the Millipore Montage SEQ96 Sequencing Reaction Cleanup kit (Millipore) and a robotic station Biomek FX (Beckman Coulter). Products were analyzed on an Automated Capillary Electrophoresis Sequencer 3730 DNA Analyzer (Applied Biosystems).

cDNA sequence analysis was routinely performed by running a basic local alignment search tool of the GenBank database.

The deduced amino acid sequences of *CiC3α* and *C3α* receptors from different vertebrate species were aligned using the Clustal X 1.8 program (29), and the resulting alignment was corrected manually.

**Phylogenetic analysis**

To construct a phylogenetic tree, a data set including *CiC3α* and selected anaphylatoxin receptor amino acid sequences, aligned with the program Clustal X 1.8, was used. The *CiC3α* and *C3α* residues in the third cysteine loop and the elongation of the carboxyl-terminal tail were not considered in the data set. The phylogenetic relationships among related sequences were inferred by the maximum likelihood method implemented in PHY-ML software (30). The Jones-Taylor-Thornton (JTT) substitution model (31), with gamma-distributed rates, was selected among
alternative models (see Table I) using Tree-Puzzle software (32). The estimated value of the gamma distribution parameter $\alpha$ was 1.11 ± 0.08. The tree topology was reconstructed by applying the quartet puzzling maximum likelihood method implemented in Tree-Puzzle.

**Peptide synthesis and Ab production**

The 17-aa peptide, CiC3aR55–81 (HELPEPTQLANAEPP), the 17-aa peptide, CiC3aR228–244 (DPMYEEYQELYTDIYIG), and the 15-aa peptide, CiC3aR398–412 (LRRSLKKAARDRGTG), representing part of the CiC3aR-deduced amino acid sequence and located in the area of the presumptive first and second ELs and the third intracellular loop (IL), respectively, were synthesized using an Applied Biosystems 430A peptide synthesizer as described previously (33). The peptide was then coupled to OVA by the glutaraldehyde method and used for the immunization of rabbits to produce the anti-CiC3aR55–81 (anti-EL1) (Primin)-, anti-CiC3aR228–244 (anti-EL2) (Primm)-, and anti-CiC3aR398–412 (anti-IL-3) (Cocalico Biologics)-specific Abs.

Columns of each peptide-specific Abs were purified by affinity chromatography using the corresponding synthetic peptides coupled to cyanogen bromide-activated Sepharose 4B (Amersham Biosciences).

**Western blot analysis**

Hemocytes, collected as indicated above, were resuspended at 1.6 × 10^6/ml in sample buffer (60 mM Tris with 2% SDS, 10% 2-ME, 10% glycerol, and 0.01% bromophenol blue (pH 6.8)) containing the protease inhibitor mixture Mini Complete (Roche Applied Science). Samples were homogenized by hand in a Potter-Elevehjem homogenizer with a Teflon pestle, boiled for 10 min, and centrifuged at room temperature for 30 min at 13,400 rpm in a Mini Spin centrifuge (Eppendorf). The supernatant was recovered and stored as aliquots at −20°C. The reactivity of Ciona hemo-ocyte proteins with the anti-EL1, anti-EL2, and anti-IL3 Abs was assessed by Western blotting analysis: 80 μg of protein, corresponding to 4.8 × 10^9 hemocytes, was loaded onto a 7.5% SDS-PAGE gel under reducing conditions. The proteins were then electroblotted onto a nitrocellulose membrane (Amersham Biosciences), and the CiC3aR was detected using 10 μg/ml of affinity-purified anti-EL1, anti-EL2, or anti-IL3 Ab. Non-specific binding to the membrane was prevented by incubation in blocking buffer (PBS containing 5% milk and 0.05% Tween 20). Bound polyclonal Abs were detected with a protein A-HRP conjugate (Bio-Rad), according to the manufacturer’s instructions. All the incubation steps were performed for 1 h at room temperature, and the membranes were washed with PBS containing Tween 20 (0.05%) between each incubation step. The proteins were detected using the ECL Plus kit (Amersham Biosciences). Protein determination assays were performed using a BCA kit (Sigma-Aldrich).

**Immunostaining of circulating hemocytes**

Hemocytes, collected as described above, were fixed for 30 min at room temperature in 4% paraformaldehyde with 0.5 M NaCl and 0.1 M MOPS (pH 7.5). At the end of the incubation time, 100-μl samples with ~1.5 × 10^10 cells/ml were applied onto Superfrost Plus slides (VWR International) by cytocentrifugation (Cytoprep: Wescor) at room temperature for 6 min at 400 rpm. Following three washings of 5 min each with PBS containing 0.1% Tween 20 (PBT), slides were incubated for 20 min with 0.3% (v/v) hydrogen peroxide in PBT and then washed again in PBT for 5 min, in PBT containing 10% normal goat serum for 10 min, and three times for 5 min in PBT. The incubation with the affinity-purified anti-EL1, anti-EL2, or anti-IL3 was performed for 1 h at room temperature in a humidified chamber at protein concentrations of 15, 45, and 30 μg/ml, respectively, in PBT containing 1% BSA. The slides were immunostained using the Vectastain Elite ABC kit (Vector Laboratories) and the Sigma Fast Dab set (Sigma-Aldrich), according to the manufacturer’s instructions. Controls were run in parallel with preimmune rabbit IgG at the same concentrations. Staining was observed under a Zeiss Axioskop microscope with Nomarski optics. Immunofluorescence microscopy was performed using the same proce- dure as described above, omitting the hydrogen peroxide treatment. The immunostaining was conducted for 1 h with the affinity-purified fluorescein-labeled goat anti-rabbit IgG (Vector Laboratories) diluted 1:250, followed by two 30-min washes and an overnight wash with PBS.

**Chemotaxis assay**

Chemotaxis experiments with *C. intestinalis* hemocytes were performed in micro Blind Well Chambers, model BW200L (Neuromax). A single Blind Well Chamber (13-mm diameter, 5.0-μm pore size; Whatman) was used to separate the upper and lower wells. The lower well was then loaded with 200 μl of 10 μM chemotaxant in the form of the carboxyl-terminal synthetic peptide CiC3a436–455 (55844–58678) (IARLNSGTRQRVGRQ). Hemocytes (300 μl, 2.5 × 10^6 cells/ml) were added to the upper cell well and allowed to migrate for 2 h at 20°C. At the end of the incubation time, the fluid in the upper chamber and the nonmigrated cells on the top surface of the filter were gently wiped off with a cotton swab. The number of cells that had migrated to the lower well was estimated by counting 16 randomly chosen fields in a hemocytometer, and their percentage of the total number of cells added to the chemotaxis chamber was calculated. Chemotaxis control experiments were conducted with two unrelated peptides substituting the specific chemotactic agents in the lower cell well.

For each inhibition experiment using anti-EL1, anti-EL2, or anti-IL3 Abs, hemocytes were resuspended in Eppendorf test tubes at 2.5 × 10^6 cells/ml in marine solution containing the Abs in a concentration range from 0.5 to 15 μg/ml. Following a 20-min incubation at 20°C, 300-μl samples were moved to the upper well of the chemotaxis chambers, and the chemotaxis assay was run and evaluated as described above. Control che- motaxis inhibition experiments were run in parallel with the same amounts of the corresponding preimmune rabbit IgG or an affinity-purified Ab of unrelated specificity (anti-trout factor H Abs). The results of inhibition experi- ments are presented as the percentage of inhibition of cell migration and are expressed as the mean ± SD of five independent experiments conducted in duplicate. A further control was also performed using the Ab-related peptides: the synthetic peptides used to raise rabbit Abs against the first and the second ELs were used in competition experiments in which the hemocytes were incubated with 5 μg/ml of the Ab in the presence of 0.06 μM of the corresponding peptide, then used in chemotaxis experiments.

**Results**

**Identification and sequence analysis of the CiC3aR gene**

To identify the CiC3aR gene, we conducted an in-depth search of the *C. intestinalis* genome web site ([http://genome.jigspf.org/ Ciona4/Ciona4.home.html]) for sequences showing significant homology to mammalian C3aRs. We found only one homologous gene model (ci0100141062), located at the 3′-end of the scaffold 455 (55844–58678) and including three exons, for total of 381 bp. The analysis of the deduced amino acid sequence indicated that it includes the second, third, and fourth transmembrane domains, as well as a few amino acid residues of the presumptive second EL of a G protein-coupled seven-transmembrane receptor.

To verify the presence in *Ciona* hemocytes of transcripts repro- ducing the identified sequence, we prepared total RNA and syn- thesized single-strand cDNA, which was then PCR amplified using two specific primers designed on the basis of the gene model se- quence. A single band of the predicted size (333 bp) was detected. This product was cloned, sequenced, and found to correspond to the gene model sequence. Because our search of the *Ciona* genome for scaffolds, including the 3′-terminal portion of this sequence, did not produce any results, we used 3′-RACE to determine the sequence of the 3′-end of the presumptive *Ciona* C3aR gene. We obtained a 2894-bp amplification product that allowed us to complete the 3′-end. A single cycle of 5′-RACE was conducted to

### Table I. Comparison of maximum log-likelihood values for the anaphylatoxin receptor data set under different models of evolution

| Model      | Rate   | Maximum Likelihood (logL) |
|------------|--------|---------------------------|
| Blosum     | Uniform| −10,773.58                |
| Blosum     | Gamma  | −10,577.95                |
| Dayhoff    | Uniform| −10,791.83                |
| Dayhoff    | Gamma  | −10,537.88                |
| JTT        | Uniform| −10,694.96                |
| JTT        | Gamma  | −10,448.11                |
| Variable 1 | Uniform| −10,912.42                |
| Variable 1 | Gamma  | −10,633.35                |

*Maximum likelihood estimates of anaphylatoxin receptor phylogeny under different models. Entries are: maximum log-likelihoods under the models in the first column, considering the rate of evolution of the sites distributed as a uniform or gamma distribution as indicated in the table.*
cover the 5'‐end. The total transcript was 3071 bp (European Molecular Biology Laboratory database accession number AJ966353 located online (www.ebi.ac.uk/) and included a 52‐bp 5′‐untranslated region (UTR), a 2544‐bp coding region with the initial methionine codon in the context of a canonical Kozac motif, and a 475‐bp 3′‐UTR, with two possible consecutive polyadenylation signals, at nt 3030 and 3037, and a poly(A) tail. A basic local alignment search tool analysis of this sequence in the Ciona genome, version 1.0, revealed that the CiC3aR regions between the nt 1–750 and 1673–3053 were present in scaffolds 455 and 438, respectively, whereas the region between nt 751–1672 was not found in any scaffold. A further analysis conducted in the National Center for Biotechnology Information database revealed the presence of a 1536‐bp C. intestinalis cDNA clone (ciht004g05; accession number AK174732) (34) corresponding to the 1534–3071 region of the presumptive CiC3aR.

To determine whether CiC3aR was present as a single copy or in the form of multiple copies per haploid genome, Southern blot analysis of genomic DNA from three animals’ sperm was performed. Each sample was digested with two endonucleases (EcoRI and HindIII), and the blot was hybridized with an 843‐bp probe corresponding to the fragment 597‐1439 of the coding sequence. Only a single hybridizing band was detected for the DNA of animals 1 and 3, allowing us to conclude that this gene is present as a single copy per haploid genome. Digestion with the same panel of restriction enzymes of the DNA from animal 2 revealed three bands per lane, reflecting the presence of allelic polymorphic sites within the gene (Fig. 1).

The alignment between the CiC3aR transcript sequence and the Ciona genome assembly, version 2.0 (http://genome.jgi-psf.org/Cioin2/Cioin2.home.html), released in March 2005, indicated that the entire CiC3aR sequence was located on scaffold 187 (reverse strand) between nt 27,219 and 39,709. According to this alignment, 11 exons contribute to the gene structure (Fig. 2). The first exon encodes the 52 bp of 5′‐UTR and the first 96 bp of the open reading frame, which spans from exon 1 to exon 10; the 475 bp of the 3′‐UTR are encoded by exons 10 and 11. This genomic organization is significantly different from the mammalian C3aR gene structure, which includes only two exons encoding the entire transcript, with the whole coding sequence contained on a single exon (12, 14).

Several tissues from adult animals and the larval stage were analyzed by Northern blot to evaluate the expression of the CiC3aR gene. A 747‐bp 32P‐labeled cDNA probe detected a single band of ~3.1 kb in all tissues analyzed (Fig. 3), thus confirming the size of the total transcript (3071 bp) obtained by sequence analysis of genomic DNA from three animals’ sperm was performed. The Ciona2/Cioin2 home.html), released in March 2005, indicated that the entire CiC3aR sequence was located on scaffold 187 (reverse strand) between nt 27,219 and 39,709. According to this alignment, 11 exons contribute to the gene structure (Fig. 2). The first exon encodes the 52 bp of 5′-UTR and the first 96 bp of the open reading frame, which spans from exon 1 to exon 10; the 475 bp of the 3′-UTR are encoded by exons 10 and 11. This genomic organization is significantly different from the mammalian C3aR gene structure, which includes only two exons encoding the entire transcript, with the whole coding sequence contained on a single exon (12, 14).

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FIGURE 1. Southern blot analysis of CiC3aR. Four micrograms of sperm DNA isolated from three different animals was digested with EcoRI and HindIII endonucleases, run on an agarose gel, and transferred to a nylon membrane. Hybridization was conducted with a probe corresponding to the fragment 597-1439. Values on the left of the blot represent molecular mass markers in kilobases.

FIGURE 2. Genomic organization of the CiC3aR gene. Shown is a schematic representation of the gene encoding the CiC3aR. Exons are indicated by closed boxes. The lines connecting boxes represent the intervening introns. Scale bar: 1 kb.

FIGURE 3. Tissue distribution of the CiC3aR by Northern blot analysis. Total RNA (6 μg/lane) from the indicated C. intestinalis tissues was electrophoresed, blotted onto a nylon membrane, and hybridized with a 32P-labeled 747-bp CiC3aR cDNA probe. The CiC3aR transcript was detected as a single band of ~3.1 kb in all samples. The corresponding ethidium bromide-stained 28S bands are shown in the lower part of the figure.

FIGURE 4. Schematic representation of the CiC3aR. The seven-trans-membrane domains of the CiC3aR are depicted as cylinders connected by ribbons representing the IL and EL.
The open reading frame of 2,544 nt encodes a protein of 848 aa with a deduced molecular mass of 95,394 Da. Hydropathy analysis (www.ch.embnet.org/software/TMPRED_form.html) of the predicted amino acid sequence of CiC3aR suggested the presence of seven-transmembrane domains (as schematically represented in Fig. 4), a feature common to G protein-coupled receptors belonging to the rhodopsin family. The alignment of CiC3aR with homologous sequences from other vertebrate species (Fig. 5) revealed a 276-aa insertion in the third cytoplasmic loop and a 105-aa elongation of the cytoplasmic tail. Not taking into account these regions, the Ciona sequence preceding the insertion shares ~35% similarity with the other species, while the similarity is ~45% for the region following the insertion.

The long hydrophilic region between the fourth and fifth transmembrane domain, EL2, a common feature of all C3a receptors, consisted of 162 aa in Ciona. Five of the seven identified N-glycosylation sequons (www.cbs.dtu.dk/services/NetNGlyc/) were localized to this loop.

Scanning the CiC3aR sequence for putative phosphorylation sites (35) revealed 43 serine, 15 threonine, and 11 tyrosine residues. Interestingly, the majority of serine (39 of 43) and threonine (12 of 15) residues were located in the third IL or the cytoplasmic tail. In the same regions, six sequences reproducing substrate motives of protein kinase C, (R/K1–3,X2–0)S*/T*-(X2–0,R/K1–3) (36), were found.

A phylogenetic tree was constructed based on the alignment of CiC3aR with vertebrate C3aR and C5aR sequences. As shown in Fig. 6, CiC3aR did not cluster with any of the C3aR or C5aR clades, and its position suggests that it is the most basal of the anaphylatoxin receptors.

Western blot analysis of circulating hemocytes
To assess the presence of a CiC3aR gene product in circulating hemocytes of Ciona, we examined the reactivity of the three specific Abs with hemocyte protein extracts by Western blotting. The

FIGURE 5. Alignment of the CiC3aR amino acid sequence with C3aR from other species. The deduced amino acid sequence of CiC3aR was aligned with rat, human, and chicken C3aR sequences, using the Clustal X (1.8) program, with adjustments made by hand. Identical residues in all sequences are shown in bold. The seven-putative transmembrane (TM) domains are overlined. ELs and ILs and the cytoplasmic tail are indicated by EL, IL, and CT, respectively. The three underlined sequences correspond to the synthetic peptides CiC3aR65–81, CiC3aR228–244, and CiC3aR398–412 used as Ags to produce specific anti-CiC3aR Abs. Accession numbers: Ciona intestinalis, A966353; Rattus rattus, AAC40071; Homo sapiens, CA97504; and Gallus gallus, CAG32407.
three affinity-purified Abs, anti-EL1, anti-EL2, and anti-IL3, reacted with the same protein band, which exhibited an apparent molecular mass of 98–102 kDa under reducing conditions (Fig. 7, A–C). The electrophoretic pattern of the hemocyte protein extract after Coomassie brilliant blue staining is shown in Fig. 7D. Glycosylation or other posttranslational modifications could account for the slight difference between the deduced (95,394 Da) and the SDS-PAGE-calculated (~100 kDa) molecular size of CiC3aR.

Hemocyte immunostaining

To determine which blood cell types express CiC3aR, circulating hemocytes were immunostained with the affinity-purified anti-EL1, anti-EL2, and anti-IL3 Abs. Staining of hemocytes revealed identical expression patterns for the three Abs. A clear staining pattern was found in the majority of hemocytes identified as granular and hyaline amoebocytes, according to the classification scheme of Rowley et al. (37). In Fig. 8A, immunostaining performed with the affinity-purified anti-IL3 Ab and the DAB substrate is shown. The cytoplasm of the granular and hyaline amoebocytes is characterized by the presence of granules that often appeared intensely stained. The cell types classified as vacuolated cells were never stained.

To extend the analysis of CiC3aR expression pattern, immunofluorescence experiments were conducted using the affinity-purified Abs and a fluorescein-conjugated secondary Ab. These experiments clearly indicated that CiC3aR is expressed not only in the cytoplasm but also on the cell surface of both granular and hyaline amoebocytes (Figs. 8, C–F).

Inhibition of hemocyte chemotaxis by specific Abs

To provide conclusive evidence that the identified and sequenced presumptive C3aR gene encoded the CiC3aR, we performed chemotaxis experiments using the three polyclonal Abs, anti-EL1, anti-EL2, and anti-IL3, raised against the synthetic peptides CiC3aR65–81, CiC3aR228–244, and CiC3aR398–412, representing deduced amino acid sequences in the first and second ELs and the third IL, respectively. We tested the ability of the affinity-purified Abs against the extracellular portions to inhibit the hemocyte chemotaxis elicited by the C3a59–76 synthetic peptide. We previously demonstrated that this peptide representing the CiC3aR65–81 terminus is able to promote specific hemocyte chemotaxis in a dose-dependent manner (26). In these experiments, we used as negative control the Ab directed against the cytoplasmic loop. Preincubation of the hemocytes with the anti-EL Abs (anti-EL1 and anti-EL2) almost completely (>90%) inhibited their migration to the lower well of the chemotaxis chamber containing the chemotractant peptide (Fig. 9). In contrast, preincubation of the hemocytes with the anti-cytoplasmic loop Ab (anti-IL3) had no inhibitory effect. The anti-EL1 and anti-EL2 Abs inhibited chemotaxis in a dose-dependent manner and produced 50% inhibition at concentrations of 1.2 and 3.7 μg/ml, respectively. Control experiments were run in parallel using either the preimmune polyclonal rabbit IgG or an affinity-purified polyclonal anti-trout factor H; both had no inhibitory effect. A further control was performed using the Ab-related peptides in competition experiments. When the hemocytes were incubated with the Ab in the presence of the corresponding peptides, the inhibitory activity was completely abolished, thus demonstrating the specificity of the Ab recognition (data not shown).
The chemotaxis experiments also allowed us to identify the cell types involved in the chemotaxis process. Microscopic analysis of the migrated hemocytes revealed that the chemoattracted cell types were hyaline and granular amoebocytes and univacuolar refractile granulocytes. Small hemocytes that could not be classified were also recovered in the lower chemotaxis chamber.

Discussion

In a previous paper (26), we demonstrated a chemotactic activity elicited by C3a in Ciona intestinalis, clearly indicating the presence of a complement-mediated inflammatory pathway in deuterostome invertebrates. Pertussis toxin inhibited this chemotactic activity, suggesting that the receptor molecule mediating the chemotactic effect is G protein coupled (26). However, the nature of the receptor molecule mediating this chemotaxis was unknown. In the present study, we report the primary structure of a transcript homologous to the mammalian C3aRs and demonstrate that its product is the membrane molecule interacting with the Ciona C3a fragment and mediating hemocyte chemotaxis. This information is relevant from an evolutionary point of view because of the phylogenetic position of the ascidians at the base of the vertebrate lineage.

Our Northern blot analysis of the transcripts showed that the CiC3aR is constitutively expressed in circulating hemocytes and the ovary, heart, and neural complex of the adult animals, as well as in the larval stage. The detection of a single band of ~3.1 kb in all tissues examined allows us to exclude the presence of different transcripts generated as a result of alternative RNA splicing. The wide CiC3aR distribution pattern is in agreement with those of mammals (8, 10–13) and fish (20) and is consistent with the role of C3a as mediator in inflammatory reactions.

The domain structure analysis of the deduced amino acid sequence indicated that CiC3aR, like the mammalian counterparts, is a G protein-coupled seven-transmembrane receptor. From the alignment shown in Fig. 5, it is evident that in Ciona the most relevant differences from other C3aRs involve the carboxyl-terminal tail and the third cytoplasmic loop, which are respectively 105 and 276 aa residues longer than the corresponding regions of vertebrate C3aRs. Although different in size, the C-terminal end of the Ciona cytosolic tail, like those in mammals, contains many serine and threonine residues (eight residues in the segment 808–819) that represent tentative phosphorylation sites. A further difference between Ciona and the mammalian C3aRs is related to the extracellular amino-terminal sequence, which in Ciona is shorter and lacks the tentative N-glycosylation site. In mammals, this site, together with another glycosylation site in the second EL, is thought to be responsible for the carbohydrate content, accounting for the difference between the deduced (~54 kDa) and the SDS-PAGE (83–114 kDa) estimated molecular size (7). In Ciona, the difference between the electrophoretic mobility (98–102 kDa) and the deduced molecular mass (95,394 Da) is small, even though five presumptive N-glycosylation sites have been identified, all within the second EL.

FIGURE 8. Immunostaining of C. intestinalis hemocytes. Circulating hemocytes were tested for the presence of CiC3aR with the affinity purified anti-EL1, anti-EL2, or anti-IL3 Abs. A, The immunostaining reaction performed with the affinity-purified anti-IL3 Ab, using diaminobenzidine as a substrate. Staining revealed the intracellular localization of CiC3aR Ags in hyaline amoebocytes and granular amoebocytes. No labeling was detectable in morula cells and univacuolar refractile granulocytes. B, Control reaction with the corresponding preimmune IgG. The immunofluorescence staining performed with the affinity-purified anti-IL3 Ab, in combination with an FITC-conjugated secondary Ab, indicates that CiC3aR is expressed in the cytoplasm and on the cell surface of granular and hyaline amoebocytes (C and E). D and F, The same fields with Nomarski optics. G and H, The fluorescence and Nomarski optic field of the control conducted with the corresponding preimmune Ab. HA, hyaline amoebocyte; GA, granular amoebocyte; MC, morula cell; URG, univacuolar refractile granulocyte. Scale bar, 10 μm.
The most distinctive feature of all the C3aRs is the length of the second EL, which is generally composed of ~170 residues, with only two exceptions: one of the two isoforms in the guinea pig, generated by alternative splicing (35 aa residues shorter) (13), and that in the trout C3aR, which is only 36 residues long (20). The length of the second EL of CiC3aR, 162 aa residues, is comparable to those in mammals. This loop has been shown to be critical for ligand binding involving both sites of the proposed two-site interaction model (15). In Ciona, the situation may be quite different; in fact, although many charged residues are present at the N and C termini of the second EL, there is no net negative charge, which has been suggested to be important in the interaction in mammals. However, these differences are not surprising because the CiC3aR ligand, i.e., CiC3a, also differs from its mammalian homologs (26). These differences at both the ligand and receptor levels could account for a different model of interaction.

To approach this topic, we explored the functional activity of CiC3aR by means of chemotaxis inhibition experiments using Abs against sequences in the first and second extracellular or the third IL. High levels of inhibition of the chemotactic activity (>90%) were achieved only with Abs against the ELs of the receptor. As expected, the Ab against the third IL, while not affecting the chemotactic activity, was equally efficient in Western blotting and hemocyte immunostaining. Our results do not allow us to distinguish between the contributions of the first and second ELs to the functional activity. In fact, the Ab inhibitory activity could have resulted either from steric hindrance at the interaction site(s) or by its direct engagement.

A comparative analysis of the sequence of the first EL of the Ciona receptor with those of mammals revealed interesting differences: it is longer and shows a net negative charge that in mammals is a feature of the N and C termini of the second EL. This finding could suggest a specific role for the first EL in the functional activity of Ciona.

The phylogenetic analysis we have conducted indicates that CiC3aR sequence does not cluster with any of the vertebrate C3aR and C5aR clades and supports the hypothesis that, if C3aR and C5aR genes originated from a common ancestor, this event occurred after the emergence of the urochordates. In agreement with this hypothesis is the absence from the Ciona genome of both C5 and a C5aR (38).

In the present study, we have also determined which hemocyte types in the circulating hemolymph express the C3aR. In fact, the immunostaining analysis performed with the anti-CiC3aR Abs showed a clear labeling only of hyaline and granular amoebocytes, which play a crucial role in complement-mediated inflammatory reactions being able to synthesize C3aR as well as C3, the precursor of C3a. In addition to these cell types, we also saw migration of the univacuolar refractile granulocytes, which do not express C3aR, toward the chemotaxtractant in chemotaxis experiments. This cell type has also been shown to be involved in local inflammatory reactions, with its concentration being significantly increased in tunic sections of LPS-injected Ciona (26). This behavior, in the absence of CiC3aR expression, could be explained by a cooperative effect: amoebocytes, the direct target of C3a, could, in turn, produce a mediator that can activate the univacuolar refractile granulocytes. This situation is somewhat reminiscent of the C3aR activity described by Daffern et al. (39) in humans. They demonstrated that eosinophils are directly activated by C3a, whereas neutrophils are only indirectly stimulated by C3a in the presence of eosinophils.

Taken together, our findings provide evidence for the first time of the presence of a functional C3aR in an invertebrate species. The identification in C. intestinalis of both the partners participating in the complement-mediated inflammatory response provides the impetus for further investigating the C3a-C3aR interaction at both the molecular and cellular levels.

In this context, a point that we consider relevant concerns the possibility of unraveling the mechanisms that regulate the cross-talk among various hemocyte types during the progression of the inflammatory process. Other points of interest pertain to the identification of the amino acid residues involved in the ligand-receptor recognition and binding, as well as the study of signal transduction and internalization. All this information, in a species at the crossroads between the invertebrate and vertebrate lineages (40), could help to shed light on the evolution of the inflammatory reaction.

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Disclosures
The authors have no financial conflict of interest.

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