Changes in reflectin protein phosphorylation are associated with dynamic iridescence in squid

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Manuscript information: Number of text pages; 21

Number of figures; 6

Abbreviations: ACh, acetylcholine; mAChRs, muscarinic acetylcholine receptors; DTT, dithiothreitol; HPLC, high performance liquid chromatography; IEF, isoelectric focusing; PVDF, polyvinylidene fluoride; HRP, horseradish peroxidase; MALDI-TOF MS, matrix-assisted laser desorption time-of-flight mass spectroscopy; RM, reflectin motif; PKC, protein kinase C
Abstract

Many cephalopods exhibit remarkable dermal iridescence, a component of their complex, dynamic camouflage and communication. In the species *Euprymna scolopes*, the light-organ iridescence is static and is due to reflectin protein-based platelets assembled into lamellar thin-film reflectors called iridosomes, contained within iridescent cells called iridocytes. Squid in the family Loliginidae appear to be unique in that the dermis possesses a dynamic iridescent component, with reflective, colored structures that are assembled and disassembled under the control of the muscarinic cholinergic system and the associated neurotransmitter acetylcholine (Mathger *et al.* 2004). Here we present the sequences and characterization of three new members of the reflectin family associated with the dynamically changeable iridescence in *Loligo* and not found in static *Euprymna* iridophores. In addition, we show that application of genistein, a protein tyrosine kinase inhibitor, suppresses acetylcholine- and calcium-induced iridescence in *Loligo*. We further demonstrate that two of these novel reflectins are extensively phosphorylated in concert with the activation of iridescence by exogenous acetylcholine. This phosphorylation and the correlated iridescence can be blocked with genistein. Our results suggest that tyrosine phosphorylation of reflectin proteins is involved in the regulation of dynamic iridescence in *Loligo*.
**Introduction**

Structural coloration, or iridescence, is widespread across metazoa, from insects and marine invertebrates to fishes and birds (Denton 1970, Mathger et al. 2003, Osorio and Ham 2002, Vukusic and Sambles 2003). Reflective cells, called iridocytes, typically contain stacks of platelets, or iridosomes, composed of different high refractive index materials in different organisms. These stacks of platelets function as iridescent Bragg reflectors (Cloney and Brocco 1983, Huxley 1968, Land 1972, Mirow 1972). The iridosomes in fish, amphibians and reptiles contain oblong, extracellular purine crystals nested in spaces in reticulated cells (Menter *et al.* 1979, Mathger *et al.* 2003, Rohrlich and Rubin 1975, Kobelt and Linsenmair 1986), whereas in insects, layers of microstructured cuticular chitin serve the same optical function (Vigneron *et al.* 2005). In organisms such as lizards, iridescence can be modulated with mobile chromatophore cells acting as shutters, but the iridescent structures themselves remain immobile (Carlton 1903). Uniquely, loliginid squid can actuate the structures responsible for iridescence, turning the optical effect off and on under the control of muscarinic acetylcholine receptors (mAChRs). Accordingly, exogenous acetylcholine and calcium ionophores induce iridescence in these species (Mathger *et al.* 2004) (Fig. 1). We know of no other organisms in which the iridescent structures themselves are assembled and disassembled under fast control, rather than slightly modulated or shuttered (Mathger *et al.* 2003).

In contrast to the purine crystals found in vertebrates and layered chitin found in insects, squid iridophore platelets are proteinaceous. Crookes and colleagues characterized the proteinaceous constituents of the static light-organ iridosomes of the bobtail squid *Euprymna scolopes*, and named this group of proteins ‘reflectins’ (Crookes *et al.* 2004). Subsequent work has shown that the dynamic iridosomes of the loliginids also are proteinaceous, and that their dynamic be-
Behavior is regulated by acetylcholine-dependent muscarinic-type G-protein-coupled receptors (Mathger et al. 2004). Activation of the loliginid iridophore layer by the neurotransmitter acetylcholine (ACh) leads to a dramatic increase in reflectance. Because nicotinic-type receptors that characterize the neuromuscular junction do not affect dynamic iridescence, this optical change is not attributed to muscular contraction, as is the case for chromatophore signaling in squid (Mathger et al. 2004). In addition, isolated reflectin proteins exhibit unusual solubility and self-association properties (Kramer et al. 2007). These data all suggest that the proteinaceous nature of iridosomes in loliginids may contribute to their dynamic ability to modulate their iridescent effects. To better understand the biochemical and biophysical bases of dynamic iridescence in loliginids, we have sought to characterize the protein targets of the muscarinic ACh signal transduction cascade as well as the downstream effects on reflectins resulting from this cascade.

Here we report that ACh-mediated signaling induces changes in the phosphorylation state of Loligo reflectin proteins, and that phosphorylation is concomitant with the dynamic changes in the optical properties of Loligo skin. These changes in phosphorylation can be inhibited with genistein, a potent kinase inhibitor. We hypothesize that the mechanism leading to reflectin aggregation and increased skin iridescence may be linked to protein charge neutralization via addition of phosphates upon ACh activation, suggesting a biochemical and biophysical mechanism for dynamic iridescence in loliginid squid. Although it has long been known that post-translational protein modifications can lead to aggregation, opacity and disease in ocular and neuronal tissues (e.g. Takemoto and Boyle 2000, Haass and Selkoe 1993), this is the first suggestion of a functional and reversible post-translational modification leading to aggregation and the consequent control of an important optical function.
Materials and Methods

Biological material

Specimens of the Longfin Inshore Squid, *Loligo pealeii*, collected from coastal waters at Woods Hole, MA were used for all experiments. *Escherichia coli* strain TOP10 (Invitrogen, Carlsbad, CA) cells were used for all recombinant DNA manipulations.

Protein preparation

Freshly isolated and physically responsive iridophore layers from *L. pealeii* were homogenized by mortar and pestle in liquid nitrogen. Tissue homogenates were resuspended in suspension buffer (phosphate buffered saline containing 1mM dithiothreitol (DTT), Phosphatase Inhibitor Cocktails I and II (EMD Bioscience, San Diego, CA) and Complete protease inhibitor (Roche, Indianapolis, IN), and then centrifuged at 13,500 rpm for 15min at 4°C in a F45-30-11 rotor (Eppendorf, Westbury, NY). The resulting pellet containing the reflectin proteins was resuspended and washed with suspension buffer to remove any residual soluble proteins. This crude reflectin extract was further processed by dissolving the pellet in Buffer A (8M urea, 2mM DTT, Complete protease inhibitor and 30mM Z-(N-morpholino) ethanesulfonic acid (MES)-NaOH pH 5.5) and centrifuging as above. The residual insoluble material was hydrolyzed with 6N hydrochloric acid (110°C; 16 h) and analyzed with a Beckman System 6300 amino acid analyzer with a cation exchange column, HiTrapSP, connected to ÄKTA purifier (GE Healthcare, NY) with 0-50% of Buffer B (8M urea, 2mM DTT, Complete protease inhibitor (Roche), 2M NaCl and 30mM MES-NaOH pH 5.5). Fractions containing Reflectin A1 and A2 were identified by Western blotting with an antibody against the reflectin first purified and identified from *Euprymna scolopes* (Crookes *et al.* 2004) and collected for further purification. Pooled fractions were concentrated by centrifugation (Ultracon; Millipore Corp., Billerica, MA) and further puri-
fied with a C18 reverse phase column equilibrated with Buffer C (0.1% TFA in water) and a 30-50% gradient of Buffer D (0.1% TFA in 95% acetonitrile). The resulting purified proteins were lyophilized and stored at -80°C until use. Purification was monitored by sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) with a 10-20% acrylamide gradient gel. Prior to SDS-PAGE, the proteins were dissolved in SDS-PAGE sample buffer (Invitrogen). Protein concentration was measured using Bio-Rad Protein Assay Kit (Bio-rad) or 2-D Quant Kit (GE Healthcare).

To prepare samples for quantification of phosphorylation level, powdered frozen iridophore layers were directly dissolved in SDS-PAGE sample buffer containing DTT, protease inhibitor and phosphatase inhibitors. Following incubation at 98°C for 10 min, the samples were centrifuged as mentioned above, and the resulting supernatants were recovered for analysis.

**Preparation of RNA and cDNA**

Messenger RNA (mRNA) was purified from the freshly isolated iridophore layer of *L. pealeii*. Iridophore layers were dissected from live squid and stored in RNAlater (Qiagen, Valencia, CA) at 4 or -80 °C until use. Total RNA was isolated using TriZol (Invitrogen) followed by RNeasy (Qiagen) for further purification. First-strand cDNA was synthesized using SuperScript™ III RT (Invitrogen) with an oligo(dT)20 primer. BD SMART RACE cDNA amplification kit (Clontech, Mountain View, CA) was used to prepare rapid amplification of complementary DNA ends (RACE)-ready cDNA. All procedures were performed according to the manufacturers’ instructions.

**Amino acid sequencing and gene cloning**

To clone Reflectin A2, several primers were designed based on the consensus sequence information obtained from *E. scolopes* and *Loligo forbesi* reflectins, and the following primer
was successfully used to obtain the sequence in this manuscript (Lp2-1, AT-GAACCAGCTCTATGAACAGATACC). PCR using primer Lp2-1 and oligo-dT was performed using single strand cDNA as a template. The resulting PCR product contained the 3´ region of the Reflectin A2 gene. To clone the 5´ end of the gene, an internal primer from the resulting PCR product was designed for rapid amplification of cDNA ends (RACE) (GTCCATC-CAGCGTCCCTGCATGTCCATCTGATAGCCGACAT).

To obtain the nucleotide sequence of Reflectin A1, an internal amino acid sequence of protein purified using the method above was determined by Edman degradation following trypsin digestion and HPLC purification. This procedure was performed at the PAN facility at Stanford University (Stanford, CA). Based on the resulting internal amino acid sequences, primers were designed for RACE PCR (for 3´ RACE, TAYAAYAAYGCNTTYYTCNCARATGTGGCA and TAYAAYAAYGCNTTYYAGNCARATGTGGCA; for 5´ RACE, TGCCACATYTAGNANGC). For RACE PCR, a SMART™ RACE cDNA Amplification Kit was used. Additional sets of primers were designed to anneal regions (Lp1-F, GTCTCCTTCGAGAACG-CATGCCCTGTTGTCCGGG; Lp1-R, GGAAAATGTCCGGTTTCATTTTGAA-CATGCCACGCCC; Lp2-F, GTCTCACACGAGAACAAAACCTCGAAGCCACC; Lp2-R, CCCACAATTCCACGTTGA CATATATCCCGACCGCC). PCR products were cloned into pCR®2.1 vector (Invitrogen) using TA cloning. The sequence of the inserted fragment was analyzed with an ABI 3730 Capillary Electrophoresis Genetic Analyzer (Applied Bioscience) at the DNA Sequencing Facility at the University of California, Davis. Sequence data were analyzed using MacVector (Accelrys Software Inc.).

To determine the sequence of Reflectin B1, we obtained de novo sequence using LC-MS/MS (liquid chromatography-followed by 2-dimensional mass spectrometry). Following
HPLC, purified Reflectin B1 was digested with sequence-grade trypsin (Roche) in 0.02% SDS and 50mM Tris-HCl, pH 8.5 at a 50:1 ratio. Peptides derived from this digestion were separated with a PepMapC18 HPLC column. Each fraction was applied to a Waters Micromass QTOF2 tandem mass spectrometer equipped with a nanoflow electrospray ionization source coupled to an Agilent 1100 nano LC system with a Zorbax 300SB-C18 column that was 150 x 0.075 mm with 3.5 um particle size. Based on the obtained amino acid sequences, (TTYGCNGAYGG-NATGTAYMG) and (TTYATGGAYATGCAYTAYGAYGGNATGGGNATG) were used as primers for RACE-PCR performed against single-strand iridophore cDNA as described above. The resulting DNA sequences were deposited in GenBank under accession numbers FJ824804, FJ824805, and FJ824806.

Computational methods

We computationally predicted the potential phosphorylation sites of our novel reflectin sequences using NetPhos 2.0 (Blom et al. 1999, Blom et al. 2004). Hydropathy profiles were generated and transmembrane regions were predicted using TMHMM (Sonnhammer et al. 1998). Calcium-binding sites were predicted using Pfam (Finn et al. 2008). The cytoplasm/membrane interface affinities of the new reflectin protein sequences were analyzed using MPEx software (Jaysinghe et al. 2006) with water-to-lipid partitioning and a 13 amino acid window.

One- and two-dimensional polyacrylamide gel electrophoresis (2-D PAGE)

Protein samples were prepared from either ACh-treated (iridescent) or untreated (not iridescent) iridophore layers of *L. pealeii*. Iridophore layers were powdered using a mortar and pestle under liquid nitrogen, and then solubilized with DeStreak Rehydration Solution (GE Healthcare), followed by centrifuging to remove the residual insoluble materials. Following treatment with DeStreak Rehydration Solution, protein concentration was determined with 2-D
Quant Kit (GE Healthcare). For first-dimension isoelectric focusing (IEF), the soluble component was loaded onto an Immobiline™ DryStrip gel (GE Healthcare) (pH 6-11) (60µg of total protein), followed by SDS-PAGE separation with 10-20% gradient gel (Bio-Rad). For IEF, a Multiphor II Electrophoresis Unit (GE Healthcare) was used. Following 2-D PAGE, gels were analyzed either by staining for phospho-amino acids with Pro-Q Diamond (Molecular Probes, Inc.) or immunodetection of phosphotyrosine. All experiments were performed in triplicate and the results reported are representative of those obtained. One-dimensional PAGE was performed using only the second step of this 2D PAGE protocol.

**Immunoblotting**

Immunoblotting was performed with either anti-reflectin polyclonal antibodies or horseradish peroxidase-conjugated anti-phosphotyrosine primary antibodies (HRP-PY20) (BD Transduction Laboratories™, San Jose, CA). Following one-dimensional SDS-PAGE or 2-D PAGE, proteins were electroblotted from the acrylamide gel to a polyvinylidene fluoride (PVDF) membrane. This PVDF membrane was blocked with 1% BSA in PBST (0.1% Tween-20 in PBS) for 1h at room temperature, then incubated in PBST containing a 1:5,000 dilution of primary antibody for at least 1h at room temperature on a rocking platform. The membrane was then washed three times with PBST to remove unbound and non-specifically bound antibodies. For the use of anti-reflectin polyclonal antibodies, a secondary antibody (alkaline phosphatase-conjugated anti-rabbit) was used to identify immunopositive bands. For alkaline phosphatase detection, a 1:50 dilution of NBT/BCIP Stock Solution (Roche) in alkaline phosphatase buffer (10mM MgCl₂, 100mM Tris-HCl, pH 9.0) was used. Following sufficient development, the reaction was terminated by adding Stop solution (10mM Tris-HCl, 1mM EDTA, pH 8.0). For HRP detection, an ECL kit was used (GE Healthcare). All experiments were performed in triplicate and the results
reported are identified either as the averages of the normalized results or as representative of those obtained.

**Imaging analyses with fluorescent dye**

For analysis of staining with fluorescent dyes Molecular Imager FX™ with Quantity One software for image analysis (Bio-Rad Laboratories, Hercules, CA) was used. ImageJ (NIH) was used to quantify the immuno-blot signal intensity.

**Quantification of phosphorylation level**

To quantify relative levels of reflectin phosphorylation, two different techniques were used: signal intensities of reflectin staining obtained with Pro-Q Diamond stain or normalized signal intensities obtained by Western blotting with HRP-PY20 were measured and normalized to the respective signal intensities obtained from protein quantification with SYPRO Ruby (Molecular Probes) stain, thereby providing the relative phosphorylation level per protein. Values for samples treated with ACh and/or genistein were further normalized to those obtained for the corresponding non-ACh treated samples.

**MS mapping for phosphorylation sites**

Mass Spectroscopy was used to identify the specifically phosphorylated residues. Partially purified Reflectin A1 and Reflectin A2 were subjected to trypsin digestion followed by matrix assisted laser desorption/ionization-time of flight mass spectroscopy (MALDI-TOF MS). All mapping analyses were performed at the Stanford Proteomics and Integrative Research Facility, Stanford University.

**Activation of adaptive iridescence and the effects of protein kinase inhibitors on iridescence**

Squid tissue samples were prepared identically for all adaptive iridescence experiments. The superficial chromatophore- and iridophore-containing layers of skin were surgically re-
moved from freshly killed squid and the iridophore-containing layer trimmed to size to fit the Sylgard-lined incubation tray used in each experiment.

To activate iridescence, specimens of explanted iridophore-containing layers were transferred either to artificial seawater (500 mM NaCl, 10 mM CaCl₂, 10 mM KCl, 12 mM MgCl₂ and 10 mM HEPES, pH 8.0) or 0.2 micron-filtered natural seawater followed by addition of acetylcholine (Sigma, St. Louis, MO) to a final concentration of 10µM. To investigate the effects of various kinase inhibitors, iridophore layer samples were pre-soaked with individual kinase inhibitors prior to the addition of ACh for at least 20 min at room temperature at concentrations indicated in Figure 2. All kinase inhibitors and the calcium ionophore A23187 were purchased from EMD Bioscience.

Spectral reflectance measurements before and after the addition of acetylcholine were obtained with a fiber optic spectrometer (USB2000, Ocean Optics, USA; spectra recorded on PC, using OOIBase 32 software, which automatically plotted reflectance) connected, via a 1-mm-diameter fiber, to the c-mount of a dissecting microscope (Zeiss; angle of acceptance: 13°; distance to specimen approximately 9 cm). At the highest magnification of the microscope, the area of the measured field was approximately 0.3 mm in diameter. Illumination was provided by a Schott fiber-optic microscope-light source, illuminating the iridophores at an angle of incidence of approximately 10 degrees. A diffuse reflection standard (WS-1, Ocean Optics) was used to standardize measurements. For iridophore reflectance measurements, squid were killed by decapitation. Small specimens of skin from the dorsal side of the mantle, containing active iridophores with known optical properties (see Mäthger & Hanlon 2007), were dissected, stretched to original size and pinned onto a Sylgard-covered petri dish. The chromatophore layer was then peeled away, exposing the underlying iridophore layer. To quantify the intensity of iridescence
following genistein treatment, the average digital image grayscale brightness values of individual iridophore patches were normalized to the non-iridescent background and measured as a function of time. Since each iridophore patch is largely elliptical, an elliptical pixel selection tool was used to sample the interior region of each iridophore. In order to eliminate problems associated with edge effects, following identification of the iridophore patch periphery, the selected area was decreased by 20%. The average grayscale brightness intensity value of each iridophore patch was determined through averaging of the brightness intensity value of each pixel contained within the selected area (on a scale of 1 to 256) using Adobe Photoshop. These values were averaged for at least 6 separate iridophore patches within each field of view. These average grayscale brightness intensity values were normalized to take into consideration small local differences in background tissue brightness by subtracting the grayscale brightness values of four adjacent skin regions immediately surrounding each iridophore of interest (each of which had a selected area approximately the same as that of the selected iridophore area). All experiments were performed in triplicate and the results reported are representative of those obtained. The effect of genistein was analyzed using the closely related *L. opalescens* rather than *L. pealeii* due to differences in seasonal availability.

**Polysaccharide analysis**

To determine whether chitin was present in iridescent *Loligo* tissues, the iridophore layer was ground in liquid nitrogen and then acid digested under vacuum in 6M HCl at 110°C for 24h. After acid was removed, digested samples were subjected to amino acid analysis using a Beckman Autoanalyzer 6300 to detect N-acetyl glucosamine, the signature product of acid degradation of chitin. This experiment was repeated three times.
Results

Iridescence in *L. pealeii* skin can be induced by the exogenous addition of the neurotransmitter, ACh (Fig. 1). Rapid changes in both the intensity and color of reflectance are observed following the addition of ACh (Fig. 2). The calcium ionophore A23187 also induces iridescence in *L. pealeii* without the addition of ACh (Fig. 1, right panel) (cf. Cooper and Hanlon 1986). The spectral shift following the initiation of iridescence via the addition of ACh was observed to proceed from 680 nm to 650 nm (Fig. 2) resulting in a visible change in iridophore color from red to orange-yellow.

We extracted and characterized four insoluble proteins expressed only in the iridophore-containing layer of the skin (Fig. 3). Two of these proteins, with apparent molecular masses of 40 kDa and 25 kDa based on SDS-polyacrylamide gel electrophoresis (SDS-PAGE) mobility, strongly cross-react with anti-reflectin antibodies generated against *E. scolopes* reflectins. The remaining two proteins exhibit SDS-PAGE molecular masses of 30 kDa and 20 kDa and only weakly cross-react with the anti-*E. scolopes* reflectin antibody (Fig. 3, right).

We cloned and sequenced the genes coding for three of these proteins and designated them as follows: the strongly immunoreactive proteins Reflectin A1 (44 kDa) and Reflectin A2 (25 kDa); and the weakly immunoreactive protein Reflectin B1 (30 kDa). Their deduced amino acid sequences (Fig. 4) revealed similarity to the reflectin proteins of *E. scolopes* and *Loligo forbesi* (Weiss et al. 2005, Crookes et al. 2004), with Reflectins A1 and A2 exhibiting greater sequence homology to the previously characterized proteins than Reflectin B1. Interestingly, Reflectin B1 appears to be the most abundant protein we have extracted from the dynamically iridescent tissue (Fig. 3). Although seven unique reflectin gene variants were identified in *E. sco-
lopes (Crookes et al. 2004), a similar number of reflectin sequences has not yet been found in *L. pealeii*.

As is the case for the other reflectin proteins, Reflectin A1 and Reflectin A2 contain a series of conserved reflectin motifs (RMs) interspersed throughout the protein, previously defined as 

\[\text{[M/FD(X)\textsubscript{5}MD(X)\textsubscript{5}MDX\textsubscript{3/4}] (Crookes et al. 2004).}\]

There is an additional conserved N-terminal motif 

\[\text{[MEPMSRM(T/S)MDF(H/Q)GR(Y/L)(I/M)DS(M/Q)(G/D)R(I/M)VDP(R/G)]]}\]

in both *Loligo* and *Euprymna* reflectins. This N-terminal region is more evolutionarily conserved across species and reflectin isoforms than the canonical reflectin motif (RM). This highly conserved N-terminal domain aligns poorly with the previously defined repetitive motif, and is not repeated within the reflectin proteins. The N-terminal region is the only conserved motif present in Reflectin B1, which does not contain any of the other canonical RMs as previously defined (Fig. 4). In contrast, reflectin A1 contains five canonical RMs and Reflectin A2 contains three canonical RMs in addition to the N-terminal motif.

All three of the reflectins we characterized had exceptionally high aromatic amino acid residue content: 19% in Reflectin A1, 19% in Reflectin A2, and 19% in Reflectin B1. We have included histidine in our accounting of aromatic residues, since for reasons discussed below, it appears to behave like an aromatic residue in reflectin proteins. In contrast, the average content of aromatic residues in most other proteins is approximately 10%. The total content of arginine and methionine also is quite high in our reflectin proteins: arginine content was 12%, 11% and 10% for Reflectins A1, A2 and B1, respectively. Methionine content was 16%, 14% and 11% for Reflectins A1, A2 and B1, respectively. Notably, these reflectins have extremely few non-aromatic hydrophobic residues that are usually required for protein folding. Reflectin B1, although it contains no internal reflectin motifs as previously defined, shares the N-terminal motif.
and other properties such as high methionine content and high positive charge with previously identified reflectins (Fig. 4).

After complete acid hydrolysis of the dermal iridophore layer under vacuum, no glucosamine was detected by amino acid analysis, indicating the absence of chitin.

Hydropathy plots of the *L. pealeii* reflectins revealed that the three proteins are globally hydrophilic (data not shown), lacking any distinct hydrophobic regions. Computational predictions of trans-membrane helices were also negative. Although hydrophobicity calculations predict that reflectins should be water-soluble, upon purification they co-migrate with the cellular membrane fraction. Accordingly, plots of membrane-interface affinity showed that the non-conserved regions between reflectin motifs are highly energetically stable in membrane interface regions (Fig. 5). The predicted isoelectric points for the three new reflectin proteins are >8.5; these high isoelectric points are largely due to high arginine content, indicating these proteins are likely to be positively charged in the absence of extensive phosphorylation and under physiological conditions.

Using neural network predictions (Blom *et al.* 1999), we predicted numerous potential sites of phosphorylation for the three *Loligo* proteins. For Reflectin A1, we identified 10 serine (Ser), 13 tyrosine (Tyr) and zero threonine (Thr) sites; for Reflectin A2, we identified 5 Ser, 11 Tyr and 2 Thr sites; for Reflectin B2, we find 20 Ser, 6 Tyr and 1 Thr sites. Analysis of the all three reflectin sequences using PROSITE (Hulo *et al.* 2006) and Pfam (Finn *et al.* 2008) predicted no calcium-ion binding motifs (such as EF-hand motifs).

We tested the effects of several tyrosine kinase and protein kinase C (PKC) inhibitors on ACh-induced iridescence in *Loligo*. Genistein, a broad-range tyrosine kinase inhibitor, dramatically suppressed ACh-induced dynamic iridescence, while PKC-specific inhibitors did not. The
effect of genistein on iridescence intensity exhibited a clear dose dependency: higher doses of genistein resulted in progressively dimmer iridocyte reflectance induced by ACh (Fig. 2).

We confirmed and quantified active phosphorylation of the three *Loligo* reflectins by staining SDS-PAGE separated reflectins with Pro-Q Diamond, a stain specific for phosphoamino acids. We compared staining intensities of ACh-treated tissue to those in the ACh + genistein treated tissue. These intensities were normalized to those of untreated control tissue. We found that both Reflectin A1 and Reflectin A2 showed higher phosphorylation levels in the ACh-only treatment (Fig. 6) than in either the ACh+genistein treatment or the control treatment. The ratio of normalized staining intensity between the ACh-only and the ACh+genistein treated tissues increased by 130% and 240% for Reflectins A1 and A2 respectively (Fig. 6). In contrast, the net phosphorylation staining intensity of Reflectin B2 decreased by 25% (Fig. 6).

Since Pro-Q Diamond cannot differentiate between different phosphorylated amino acids, we specifically quantified levels of tyrosine phosphorylation via western blotting with PY-20, a phosphotyrosine-specific antibody. Consistent with results from Pro-Q staining, phosphotyrosine levels for the ACh-treated tissue increased by 170% and 290% for Reflectins A1 and A2, respectively, when compared to the ACh+genistein treatment (Fig. 6). The control values are included in the data presented in Figure 5, as the data for the experimental treatments are plotted as the percent change relative to the control values obtained from the corresponding (control) lane of the gel. We were unable to detect Reflectin B1 with PY-20 in any tissue or treatment, indicating that tyrosine phosphorylation is low, not present, or especially labile for this protein. PY-20 sensitivity is known to be dependent on sequence context, possibly contributing to this observation. Alternatively, this result may suggest that phosphorylation of serine and/or threo-
nine dominate in the active state for Reflectin B1, in contrast to the phosphorylation of tyrosine residues in the Reflectin A sequences.

To identify the locations and identities of the reflectin residues phosphorylated in the ACh-activated tissue, MALDI-TOF mass spectroscopy was performed on trypsin digests of Reflectins A1 and Reflectin A2 (Table 1). This technique revealed that Tyr14 and Tyr127 of Reflectin A1 and Tyr12, Tyr 214, Ser218 and Tyr223 of Reflectin A2 were phosphorylated in the active state (Fig. 4), consistent with our stain-based analysis of phosphorylation. Reflectin A1 had an additional peptide (WMDAQGRFNNQFGQMWHGR) that showed a mass shift consistent with phosphorylation despite the absence of tyrosine, serine or threonine. This may indicate a histidine phosphorylation, as indicated in Figure 4. In those cases in which more than one phosphorylatable amino acid was present in a mass-shifted peptide, we used the congruence of our computational phosphorylation predictions and our mass spectroscopy data to identify the specific residues that were phosphorylated. Interestingly, all but one phosphorylation event occur outside the conserved reflectin domains. It is possible that additional non-tyrosine phosphoamino acids may have been missed by our analyses due to their lability during protein isolation and purification (Mikesh et al. 2006, Sickman and Meyer 2001).

Two-dimensional polyacrylamide gel electrophoresis (2-D PAGE) revealed the populations of phosphorylated states of the reflectins before and after iridescence activation (Fig. 3). The multiple phosphorylated states of each isoform reflect the fact that each reflectin contains multiple residues that can be phosphorylated. However, because reflectin A1 has very limited solubility in the IEF buffer in the absence of detergent (as required for 2D-PAGE), and the different reflectin isoforms exhibit variable solubilities in detergent-free buffer, it proved difficult to quantitatively compare phosphorylation levels between reflectin species in the same 2D-PAGE
experiment. In contrast, we found that reproducible comparisons of the levels of phosphorylation can be obtained from analyses of 1D gel electrophoresis conducted in the presence of SDS. Our 2D-PAGE data enabled us to evaluate the shifts in the populations of differentially phosphorylated states of the reflectins from untreated and ACh-treated samples. Thus, staining with Pro-Q Diamond and PY-20 revealed that Reflectin A1 and Reflectin A2 each consist of populations with several discrete phosphorylated states for both activated and inactivated tissues (Fig. 3). Immunodetection with PY-20 (and to a lesser degree, staining with Pro-Q Diamond) revealed that the most acidic, extensively phosphorylated Reflectin A2 molecules possess an isoelectric point of 7, and are in significantly higher abundance in the activated tissue. Reflectin B2 also appeared to exhibit a higher degree of phosphorylation, as determined with Pro-Q Diamond, following addition of ACh (Fig. 3). In contrast, Pro-Q Diamond staining shows that Reflectin B1 became more basic (less phosphorylated) upon activation, and also is present in several distinct phosphorylated populations (Fig. 3). In general, the reflectin proteins extracted from the acetylcholine-treated tissues appeared to migrate slightly farther than did the proteins from the controls. Although we cannot rule out imperfect alignment of gel images, it seems probable that this is caused by increased negative charge on the activated proteins due to phosphorylation.

**Discussion**

We tested the hypothesis that in loliginid squid, the muscarinic ACh cascade leading to dynamic iridescence has direct biochemical effects on the reflectin proteins comprising the squid iridosomes. We characterized the reflectins found in the dynamically tunable iridosomes and found that these reflectin proteins appear to be progressively phosphorylated or dephosphorylated upon stimulation of iridocytes with ACh. These changes in phosphorylation are concurrent with the activation and deactivation of iridescence. The dose-dependent ability of genistein, a
broad-spectrum tyrosine kinase inhibitor, to quench the stimulation of iridescence as well as prevent the phosphorylation of reflectin, further supports our hypothesis.

Genistein lowered the phosphorylation levels of reflectin proteins below those of non-iridescent, untreated controls. Because baseline protein phosphorylation states are maintained by active homeostasis in cells, this finding supports an active role for reflectin phosphorylation in iridescence modulation. In general, cells exert multiple mechanistic layers of intricate control over protein phosphorylation state (Ubersax and Ferrell 2007). This phenomenon, together with our findings showing that reflectin phosphorylation is dynamic, provides a model for the intricate homeostatic control by which iridocytes regulate the brightness and color of iridescence.

Because PKC inhibitors did not achieve the same iridescence-suppression effect as genistein, we hypothesize a key role for tyrosine kinases in iridescence activation. However, because commercially available PKC inhibitors may be specific only to mammalian proteins, and because their effects on cephalopod protein kinases are unknown, we cannot yet identify the specific protein kinases that mediate the ACh-induced phosphorylation of the reflectins studied here.

Phosphorylation is known to trigger dramatic changes in the conformations and assembly of many proteins and extensive tyrosine phosphorylation is largely responsible for the many rapid changes in cell physiology, such as in insulin signaling (Karim et al. 2006, Schmelzle et al. 2006). This signaling paradigm is a possible precedent for the dynamic iridescence mechanism outlined below.

We suggest three possible mechanisms that may work independently or in concert to account for the observed changes in iridescence. First, phosphorylation may induce changes in the net charge of the reflectins, which may directly alter reflectin protein-protein interaction and assembly. These changes could play a role in altering the refractive index, thickness and/or spac-
ing of the reflectin-containing iridosomes, leading to the observed changes in reflectance.
Second, phosphorylation of reflectins may trigger changes in the Gibbs-Donnan equilibrium across the iridosome membranes, affecting a number of variables such as protein aggregation, protein mobility, turgor pressure, or thickness of the membrane-bound iridosomes. Third, phosphorylation of reflectins could modulate the affinity of reflectins for cell membranes, shifting assembly dynamics toward auto-aggregation and hierarchical self-assembly (cf. Hanlon et al. 1990). Consistent with the third hypothesis, reflectins are characterized, in part, by their extremely high content of aromatic and arginine residues. These residues are both preferentially soluble in membrane/cytoplasm interfaces (Wimley and White 1996).

Membrane/cytoplasm interfaces are known to exhibit unique solubility properties relative to aqueous cytoplasm. Polar aromatic residues are particularly stable in this microenvironment, while very hydrophobic and very hydrophilic residues are not (Granseth et al. 2005; Yau et al., 1998). Arginine, because of its positive charge and long hydrophobic stem exhibits “snorkeling” behavior from the interface into the membrane (Killian and von Heijne 2000). Reflectins are, indeed, characterized by their high content of polar aromatic residues and arginines, and several lines of evidence suggest that they are intrinsically unstructured, with no likely transmembrane, alpha-helix or beta-sheet regions. *E. scolopes* reflectins also have high arginine, hydrophobic and aromatic content in their (Crookes et al. 2004). The first study to characterize the reflectins described them as “methionine-rich membrane-associated proteins” and demonstrated that they have a high affinity for assembly with microsomal membranes when translated *in vitro* (Weiss et al. 2005). Given these data, and what is known about the biophysics of membrane interfaces, we hypothesize that reflectins may exhibit energetically favorable interactions with membrane interfaces. This suggestion is consistent with our observation that the reflectins co-purify with cell
membranes, yet lack any traditionally recognized transmembrane domains and small hydrophobic amino acids. Reflectins A1 and A2 exhibit an interesting periodicity of domains potentially capable of interacting with membrane interfaces, with the proteins apparently alternating between interface-stable regions and hydrophilic domains approximately every 50 amino acids (Fig. 5). Reflectin B2, which exhibits the opposite phosphorylation pattern upon ACh stimulation from Reflectins A1 and A2, does not show the same periodicity, consistent with the possibility of a different assembly dynamic for this protein. Interestingly, the calculated affinity of specific domains of Reflectins A1 and A2 for membrane is predicted to be abolished upon the observed phosphorylation of those sites (Fig. 5). Phosphorylations occur in regions of the protein that prior to phosphorylation are predicted to be highly stable in membrane interfaces, while the conserved reflectin motifs are not phosphorylated, and are predicted to be more stable in the cytoplasmic compartment of the cell. After phosphorylation (mimicked for these calculations by substitution of the original residues with negatively charged glutamate residues), the phosphorylated protein regions are predicted to change from being interface-stable to being stable in the cytoplasm.

Although iridescence is induced by both ACh and elevation of intracellular calcium concentrations via administration of a calcium ionophore (cf. Figure 1), we find no evidence for any calcium ion binding motifs in the newly identified reflectins. Instead, our result suggests that calcium functions as a second messenger in the signal transduction cascade rather than as a direct modulator of the reflectin proteins. Support for our suggestion of a receptor-activated G protein-dependent signal transduction pathway is provided by our observation that iridescence is activated by cholera toxin (an activator of the G protein), in the absence of ACh (data not shown).
Reflectin A1 and A2 from *Loligo* are most similar to the Reflectin 1 group from the static iridophores in *Euprymna*, both in primary sequence and immunoreactivity to antibodies against *Euprymna* reflectin. Therefore, the protein Reflectin B1 may be especially interesting in discerning important functional and evolutionary differences between static iridescence in *E. scolopes* and dynamic iridescence in *Loligo*. Unlike the Reflectins A1 and A2, Reflectin B1 has no conserved reflectin motifs and instead only contains the initial N-terminal peptide characteristic of reflectins. Reflectin B1 is more acidic than Reflectin A1 or A2, and it exhibits the opposite phosphorylation response to ACh stimulation relative to Reflectin A1 or A2. In addition, a comparison of Pro-Q staining to PY-20 staining suggests that Reflectin B1 may be predominantly serine/threonine phosphorylated, rather than tyrosine phosphorylated. Given that Reflectin A1 and A2 and Reflectin B1 have differing phosphorylation responses to ACh, the Reflectin As and Reflectin Bs could operate under separate control mechanisms within a single cell. These characteristics may provide important clues for future work in understanding how reflectin motifs, phosphorylation, and charge modulation of reflectins contribute to the ability of loliginid iridocytes to modulate their reflectance.

Our data demonstrate changes in the relative levels of phosphorylation of the reflectin proteins upon ACh stimulation of the adaptive iridophores of *Loligo*. Detailed quantitation and mapping of the sites of phosphorylation by mass spectroscopy are now in progress to help provide a deeper mechanistic understanding of the biophysical role of phosphorylation in tuning the Bragg reflectors in this system.

Finally, in contrast to prior reports (cf. Denton and Land 1971, Cloney and Brocco 1983), we have shown that chitin is not present in measurable quantities in iridescent squid skin, as indicated by the complete absence of glucosamine after complete acid hydrolysis (data not shown).
This should help end historical speculation that small movements of chitin platelets could be responsible for dynamic iridescence in squid.

Our observation of ACh-dependent changes in reflectin phosphorylation accompanying increased reflectance suggests a molecular mechanism for adaptive control in this biophotonic system. The reciprocal nature of the phosphorylation of Reflectin A1 and A2 vs. Reflectin B presents interesting possibilities for further probing the biophysical phenomena that promote dynamic iridescence. We have traced the signal transduction cascade in this system from the transmitter to a likely final target: the reflectin proteins that apparently change their conformation or assembly to reversibly create the photonic structure. Further elucidation of the underlying molecular mechanisms regulating the intensity and color of the tunable Bragg reflectors in this system can be expected to shed new insight into the evolutionary origin of adaptive reflectance in cephalopods, and may also inspire novel synthesis strategies for production of the next generation of tunable photonic materials for advanced applications.

Acknowledgements

We gratefully acknowledge support from Anteon contract F33615-03-D-5408 to the Marine Biological Laboratory, Woods Hole, MA and grant # W911NF-06-1-0285 from the Army Research Office to D.E.M. We also thank Margaret McFall-Ngai (NIH AI50611) and Edward G. Ruby (NIH RR 12294) for providing the anti-reflectin antibodies, Tim Athens for providing live squid for analysis, Dr. James Pavlovich for de novo amino acid sequence analyses and Drs. Paul Hansma, Igor Mezic, Yoshiko Okamura, Kathy Foltz and Daniel Alkon for their helpful discussions and suggestions.
References


Table 1. Reflectin phosphorylations predicted from MALDI-TOF mass spectroscopy. Peptide sequences identified by MALDI-TOF mass spectroscopy, with asterisks indicating identified phosphorylated residues.

<table>
<thead>
<tr>
<th>Protein ID</th>
<th>Residues</th>
<th>Mass (amu)</th>
<th>Predicted phosphorylated tryptic peptide</th>
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<tbody>
<tr>
<td>Refl A1</td>
<td>10 – 17</td>
<td>1181.564</td>
<td>LYNMY*RNK</td>
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<td></td>
<td>102 – 120</td>
<td>2415.548</td>
<td>WMDAQGRFNNPFGQMWH*GR</td>
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<td>121 – 136</td>
<td>1936.058</td>
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<td>Refl A2</td>
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<td>121 – 136</td>
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<td>QGHYPGY*MSHSMYGR</td>
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<tr>
<td></td>
<td>207 – 230</td>
<td>3350.899</td>
<td>WMDTQGRY<em>MDPS</em>WSNMY*DNYSWY</td>
</tr>
</tbody>
</table>
Figure legends

Figure 1. **Effect of ACh and calcium ionophore on Loligo dynamic iridescence**

Left column: Reflectivity of squid skin increases upon addition of exogenous 10µM ACh. Right column: The same effect is observed upon addition of a calcium ionophore (A23187, 15 µM final conc.) supplemented with calcium chloride (4 mM). Time following addition indicated in lower left of each panel. Each panel shows approximately a 1 cm² patch of skin.

Figure 2. **Quantification of effects of ACh and genistein on dynamic iridescence.**

Top panel: Dose dependency of dynamic iridescence on genistein, a broad-spectrum protein kinase inhibitor. Brightness is in arbitrary units. Lower left panel: Reflectance of *Loligo* skin as a function of time following the simultaneous addition of ACh and genistein. Lower right panel: Reflectance of *Loligo* skin as a function of time following the addition of ACh alone. While error bars are not possible because no two patches of squid skin have identical iridophore patterning, the results shown here are representative of a typical patch of squid skin.

Figure 3. **Electrophoresis, immunoreactivity and phosphorylation of Reflectins.**

Center panels: Overlay of two artificially colored 2D electrophoretic blots of reflectin proteins stained with Pro-Q Diamond stain for phosphoamino acids (top) or PY-20 antibody against phosphotyrosine (bottom). Grey blot shows protein from unactivated control skin. Red blot shows protein from ACh-activated, iridescent skin. Populations of multiple phosphorylation states exist for each of the major reflectins in both the control and the treated state. Upon ACh-activation, these populations of phosphorylation states apparently shift toward charge neutrality.
Left panel: SDS-PAGE of iridophore layer. Right panel: Immunoblot of iridophore layer with antibody to *E. scolopes* reflectins, showing that A-group reflectins are strongly cross-reactive, while B-group reflectins are weakly cross-reactive.

**Figure 4.** Sequence alignment and MALDI-TOF confirmed phosphorylation sites of novel reflectin proteins.

The three novel reflectin proteins characterized here are aligned with *E. scolopes* Reflectin 1B. Phosphorylated residues identified in MALDI-TOF MS analysis are highlighted in red. The dark grey box and white text indicates the conserved N-terminal peptide common to all reflectins. The lighter grey boxes with black text indicate the conserved reflectin motif found in all *E. scolopes* reflectins, but only the Reflectin A sequences of *Loligo*.

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Traces indicate of the calculated energy of interface interaction between a hypothetical membrane surface and 13-residue “windows” of the reflectin proteins as a function of position in the proteins. Values are calculated by the method of Jaysinghe *et al.* (2006). Grey scale bar shows 2 kcal/mol; negative values indicate regions of potential membrane association, while positive values indicate regions predicted to be cytosolic. Blue regions are the conserved “reflectin motifs”. Black regions are the intervening non-conserved regions. Dashed lines show the hypothesized interface interaction energy after phosphorylation. Red dots indicate residues phosphorylated concomitant with the activation of iridescence by ACh. When a residue becomes phosphorylated, the surrounding regions of the protein are predicted to shift from being membrane-associated to cytosolic. Sites of phosphorylation in reflectin 2B have not yet been identified.
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**Left panel.** Changes in Pro-Q Diamond staining intensity of reflectin proteins upon treatment with 10µM ACh (red columns) or 10µM ACh + genistein at indicated concentrations (grey columns). Data are percent differences relative to an untreated control, here shown as zero. All analyses were done in triplicate, and error bars show ± standard deviation. **Right panel.** Changes in PY-20 anti-phosphotyrosine antibody upon treatment with ACh or ACh + genistein. Colors and values are the same as in left panel.
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Reflectin A1: MNRYLRN------------------QR----------LY-NMFRN--KYR--GVMEMPR
Reflectin A2: MNRYMMR------------------HRP----------MSMYRTGRKYR--GVMEMPR
E.s. 1B: MNRFMNK------------------YRPMFNNMSYNRGR--RNR----------GMMEPR
Reflectin B1: MSSFMDPMHYDMGMSHKTGDFSHNCMRSPHSQDRVMDMKGSSKNRRGFNDMMEPR

Reflectin A1: SMTMDPQGYNMDGQSRGAVPDRYDYGRMHDHDYGRSMFNQNGHSMDS----------QRYGWW
Reflectin A2: SMTMDPQGNYMDGQSRGAVPDRYDYGRMHDHDYGRSMFNQNGHSMDS----------QRYGWW
E.s. 1B: SMTMDPQGNYMDGQSRGAVPDRYDYGRMHDHDYGRSMFNQNGHSMDS----------QRYGWW
Reflectin B1: SMTMDPQGNYMDGQSRGAVPDRYDYGRMHDHDYGRSMFNQNGHSMDS----------QRYGWW

Reflectin A1: DNPQRYMDGQYMDQMRWMDAQPFRPNFPGQMM----------GRQGHYPG---------MSSHSMGRNM
Reflectin A2: DNPQRYMDGQYMDQMRWMDAQPFRPNFPGQMM----------GRQGHYPG---------MSSHSMGRNM
E.s. 1B: DNPQRYMDGQYMDQMRWMDAQPFRPNFPGQMM----------GRQGHYPG---------MSSHSMGRNM
Reflectin B1: DNPQRYMDGQYMDQMRWMDAQPFRPNFPGQMM----------GRQGHYPG---------MSSHSMGRNM

Reflectin A1: YNPYHSYASRHFDSPEWMDGQYMDQMRWMDNRYVNPFFNHIM----------YGRNMCPY--Y
Reflectin A2: YNPYHSYASRHFDSPEWMDGQYMDQMRWMDNRYVNPFFNHIM----------YGRNMCPY--Y
E.s. 1B: YNPYHSYASRHFDSPEWMDGQYMDQMRWMDNRYVNPFFNHIM----------YGRNMCPY--Y
Reflectin B1: YNPYHSYASRHFDSPEWMDGQYMDQMRWMDNRYVNPFFNHIM----------YGRNMCPY--Y

Reflectin A1: GNHYNRHMHPEPERYMDGQYMDQMRWMDNRYVNPFFNHIM----------YGRNMCPY--Y
Reflectin A2: GNHYNRHMHPEPERYMDGQYMDQMRWMDNRYVNPFFNHIM----------YGRNMCPY--Y
E.s. 1B: GNHYNRHMHPEPERYMDGQYMDQMRWMDNRYVNPFFNHIM----------YGRNMCPY--Y
Reflectin B1: GNHYNRHMHPEPERYMDGQYMDQMRWMDNRYVNPFFNHIM----------YGRNMCPY--Y

Reflectin A1: FQPERWMDGQYMDQMRWMDNRYVNPFFSHNYGRHNMYPGHYYYHPGMHHYM
Reflectin A2: FQPERWMDGQYMDQMRWMDNRYVNPFFSHNYGRHNMYPGHYYYHPGMHHYM
E.s. 1B: FQPERWMDGQYMDQMRWMDNRYVNPFFSHNYGRHNMYPGHYYYHPGMHHYM
Reflectin B1: FQPERWMDGQYMDQMRWMDNRYVNPFFSHNYGRHNMYPGHYYYHPGMHHYM

Reflectin A1: RHMDSSYQMDHGRWMDNQGRYIDNDYNYDHYMY
Reflectin A2: RHMDSSYQMDHGRWMDNQGRYIDNDYNYDHYMY
E.s. 1B: RHMDSSYQMDHGRWMDNQGRYIDNDYNYDHYMY
Reflectin B1: RHMDSSYQMDHGRWMDNQGRYIDNDYNYDHYMY

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