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Characterization of a bile salt transport system in isolated hepatocytes from shorthorn sculpin, Myoxocephalus scorpius

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The Na+-dependent bile salt transport system plays an important role in hepatic uptake of bile salts in humans and rodents, but it is uncertain whether such a system exists in marine species. Here we demonstrate that isolated shorthorn sculpin hepatocytes specifically transported 3H-taurocholic acid in a time-, dose- and Na+-dependent manner with a \( K_m = 15 \mu M \), a much higher affinity for bile salts than in sea lamprey, suggesting that a Na+-taurocholate cotransporting polypeptide (NTCP, SLC10A1) is expressed in sculpin liver.

Sodium (Na+-)dependent bile salt transport system encoded by the Na+-taurocholate cotransporting polypeptide (NTCP/Ntcp, SLC10A1/Slc10a1) plays an important role in hepatic uptake of bile salts in humans and rodents. In contrast, studies using isolated hepatocytes from the marine skate and rainbow trout (two species that represent elasmobranchs and teleosts, respectively) failed to show Na+-dependent taurocholic acid (TCA) transport activity in these cells, indicating an Ntcp ortholog may not have evolved in these species. However, our previous studies in hepatocytes from sea lamprey, a more primitive vertebrate, have demonstrated Na+-dependent uptake activity for TCA, although with low affinity (\( K_m = 115 \mu M \)), indicating that a Ntcp ortholog evolved early in vertebrate evolution. These observations indicate transport specificity of Ntcp ortholog for TCA in evolutionarily primitive species may vary among species. In this report we examined whether a Na+-dependent TCA transport system could be identified in the liver of shorthorn sculpin (Myoxocephalus scorpius), a marine teleost, hereafter referred to as sculpin.

Sculpin were procured by MDIBL and maintained in 14°C circulating seawater tanks with a 12-hour light-dark cycle. All animals were anesthetized in 0.1 g/L Tricaine and sacrificed by detaching the head from the spinal cord using a sharp knife before liver removal for collagenase perfusion. Hepatocyte preparation was carried out as we have previously described. Briefly, a collagenase (type VIII, 0.05% in marine teleost Ringer) perfusion was carried out through the portal vein at 15°C for about 25 minutes. After collagenase digestion, the liver was disrupted using forceps. Hepatocytes were obtained by low-speed centrifugation with viability averaging ~75%. A 3H-TCA (1 µCi/ml, 10 µM) uptake assay was performed as previously reported. Total cellular protein concentration was used to normalize radioactivity. Non-specific binding activity to cell membranes was determined by subtracting radioactivity determined at 0°C and 0 minute uptake as background. To assess the Na+-dependence of the transport system, NaCl in marine teleost Ringer was replaced with equal molar amounts of choline chloride or LiCl.

TCA uptake activity occurred in a time- and dose-dependent manner at 15°C in isolated sculpin hepatocytes with \( V_{max} \) and \( K_m \) at about 80 pmol/mg protein and 15 µM, respectively (Fig 1). 3H-TCA uptake in these cells was almost completely competitively inhibited by excessive amount of the bile acid taurochenodeoxycholic acid (TCDDA, 200 µM), further suggesting there is a specific bile acid transporter in hepatocytes of this species. When NaCl in the Ringer buffer was substituted with choline chloride, the transport activities were significantly reduced by 50% after both 10 min and 30 min incubations at 15°C (Fig 2), suggesting the presence of a Na+-dependent bile salt transporter. Interestingly, when NaCl in the Ringer was replaced with LiCl the transport activities were significantly increased (Fig 2) by more than two-fold, in contrast to transport activity of mammalian NTCP/Ntcp.

Sculpin hepatocytes demonstrated TCA uptake activity in a Na+- and dose-dependent manner, with a \( K_m = 15 \mu M \) that is comparable with human NTCP (\( K_m = 6 \mu M \)). This suggests an Ntcp ortholog is expressed in this marine teleost and functions in a similar fashion as its mammalian orthologs. As compared to a similar transport system in sea lamprey, sculpin Ntcp demonstrated a much higher affinity for TCA (15 µM vs. 115 µM). The lower \( K_m \) for TCA in sculpin hepatocytes coincides with the evolution of C24 bile acids in teleosts, further suggesting that sculpin Ntcp may be the functional determinant for hepatic bile acid uptake in this species.
Interestingly, sculpin hepatocytes demonstrated enhanced uptake of TCA when Na⁺ was replaced with Li⁺ in the Ringer solution. This finding indicates that there are structural differences between sculpin Ntcp and its mammalian orthologs because NTCP/Ntcp in humans and rodents lost transport activity when Na⁺ was replaced with Li⁺. Of note, this discrepancy was described occasionally for certain other transport systems in teleosts². Future studies will be needed to sequence the sculpin Ntcp gene and characterize its transport activity in vitro in transfected cells.

Figure 1. Isolated sculpin hepatocytes demonstrated TCA uptake activity in a time (left) and dose-dependent (right) manner at 15°C. Michaelis-Menten analysis of TCA uptake at 15°C for 15 minutes; n = 3-4.

Figure 2. TCA uptake in sculpin hepatocytes is Na⁺-dependent and Li⁺-enhanced. *p < 0.05; n = 3.

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Further evidence for a transporter mediated urea uptake in the rectal gland of *Squalus acanthias*

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Cartilaginous fish such as sharks, skates, and rays prevent the water from leaving their bodies by increasing the amount of solute in their tissues and body fluids. The main solute they retain is urea. The studies reported here show that the cells of the rectal gland of the shark contribute to the retention of urea by limiting the amount of urea secreted.

The concentration of urea in the plasma of the shark is 350 mM. Isolated shark rectal glands are perfused with a similar concentration of urea. The concentration of urea in the secretion of the rectal gland is very low1. The osmolality of the rectal gland secretion is the same as that of the plasma or perfusate. Decreasing the concentration of urea in the perfusate of isolated glands decreases the concentration of salt in the secreted fluid2, while increasing the concentration of urea in the perfusate increases the concentration of salt in the fluid2. During these maneuvers, the osmolality of the secretion of the rectal gland remains isotonic with the perfusate. Because of the absence of urea from the secretion of the rectal gland the concentration of salt in the secretion almost doubles. We and others have suggested that this doubling of the concentration of salt in the secretion is the result of the virtual absence of urea from the secretion. Rectal gland cells contain urea at the same concentration present in plasma or perfusate4. We have suggested that urea enters the rectal gland cell via a urea transporter4. In the present series of experiments we re-examined the effect of lowering the concentration of urea in the perfusate of isolated rectal glands and also the effect of the inhibition of the urea transporter with phloretin, on the urea content of rectal gland cells and the secretion of chloride and urea.

Isolated rectal glands of *S. acanthias* were perfused through their single artery by gravity at 16°C and 40 mm Hg pressure with oxygenated shark Ringer’s solution containing 350 mM urea and 5 mM glucose in a single-pass perfusion. Glands were stimulated with theophylline 10^{-4} M and forskolin 5 x 10^{-8} M. In the experiments with phloretin, it was added at a final concentration of 10^{-4} M after the first thirty minutes of perfusion. Venous effluent and duct fluid were collected separately from PE-90 catheters placed in the vein and duct of the gland. Collections were made every ten minutes. At the end of each experiment, an ~100 mg piece of the rectal gland was collected, cleaned of connective tissue and frozen in liquid nitrogen for tissue urea analysis. The tissues thus collected were homogenized in 10 volumes of 100 mM phosphate buffer pH 7.6. Chloride was measured using a Buchler-Cotlove chloridometer (Labconco, Kansas City, MO). Chloride secretion was calculated from the chloride concentration in the duct fluid, the volume of the fluid, the collection time, and the weight of the gland and expressed as µEq per gram of gland per hour. The urea concentration in the dogfish blood, perfusate, rectal gland tissue homogenates, and rectal gland secretion was measured using a commercially available urea kit (QuantiChrom™ Urea Assay Kit, Hayward, CA) and expressed as mmol/l. Plasma, perfusate and rectal gland secretion osmolality were measured using a water vapor osmometer (Wescor, Logan, Utah) and reported as mOsm/l. Protein concentration in tissue homogenates was measured using Bio-Rad Protein assay (Bio-Rad Laboratories, Hercules, CA). Statistical analysis was done using Student’s t-test and ANOVA.

Reducing the concentration of urea in the perfusate resulted in a drop in the concentration of chloride in the secretion of the gland and also in significantly lower secretion of chloride, as shown in Figure 1. As expected, the osmolality of the secretion was also significantly reduced, as shown in Figure 2. The addition of phloretin to the perfusate also resulted in a reduction of the concentration of chloride in the secretion of the gland. This reduced concentration of chloride was associated with a drop in the amount of chloride secreted. However, there was no change in the osmolality of the rectal gland secretion as compared with that of control.
Figure 1. Effect of reducing urea in the perfusate and phloretin on the secretion of chloride by the rectal gland. Reducing the urea concentration in the perfusate to half 175 mM, caused a significant reduction in the secretion of chloride, \( p < 0.001 \) by ANOVA. Phloretin, \( 10^{-4} \) M added at the time indicated by the arrow also caused a significant reduction in the secretion of chloride, \( p < 0.001 \) by ANOVA. Data are expressed as mean ± SEM. \( N = 6 \) for all experiments.

The secretion of urea by these glands is shown in Figure 3. There was very little secretion of urea in the control gland perfused with 350 mM urea. Halving the urea concentration of the perfusate reduced the excretion of urea by the gland by approximately 27%, a statistical significant decrease by ANOVA, The addition of phloretin resulted in a progressive and significant increase in the amount of urea in the secretion of the gland as shown in the figure.

Figure 2. Effect of reducing urea in the perfusate on the osmolality of the rectal gland secretion. Halving the urea concentration the perfusate reduced significantly the osmolality of the secretion, \( p < 0.001 \) by ANOVA. Phloretin, \( 10^{-4} \) M added at the time indicated by the arrow, had no effect on the osmolality of the secretion. Points are mean ± SEM. Measures of dispersion cannot be seen because they are smaller than the symbols. \( N = 6 \) for all experiments.

Figure 3. Effect of reducing the concentration of urea in the perfusate and phloretin on the secretion of urea by isolated perfused rectal glands. The secretion of urea that was low in the control experiments was further reduced by decreasing the concentration of urea in the perfusate to 175 mM, \( p < 0.025 \) by ANOVA. Phloretin \( 10^{-4} \) M added at the time indicated by the arrow, caused an unexpected rise in the urea secretion by the gland, \( p < 0.0001 \) by ANOVA. Points are mean ± SEM. Measures of dispersion cannot be seen in all the series because they are smaller than the symbols. \( N = 6 \) for all experiments.

The intracellular concentration of urea is shown in Figure 4. In the glands perfused with half the usual concentration of urea there was significant drop in its intracellular concentration to 32% of that seen in the glands perfused with 350 mM. Perfusions with phloretin, in the presence of 350 mM urea also resulted in a reduction of the intracellular concentration of urea to 62% of the control glands perfused with 350 mM urea.
Figure 4. Effect of reducing the urea concentration in the perfusate of isolated perfused rectal glands and of phloretin on the intracellular urea concentration. Cutting in half the urea concentration of the perfusate reduced the intracellular urea concentration to 32% of control, \( p < 0.001 \) by Student’s \( t \)-test. Phloretin, \( 10^{-4} \) M, significantly reduced the intracellular concentration of urea, \( p < 0.001 \) by Student’s \( t \)-test. Columns are mean ± SEM. \( N = 6 \) for control and low urea and 4 for the phloretin experiments.

The observation that reducing the urea concentration in the perfusate of the isolated glands decreases the osmolality and the concentration of chloride of the secretion confirms previous observations, and supports the notion that urea plays a role in the secretion of salt by the rectal gland. The fact that urea is present in only small amounts in the secretion of the gland that is isotonic with the plasma, allows the concentration of chloride to rise in the secretion and thus increase the total amount of salt excreted.

We have previously shown that rectal gland cells contain urea at the same concentration as that in the plasma or perfusate. Phloretin, an inhibitor of urea transporters, reduced the intracellular urea concentration of the rectal gland cells, as shown here. This observation suggests that the uptake of urea by the cells is mediated by a urea transporter present in the cell membrane. The functional effects of phloretin are also of interest. Sixty minutes after starting the perfusion of the gland with phloretin, the concentration of chloride in the perfusate dropped by 20%. In association with the drop in the concentration of chloride, the total secretion of chloride also dropped. These observations support the concept that urea, present in the plasma and the rectal gland cells, but not in the secretion of the rectal gland cells, allow the concentration of chloride in the secretion to be higher than that in the plasma. An interesting observation, yet unexplained, is that phloretin increases the concentration of urea in the secretion of the rectal gland. This unexplained phenomenon has been reported in another epithelia, the gills of the \( S. \) acanthias.2

Identification and quantification of gene expression during angiogenesis in the skate, *Leucoraja erinacea*

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The skate, *Leucoraja erinacea*, provides a unique model for studies of hematopoiesis (blood cell production and differentiation) and angiogenesis (formation of new blood vessels). These animals lack bone, allowing easier access to unique and less complex hematopoietic stem cell niches; moreover, they undergo extensive angiogenesis during their reproductive cycles, thereby potentially providing novel insight into the importance of vascular niches. This study further demonstrates the comparability between mammalian and elasmobranch hematopoietic stem cell activity and begins to quantify changes in gene expression during periods of increased angiogenesis and cellular mobilization.

Bone marrow, the tissue responsible for hemopoiesis in mammals, is comprised of vascular and endosteal niches. *L. erinacea* and other elasmobranchs (sharks, skates and rays) have cartilaginous skeletons, i.e., no endosteum, and thus, house hematopoietic stem cells (HSCs) within only a vascular niche. The epigonal organ (EO) of the skate, a homolog of mammalian bone marrow in direct cellular and vascular contact with the gonads, shows a significant increase in cellular turnover (proliferation and apoptosis), as well as angiogenesis, when females transition into reproductive activity²³⁴. These interactions between primary immune tissue and primary reproductive organs in *L. erinacea* suggests further study of these neuroendocrine-immune relationships may provide novel insight into the cellular and genetic mechanisms of hematopoiesis and angiogenesis.

In 2012, our lab observed chemokine receptor 4 (CXCR4) expression in the EO of *L. erinacea*, suggesting homologous molecules are involved in chemokine signaling between elasmobranchs and mammals. While they are known to be pleiotropic molecules, CXCR4 and CXCL12 are key regulators in the retention of HSCs in the bone marrow of mammals⁵⁶. In 2013, *L. erinacea*-specific primers for a number of genes with hematopoietic and angiogenic effects were designed using newly developed databases with genomic and transcriptomic information, and the expression of CXCR4’s cognate ligand, CXCL12, was observed in the EO¹.

The goals of the current experiment were to demonstrate additional genes, as well as to assess the expression levels of genes in *L. erinacea* that are homologous to genes with known hematopoietic and angiogenic functions in mammalian bone marrow. Once identified, we aimed to quantify the change in expression of these genes between non-reproductively active (NRA) and reproductively active (RA) states. Standard PCR was used to demonstrate the expression of VEGFR1, VEGFR2, c-kit and SCF, which are receptor/ligand pairs known to play key roles in hematopoiesis and angiogenesis in mammals.⁷ qPCR was then used to quantify the expression of CXCR4 and CXCL12 in ten female skates (five NRA and five RA)¹.

PCR Primers were developed for CXCR4, CXCL12, c-kit, SCF, VEGFR1, VEGFR2 genes, and β-actin (the qPCR baseline standard) using *L. erinacea* transcriptome data. The NCBI database was used to get the FASTA code for the human sequence of each gene, which was compared to the *L. erinacea* transcriptome to find a contig (an area of sequence similarity) with the best alignment score. A reciprocal cross was performed to compare with the human genome and ensure that the correct gene had been found. The PrimerQuest tool from Integrated DNA Technologies was then used to design and order primers from the contig.

In RA *L. erinacea* we observed a trend of increased CXCR4 expression and decreased CXCL12 expression compared to the NRA group, which indicates the possibility that reproductive activity plays a role in mobilization. However, possibly due to our limited sample size of ten EO tissues (from 5 animals) for each group, the data did not indicate a statistically significant difference of increasing CXCR4 expression of decreasing CXCL12 expression as hypothesized. Data for SCF, c-kit, VEGFA, VEGFR1 and VEGFR2 genes currently show no trends or significant changes in levels of expression in RA compared to NRA individuals. The interaction between CXCR4 and CXCL12 and a panel of additional genes are currently under investigation to further elucidate the role of stem cell mobilization in angiogenesis and revascularization. In continuing this work, we will increase our sample size to reduce the impact of variation among individual specimens.
Figure 1. Representative PCR amplicons demonstrating gene expression in hematopoietic tissues of L. erinacea. Tissues from left (EOL) and right (EOR) epigonal organs as well as the spleen (SP) and Leydig organ (LO).

Figure 2. qPCR histogram plots depicting fold change (ratio of the initial value versus the final value) of gene expression from NRA to RA; positive or negative one-fold change for NRA is baseline by which RA up- or down-regulation is scaled.

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Investigation of the role of novel gene HPS3 and microRNA 636 in proteinuria on zebrafish (Danio rerio)

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As part of the study of proteinuria as a key symptom of many chronic kidney diseases, the relevance of novel gene HPS3 and microRNA 636 for the glomerular filtration barrier (GFB) is investigated by injecting morpholino oligonucleotides and microRNA mimics in zebrafish larvae to cause a knockdown of HPS3 (Hermansky-Pudlak-Syndrome Gene 3) and the target genes of microRNA 636. To understand the relevance of the genes, we examined the phenotype, the fusion of the glomeruli and the development of proteinuria in injected zebrafish. Thus far our results indicate that knockdown of HPS3 and overexpression of microRNA 636 effect the GFB through causing proteinuria and fish with edema.

Based on an up-regulation of the microRNA 636 in the urine of patients suffering from Alport’s syndrome, a rare but serious kidney disease that includes proteinuria as part of a complex of severe symptoms, one could identify HPS3 as a target of miRNA 636 through next generation sequencing. HPS3 is one of nine identified human HPS-causing genes.3 With the knowledge of the so far not published data we hypothesized that novel gene HPS3 and the microRNA 636 play a role in proteinuria.

To evaluate the hypothesis we injected three morpholinos4,5 in transgenic GFB-labeled zebrafish larvae: a morpholino to knockdown HPS3, a microRNA 636 mimic and a control morpholino without binding potential to make sure not the injection process itself is harmful. In addition we used uninjected wild type to consider the egg quality. To assess the damage the absence of the genes have for the glomerular filtration barrier we did a phenotype analysis5, sorting the fish in four main categories due to edema in heart and yolk sack, after 72 hours post fertilization (hpf). The eye-fluorescence assay6 after 96 hpf and 120 hpf is used to investigate the fluorescence level in the retinal vessels of the zebrafish to later be able to evaluate the condition of the filtration barrier. The used transgenic zebrafish Tg(l-fabp:DBP:EGFP) express a vitamin D binding protein fused with the enhanced green fluorescent protein (DBP-EGFP). If the filtration barrier is damaged, the fluorescence level in the retinal vessels due to the loss of DBP-GFP will be reduced comparable to the loss of human albumin in proteinuria. In addition to the phenotype- and eye-fluorescence assay we look at the blood flow after 48 hpf as a control experiment to distinguish between a potential cardiac phenotype and a phenotype based on an impaired filtration barrier. Moreover the wt1b-assay5 can be used to control whether the glomeruli are fused and the development of the kidney was influenced by the injected morpholino. This assay is performed 48 hpf with injected transgenic wt1b-fab-fish.

The analysis of the eye-fluorescence assay reveals for both, HPS3 and microRNA 636, a significant difference of maximum fluorescence in the retinal vessels between the control fish and fish injected with the morpholino (Fig 1 and 2). The data show that the injection occurs in a loss of fluorescence due to a lower concentration of GFB in the blood. Those results confirm our considerations that the knockdown of HPS3 and overexpression of microRNA 636 can cause damage to the filtration barrier and therefore create proteinuria. With the phenotype-analysis we could show that for the microRNA 636 the extent of edema in heart and yolk sack is dose depending: the higher the injected concentration, the stronger the phenotype. Especially in the range between 5 µM and 15 µM a change of concentration has a huge impact on the phenotype. While for 5uM injected microRNA 636 mimic mostly healthy fish without edema could be observed, for an injection with 15 µM the number of fish with a very strong edema is dominating. As well for HPS3 a dose dependence could be determined. Moreover the phenotypeanalysis revealed that the knockdown of HPS3 can cause very strong developmental problems for 25 µM and 50 µM (Fig 3). Since the ratio of fish with a very distinct phenotype and fish with no phenotype differs strongly in the different experiments, further research is necessary to be able to publish reproducible data.
for HPS3. What we could show with not only the phenotype- and eye-fluorescence assay but also our control assays is that a very high concentration has a toxic effect on the fish and causes developmental delays (seen in the wt1b-assay), the stagnation of the blood flow and a severe phenotype.

Our next step will be to use electron microscopy to better understand the impact of the knockdown of HPS3 and overexpression of microRNA 636 on the glomerular filtration barrier. Electron microscopy enables us to understand which particular part of the glomerular filtration barrier is damaged by the absence of the genes.

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The function of heparanase 2 and its connection with kidney function in zebrafish (Danio rerio)

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Heparan sulfate glycosaminoglycans are crucial components of the endothelial cell glycocalyx that constitutes together with the endothelial cells, the glomerular basement membrane and the parietal epithelial cells (podocytes) the glomerular filtration barrier in the kidney. The protein heparanase 2 was recently discovered and is speculated to regulate the activity of heparanase 1, a protein well known for cleaving heparan sulfate components of the endothelial glycocalyx. These preliminary results demonstrate that knockdown of heparanase 2 in zebrafish causes an edematous phenotype and proteinuria, probably by disrupting regulation of heparanase 1.

Heparanase 1 (HPSE1) has been studied extensively for its role in various different disease pathologies. Its physiologic function is described as an Endo-beta(1-4)-D-glucuronidase that degrades heparan sulfate polysaccharide side chains of the glycocalyx covering endothelial cells and basement membranes. Several independent groups have shown in diverse animal models and humans that heparanase expression is upregulated in several primary and secondary glomerular proteinuric diseases, including diabetic nephropathy, IgA nephropathy, the Heymann nephritis model (MGN) and models for MCD/FSGS. In children affected by steroid sensitive syndrome, and in anti-GBM disease. Whereas HPSE1 has been studied extensively, little is known about the more recently discovered heparanase 2 (HPSE2) that was cloned in 2000. HPSE2 inhibits HPSE1 in cell culture experiments, and clinical data indicate HPSE2 expression is markedly elevated in head and neck carcinoma patients, correlating with prolonged time to disease recurrence. However, HPSE2 function remains to be elucidated and has not been studied in an in vivo model. In this study, we examine the role of HPSE2 in glomerular and vascular physiology employing a zebrafish model.

Knockdown of HPSE2 gene was performed by morpholino microinjection into 1 to 4 cell stage zebrafish embryos of a transgenic fish line (Tg l-fabp:DBP:eGFP), which produced a green fluorescence vitamin-D binding protein. These fish were compared to injected control morpholino (CTRL) and uninjected wildtype (WT) group of fish. At 72, 96 and 120 hours post-fertilization (hpf); fish were assessed for generalized edema and categorized into four subgroups: P1 (no edema) to P4 (severe edema).

Injection of HPSE2 morpholino resulted in development of substantial pericardial and yolk sac edema, such that most fish could be classified as P3/4 (Fig 1). In WT1b fish, regular glomerular fusion was seen at 48 hpf (Fig 2) excluding a major disturbance in early kidney development by HPSE2 knockdown. Edema suggested loss of high molecular weight proteins from the vascular system, and loss of colloidosmotic pressure caused generalized edema. We tested this hypothesis by performing at 96 hpf and 120 hpf the fabp eye assay, which indirectly measures intravascular content of fluorescent 78 kDa high M W protein DBP in the vasculature of fish.

We observed a significant decrease in fluorescence levels in the eye as compared to wild type and control morpholino injected fish (Fig 3). There was greater loss of fluorescence at increasing morpholino
concentrations, suggesting a dose dependence of gene knockdown. These results indicated the glomerular filtration barrier is indeed compromised by HEPSE2 knockdown and that proteinuria occurs as a sequela of that.

Figure 3. HPSE2 morpholino injected groups had a lower overall fluorescence level as compared to the CTRL and WT groups, indicating loss of high molecular weight plasma proteins into the urine due to damage to the glomerular filtration barrier. Loss of fluorescence was dose-dependent. 

These data demonstrated a significant role for HPSE2 in the structural integrity of the glomerular filtration barrier. Further experiments are needed to better understand the exact function of this novel gene and will help unravel a novel mechanism involved in causing glomerular injury and acute as well as chronic kidney disease.

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Zn²⁺ increases multidrug resistance-associated protein 2 (Mrp2) transport activity in killifish (Fundulus heteroclitus) renal proximal tubules

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One function of the vertebrate renal proximal tubule is the excretion into the urine of potentially toxic metabolites and foreign chemicals. This is accomplished through the action of numerous transport proteins located in the plasma membrane. Here we show that exposing killifish renal proximal tubules to ZnCl₂ rapidly and reversibly increases the activity of one such transporter, Mrp2, and that this increase is initiated through complex intracellular signaling.

Mrp2 is an ATP-driven efflux transporter that handles a wide range of metabolites, xenobiotics and xenobiotic metabolites. In renal proximal tubule, Mrp2 is localized on the luminal plasma membrane of the epithelial cells, where it can pump chemicals into the forming urine. We previously showed that exposing killifish renal tubules to CdCl₂ or HgCl₂ rapidly decreased Mrp2 transport activity, but then increased activity and transporter expression over many hours.

Zn is in the same group of the periodic table as Hg and Cd. It is an essential trace element and a cofactor for many enzymes. Zn deficiency affects about two billion people in developing countries and is associated with many diseases. To determine whether ZnCl₂ also reduced Mrp2 activity, we exposed isolated killifish renal tubules to 0.1-1 µM ZnCl₂ and measured transporter activity using a fluorescent substrate (Texas red), confocal imaging and image analysis. The assay measured specific and concentrative accumulation of Texas red in the tubule lumens. To our surprise, ZnCl₂ exposure caused a rapid increase in Mrp2 transport activity; this effect was rapidly reversed when the ZnCl₂ was removed (Fig 1).

Previous studies with CdCl₂ indicated that reduced Mrp2 transport activity was signaled through endothelin receptor (ET₁), NO-synthetase (NOS) and soluble guanylyl cyclase and protein kinase C (PKC). The present experiments with specific inhibitors of each of these receptors and enzymes showed that the same processes signaled ZnCl₂ induced activation of Mrp2 transport activity.

Recent experiments with rat brain capillaries and killifish renal tubules show that signalling through phosphoinositide-3-kinase (PI3K, protein kinase B), Akt and mTOR regulates basal P-glycoprotein activity. In the present experiments with killifish renal tubules, we also found that sphingolipid signalling to PI3K, Akt and mTOR was also involved in the Zn-driven increase in Mrp2 activity. Based on the present findings we propose a complex signaling pathway through which ZnCl₂ activates Mrp2 transport.
increases Mrp2 activity in killifish renal tubules (Fig 2). At present, we have no data that speak to the fact that ET<sub>B</sub> receptor/NOS/PKC signaling is common to a Zn-driven pathway that increases Mrp2 activity and a Cd-driven pathway that decreases Mrp2 activity. Our current thinking is that the pathways diverge at PKC, with each activating a different isoform, with one leading to sphingolipid and protein kinase signaling and the other leading elsewhere.

![Figure 2. Signaling pathways through which ZnCl<sub>2</sub> and CdCl<sub>2</sub> regulate Mrp2 transport activity in killifish renal proximal tubules.](image)

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Biphasic, GPER-dependent regulation of Bcrp by bisphenol A in killifish (*Fundulus heteroclitus*) renal proximal tubules

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One function of the kidney is to excrete potentially toxic foreign chemicals, a process that occurs in the proximal tubule. Breast Cancer Resistance Protein (Bcrp) is an ATP-driven transport protein, expressed in proximal tubule that does just that. Here we show that bisphenol a, an environmental estrogen, regulates Bcrp activity in killifish proximal tubules through a novel mechanism involving a non-classical estrogen receptor.

Breast Cancer Resistance Protein (Bcrp) is an ATP-driven drug efflux pump that is highly expressed in barrier and excretory tissues. Estrogen regulates Bcrp expression in human placenta, breast cancer cells in culture and in rat and mouse brain capillaries. However, available evidence suggests tissue-specific modes of regulation. In addition to the hormone (estradiol, E2), persistent environmental pollutants also act through estrogen receptors to elicit estrogen-like effects. Bisphenol A (BPA) is one such environmental estrogen. It has a lower binding affinity to estrogen receptors than E2 and it exerts only a weak, genomic down-regulation of Bcrp in rat brain capillaries compared to E21,3. However, BPA also binds to G-protein-coupled estrogen receptor 1 (GPER) at pico- to nanomolar concentrations and elicits even stronger, immediate effects than E21,4. Thus, BPA is able to alter estrogenic signaling by interacting with non-classical receptors. Here we show that in killifish renal tubules BPA has a biphasic effect on Bcrp transport activity, with nanomolar concentrations reducing activity and micromolar concentrations increasing activity.

We used confocal microscopy to measure steady-state accumulation of mitoxantrone, a fluorescent Bcrp substrate, in lumens of freshly isolated renal tubules as described previously2. Exposure to 10 or 100 nM BPA reduced Bcrp-mediated mitoxantrone secretion into the lumen. In contrast, micromolar concentrations of BPA and of estradiol increased Bcrp transport activity. BPA exposure did not affect transport activity of P-glycoprotein or Mrp4, but did reduce Mrp2 transport activity. G1, an agonist for GPR, had a similar biphasic dose response. The GPER antagonist, G15, abolished the responses of Bcrp to nanomolar and micromolar G1 and to nanomolar BPA, indicating action through GPER. Effects of micromolar BPA were only partially reduced by G15. However, the nonspecific ER antagonist, ICI182.780, together with G15 completely abolished the BPA-mediated stimulation of Bcrp transport, indicating involvement of classical estrogen receptors. The increase in Bcrp transport activity caused by exposure to micromolar concentrations of BPA involved intracellular signaling, since it was reduced by Brefeldin A, an inhibitor of vesicular trafficking, and by H89, an inhibitor of protein kinase A, but not by a PI-kinase inhibitor.

To date, regulation of Bcrp activity has been lined to the action of classical estrogen receptors. The present results for renal proximal tubule provide evidence for a novel mechanism of Bcrp regulation through GPER and intracellular signaling.

Characterizing cilia in sand dollar (Echinarchnius parma) embryos after lithium ion treatment

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Cilia are whip-like, cell surface appendages that provide locomotory and sensory capabilities throughout the animal kingdom. During early echinoid development, cilia of different types rapidly form and differentiate on different germ layers. Lithium ion treatment can perturb embryogenesis by shifting germ layer boundaries and thereby aid in characterizing cilia differentiation. Our data suggest the change in cilia phenotype seen on the surfaces of lithium ion-treated “vegetalized” embryos is due to an increase in endodermal expression and endodermal cilia formation compared to untreated embryos.

The many forms and functions of cilia in humans result in a wide array of diseases, or “ciliopathies,” when cilia form abnormally. Echinoid embryos provide a valuable model for studying cilia growth and differentiation due to rapid onset of ciliogenesis and multiple cilia types that form. Echinarchnius parma globally express cilia on their outer surface 22 hours post-fertilization (hpf). Four hours later, at least three distinct ciliary phenotypes arise on the gastrula stage embryo, while a primitive gut invaginates into the blastocoel lumen. To study cilia of the gut, incubation of E. parma embryos in sea water plus 12.5 mM lithium chloride or 6 hours from 4-10 hpf “vegetalized” embryos¹ and produced embryos with exaggerated gut domains with five distinct morphological phenotypes: long endogut (63.3% of embryos), short endogut (11.9% of embryos), unconstricted exogut (1.1% of embryos), constricted exogut (1.5% of embryos), and irregular morphology (22.2% of embryos) (n = 1,449 embryos in one trial; Fig 1). We hypothesized that an exogut region would be composed of endodermal tissue, while mesodermal expression would not be affected². To test this hypothesis, vegetalized embryos were immunofluorescently stained using germ layer specific antibodies (antibody 5c7 for Endo1, gift of D. McClay, Duke University) and imaged by confocal microscopy. Our data show vegetalized embryos overexpressed endoderm in the exogut region, but also throughout the embryo (Fig 1A). Vegetalized embryos that do not exhibit the classic exogut phenotype also showed increased endodermal expression (Fig 1B). Additionally, mesodermal expression was found to be unperturbed between vegetalized and wild type embryos (not shown). These data suggest that for studying cilia types, the current model of interpreting echinoderm vegetalization based on morphology must be revised to account for variance in increased endodermal expression throughout a lithium-perturbed population.

Figure 1. Endodermal expression in vegetalized E. parma embryos exceeds the exogut region. Two phenotypically distinct vegetalized E. parma embryos illustrate two of the five morphological phenotypes we observed. Blue indicates DNA and red indicates endodermal expression of Endo1. Vegetal pole is oriented downward in images. A) Vegetalized embryo with constricted exogut exhibits a classic “vegetalized phenotype” with exogut narrower than rest of embryo as seen in bottom half of image; surprisingly, Endo1 expression reveals that endoderm domain is not limited to exogut. B) Vegetalized embryo with short endogut exhibits a short invagination at vegetal pole. Endodermal expression is concentrated near the invagination, but is also seen throughout the embryo.

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Role of genetic diversity and water quality in the 2013 decline of eelgrass (Zostera marina) populations of the Upper Frenchman Bay, ME coastlines

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To determine the role of genetic diversity in the 2013 loss of eelgrass (Zostera marina) in the upper Frenchman Bay, Mount Desert Island, a set of genetic markers were sequenced and compared against eelgrass density and tensile strength. While the genetic diversity of the eelgrass did not explain differences in health of eelgrass by site, some water quality parameters such as silica and nitrate/nitrite in the water column may be contributing factors to the decline.

Eelgrass beds (Zostera marina) serve a vital role invaluable to commercial supplies of seafood in Maine, by forming estuaries for young fish and shellfish. Since aerial fly-over imaging done in 1996, the coverage of eelgrass in the upper Frenchman Bay, ME, has diminished from 3,000 to less than 200 acres1,2. In the summer of 2013, major restoration areas around Mount Desert Island at Hadley Point, Berry Cove, and Thomas Island, established by ecologists to counter this loss, also declined. Genetic diversity of eelgrass may also explain differences in survival among populations, with higher genetic diversity potentially imparting more resistance to environmental change or physical disturbance. The genetic diversity of six populations (2 unhealthy: Berry Cove (BC) and Bar West (BW); 4 healthy: Bar East (BE), Ship Harbor (SH), Stave Island (SI), and Wonderland (W)) across Mount Desert Island was quantified using microsatellite size DNA analysis for six markers. The inbreeding coefficient (F_{IS}), a measure of genetic diversity, was calculated for each population and compared against eelgrass density; no correlation was found between the health of the site and the genetic diversity of the plants (Fig 1).

The tensile strength of the eelgrass, perhaps a measure of resilience to physical disturbance, was compared to the genetic diversity of each site. There was no correlation found between the two variables (Fig 2), and there was no difference in healthy versus unhealthy sites in the tensile strength of the plants (Fig 3).
Genetic diversity and tensile strength did not seem to correlate with the health of eelgrass populations as was expected for Berry Cove, which had high levels of intrapopulation diversity and little differentiation from healthy sites. Nevertheless, the population at Ship Harbor, a healthy site, showed the lowest intrapopulation diversity and highest interpopulation differentiation.

To determine the role of water quality in the decline, nitrate/nitrite, ammonia, phosphorus, and silicate levels were measured using a Bran Luebbe Autoanalyzer III on four separate occasions at all sites described above, plus several additional sites, over the course of the summer. There were differences in silica in the water column among sites as determined by one-way ANOVA ($F = 2.585$, $p = 0.0125$). A comparison of all sites documented to have lost eelgrass (Jordan River, Thomas Island East, Thomas Island West, Hadley Point, Berry Cove, Lamoine Shore, and Bar West) and all sites with intact eelgrass beds (Bar East, Stave Island, Wonderland and Ship Harbor) revealed a significant difference in silica in the water column over the course of the summer ($t$-
test, \( p = 0.0179 \)). Silica was highest where eelgrass was intact. There was also a significant difference among sites for nitrate/nitrite levels in the water column as determined by one-way ANOVA (\( F = 3.00, p = 0.005 \)). A comparison of sites documented to have lost eelgrass and sites with intact eelgrass as listed above revealed a significant difference in nitrate/nitrite in the water column over the course of the summer (t-test, \( p = 0.003 \)). Nitrate/nitrite levels were highest where eelgrass was intact. There was not a significant difference in ammonia levels among sites as determined by one-way ANOVA (\( F = 0.780, p = 0.667 \)) and consequently, there was not a significant difference in total dissolved inorganic nitrogen (DIN) among sites (\( F = 1.72, p = 0.099 \)). The same was true for phosphorus (\( F = 0.385, p = 0.961 \)). The affect of water column nutrient levels on eelgrass health and susceptibility to disease and damage will be the subject of future research.

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Population genetics and abundance of the invasive European green crab (*Carcinus maenas*) and its role in eelgrass (*Zostera marina*) loss around Mount Desert Island

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Following the devastating 2013 loss of eelgrass (*Zostera marina*) in the upper Frenchman Bay, Mount Desert Island, the role of the invasive green crab (*Carcinus maenas*) in the decline was investigated. Neither the genetics of the crabs nor their abundance correlated to the overall health of the eelgrass. The loss of eelgrass may be due to other factors or an interplay of abiotic and biotic factors.

*Zostera marina* or “eelgrass” is a common seagrass found widespread in northern coastlines across the world. Eelgrass beds serve a vital role in marine ecosystems of Maine, protecting shorelines against erosive forces and providing an ecosystem for larval and juvenile fish and shellfish. Between 1996 and 2013, the coverage of eelgrass in the upper Frenchman Bay area of Mount Desert Island, ME, was reduced from 3,174 to between 75 and 183 acres1,2. Contrasting sites with healthy populations to sites that have sustained significant loss is key to identifying decline causation. One potential culprit in eelgrass loss is the invasive green crab (*Carcinus maenas*); green crabs voraciously target soft shell clams3 and damage eelgrass during their foraging behavior4,5. Green crabs were first introduced to the New York Harbor from Europe in the 1800s5, and a secondary invasion of novel northern haplotypes, hypothesized to be more voracious7, occurred in Nova Scotia during the 1980s and 1990s3. To determine whether the genetics or abundance of the crabs are correlative with the health of the eelgrass, crabs were assessed at seven eelgrass sites around Mount Desert Island. Using the neutral mitochondrial genetic marker, cytochrome oxidase c I (COI), the haplotype of 15 individuals per site was ascertained. Haplotype 1, the original invading haplotype, was most frequent; novel northern haplotypes (4,5,6) were found in several individuals around the island (Fig 1).

![Figure 1. C. maenas COI haplotype frequencies per site (BC=Berry Cove; BE=Bar East; BW=Bar West; HP=Hadley Point; SH=Ship Harbor; SI=Stave Island; W=Wonderland). Sites marked in green are deemed healthy eelgrass habitats; sites in red are deemed unhealthy eelgrass habitats based on eelgrass density and biomass.](image)

Since northern haplotypes are suspected to be more voracious, and hence cause more damage than the original invading haplotype (haplotype 1), the percent of northern haplotypes per site was compared to the health of the eelgrass, as measured by eelgrass shoot density. There was no correlation of the percent of crabs of a northern haplotype with the health of the eelgrass at that site (Fig 2).
Figure 2. Linear relationship between percent of crabs identified as a northern haplotype and eelgrass density ($R^2 = 0.13$, $p = 0.38$). Sites in green are deemed healthy eelgrass habitats; sites in red are deemed unhealthy eelgrass habitats.

Figure 3. Average green crab (C. maenas) trapped over four days in September of 2014 in eelgrass habitats around Mount Desert Island. Sites in green are deemed healthy eelgrass habitats; sites in red are deemed unhealthy eelgrass habitats. Significant differences in average crabs trapped were determined with a one-way ANOVA and Tukey’s posthoc test ($\alpha = 0.05$). Letters above eelgrass habitats indicate where there was no significant difference between sites.

While it cannot be ruled out that the presence of green crabs contributed to the 2013 eelgrass decline in Frenchman Bay, the loss of eelgrass may be due to other factors or an interplay of abiotic and biotic factors.

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A survey of developmental abnormalities in *Fundulus heteroclitus* embryos from Northeast Creek

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The embryos of Northern killifish can develop immersed in seawater or in air when stranded on rocks or vegetation in estuaries. These embryos are remarkably tolerant to desiccation, thermal and salinity stress. About 6% of these embryos show developmental anomalies, of which the most common anomalies were albinism (1.6%) and microcephaly (0.7%). These data provide a baseline estimate of the rate of occurrence of developmental anomalies and/or mutations in a killifish population from a relatively pristine estuarine environment.

The Northern subspecies of killifish, *Fundulus heteroclitus macrolepidotus*, spawn in Maine in estuaries in June and July during daily high tides\(^1,2\). The adults may migrate to and from full strength seawater (SW, ~30 ppt) to freshwater (<1 ppt) but tend to spawn in brackish water (~10 ppt)\(^1,2\). As the tide ebbs, some embryos remain immersed and some embryos become stranded on rocks and vegetation. These stranded embryos are exposed to air for long periods (up to 14 days)\(^1\). In spite of enduring significant transient desiccation and thermal stress, aerially incubated embryos develop normally and hatch when flooded by high tides after 12-14 days\(^1,2\). During our ongoing studies on the physiological and molecular mechanisms these embryos employ to resist desiccation stress\(^1\) we noticed that there were a number of developmental abnormalities and/or mutants that appeared in our cultures. Northeast Creek on Mount Desert Island, ME is considered a relatively pristine environment, and animals from this population have been used for a variety of physiological studies as well as for genomic analyses. We felt a preliminary survey of the occurrences and types of embryo anomalies or mutations that spontaneously occur in this population would provide useful baseline information for research, particularly for investigations of embryonic mutations resulting from exposure to environmental pollutants.

Fish were caught in minnow traps at Northeast Creek and kept in aquaria with natural running SW (~30 ppt). Gametes were gently expressed from fish into a beaker with 25 ml of 10 ppt artificial seawater (ASW; Instant Ocean, Mentor, OH). For these experiments 10 ppt ASW was used; as it approximated the salinity at the sites in the estuary at which fish spawn. After 30 min, the embryos were rinsed with 10 ppt ASW, placed on filter paper moistened with 10 ppt ASW in Petri dishes, and cultured aerially in a chamber in which the humidity was equilibrated to that of 10 ppt ASW\(^1\). Embryos from 7 days post-fertilization (dpf) to 14 dpf were inspected daily for anomalies using a dissecting microscope. Abnormal embryos were classified, based on deformation or alteration of normal embryo body morphology. Typical examples were photographed, and an extensive reference file of the number of types of anomalies was established.

In total, 13667 fertilized embryos were examined from 11 separate batches. The average number of males and females (mean ± S.E.) used per batch was 7 ± 3 and 13 ± 5. Twenty-six individual types of morphological anomalies were identified. The overall mutation/anomaly rate was about 6%. The six most frequent anomalies were (occurrence rate shown in parentheses): albinism (1.6%), severe head and body deformation (0.8%), microcephaly (0.7%), only chromatophores but no body development (0.5%), microcephaly and missing tail (0.2%) and reduced coloration with scattered chromatophores (0.2%). A number of rare anomalies such as two-headed embryos (0.07%) were also noted. In general the types of anomalies noted also have been observed in zebrafish and other species. This preliminary survey provides valuable initial baseline data that characterize embryonic development anomalies and possible mutations in the Northeast Creek killifish population.

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