A Heterogeneous Mixture of F-Series Prostaglandins Promotes Sperm Guidance in the *Caenorhabditis elegans* Reproductive Tract

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**Abstract**

The mechanisms that guide motile sperm through the female reproductive tract to oocytes are not well understood. We have shown that *Caenorhabditis elegans* oocytes synthesize sperm guiding F-series prostaglandins from polyunsaturated fatty acid (PUFA) precursors provided in yolk lipoprotein complexes. Here we use genetics and electrospray ionization tandem mass spectrometry to partially delineate F-series prostaglandin metabolism pathways. We show that omega-6 and omega-3 PUFAs, including arachidonic and eicosapentaenoic acids, are converted into more than 10 structurally related F-series prostaglandins, which function collectively and largely redundantly to promote sperm guidance. Disruption of omega-3 F-series synthesis triggers compensatory up-regulation of prostaglandins derived from omega-6 PUFAs. *C. elegans* F-series prostaglandin synthesis involves biochemical mechanisms distinct from those in mammalian cyclooxygenase-dependent pathways, yet PGF₂α stereoisomers are still synthesized. A comparison of F-series prostaglandins in *C. elegans* and mouse tissues reveals shared features. Finally, we show that a conserved cytochrome P450 enzyme, whose human homolog is implicated in Bietti's Crystalline Dystrophy, negatively regulates prostaglandin synthesis. These results support the model that multiple cyclooxygenase-independent prostaglandins function together to promote sperm motility important for fertilization. This cyclooxygenase-independent pathway for F-series synthesis may be conserved.

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**Introduction**

The union of oocyte and sperm, known as fertilization, is paramount to the survival of animal species [1–4]. Oocytes tend to be large stationary cells filled with nutrients to support early embryonic development. Sperm, on the other hand, are smaller mobile cells capable of sensing their environment and responding with changes in motility. Sperm have a relatively simple job—find an oocyte and provide it with a second set of chromosomes. While much has been learned about sperm motility in aquatic species [5,6], the mechanisms that guide sperm to oocytes in internally fertilizing animals are less well understood. These guidance mechanisms are likely crucial for reproductive success [7,8]. Evidence is emerging that lipid signaling molecules, such as steroids and prostaglandins are important regulators of sperm motility in the human female reproductive tract [9–13].

*Caenorhabditis elegans* is a powerful model to study sperm behavior and fertilization [2,4,14–16]. Its transparent epidermis allows visualization of sperm distribution and motility in live wild-type and mutant animals (Figure 1A) [17–19]. Sperm velocity, directional velocity toward eggs, and reversal frequency within the reproductive tract can be quantified from time-lapse videos [18,20]. Genetic and biochemical analyses have shown that oocytes synthesize F-series prostaglandins (PGs) from polyunsaturated fatty acids (PUFAs) provided in yolk lipoprotein complexes [18,20,21]. PUFAs are fatty acids with two or more double bonds, which are essential for prostaglandin cyclopentane ring formation (Figure S1) [22]. Mutations in PUFA biosynthesis enzymes and predicted prostaglandin synthases cause nonautonomous defects in sperm motility, resulting in inefficient targeting to the site of fertilization and reduced reproductive output [18,20]. Introducing nanomolar concentrations of F-series prostaglandins into the uteri of prostaglandin-deficient mutants has an immediate effect on sperm motility [20]. Liquid chromatography coupled to electrospray ionization tandem mass spectrometry (LC-MS/MS) has identified multiple F-series prostaglandins in *C. elegans* extracts [20,21]. The synthesis of CePGF2, an analog of PGF2α, is dependent on 20-carbon PUFA synthesis and transport to oocytes [20]. However, the extent to which specific loss of CePGF2 or other prostaglandins affects sperm guidance is not clear. It is also not clear whether *C. elegans* F-series prostaglandins are derived from a single or multiple PUFA precursors.
Mammals lack the desaturase enzymes necessary to convert monounsaturated fatty acids into PUFA s [23,24]. A key enzyme in mammalian prostaglandin synthesis is cyclooxygenase or PG G/H synthase, which converts PUFA s into the intermediate PGH \(_2\) (Figure S1) [25]. There are three prostaglandin classes that are synthesized from distinct PUFA precursors [26–28] (Figure 1B). The F1 class is derived from dihomo-gamma-linolenic acid (DGLA), a 20-carbon PUFA with three double bonds in the carbon backbone. DGLA is also called 20:3\(_{\text{n6}}\), where the third double bond occurs at the n6 or omega-6 position. The F2 class is derived from arachidonic acid (AA, 20:4\(_{\text{n6}}\); Figure S1B) and the F3 class is derived from the omega-3 PUFA eicosapentaenoic acid (EPA, 20:5\(_{\text{n3}}\)). PGF\(_{2\alpha}\) is a member of the F2 class synthesized from AA by the sequential action of cyclooxygenase and PGF synthase (Figure S1) [22,25]. 11\(_\beta\)-PGF\(_{2\alpha}\) and 9\(_\beta\)-PGF\(_{2\alpha}\) stereoisomers are formed from PGD\(_2\) and PGE\(_2\) intermediates, respectively [29]. In addition, a wide array of F-series prostaglandin isomers is formed via cyclooxygenase-independent mechanisms mediated by free radical-initiated peroxidation [30]. In most cases, the biological activities of F-series prostaglandins are not known.

The \(C.\) elegans PUFA biosynthetic pathways have been delineated (Figure 1C) [31,32]. Four fatty acid desaturases encoded by \(fat\) genes convert monounsaturated fatty acids into omega-3 and omega-6 PUFA s. Here we use mutations in \(fat\) genes, together with LC-MS/MS, to delineate F-series prostaglandin synthesis pathways. We show that multiple F-series prostaglandins, whose formation in large part requires an adult germ line, act collectively to promote sperm guidance. F2 class prostaglandin synthesis does not involve H\(_2\), D\(_3\), and E\(_2\) intermediates that are characteristic of mammalian cyclooxygenase-dependent pathways (Figure S1). We identify a conserved cytochrome P450 enzyme that functions to inhibit prostaglandin metabolism, possibly by diverting their precursors into an alternative pathway(s). Our results support the model that \(C.\) elegans oocytes secrete a blend of F-series prostaglandins for guiding sperm to the fertilization site.

**Figure 1.** \(C.\) elegans gonad, F-series prostaglandin structures, and PUFA metabolism pathways. (A) The \(C.\) elegans hermaphrodite reproductive tract. A differential interference contrast (DIC) image of an adult hermaphrodite is shown above with a diagram of half the uterus below. Males inject sperm through the vulva (V) into the uterus. Sperm become motile by extending a pseudopod and crawl around fertilized embryos (E) toward the spermatheca (S, yellow outline). Oocytes (O) enter the spermatheca during oocyte maturation and ovulation. The uterus is divided into 3 zones for quantification of sperm distribution. Scale bar, 0.1 mm. (B) Human F-series prostaglandin structures showing carbon numbering and important energy-induced cleavage sites characteristic of F-series prostaglandins. \(z_1\) cleavage causes loss of 44 Da from the parent ion m/z 193. Prostaglandin precursors for each class are shown in parentheses. (C) Diagram of \(C.\) elegans PUFA synthesis pathways. \(\Delta_9\) and \(\Delta_{12}\) desaturase activities generate substrates for the n3 desaturase FAT-1, the \(\Delta_6\) desaturase FAT-3, and the \(\Delta_5\) desaturase FAT-4 [31]. Fatty acids are abbreviated as 20:4\(_{\text{n6}}\), which has 20 carbons with four double bonds, the first occurring at the n6 position. LA, linoleic acid; ALA, linolenic acid; GLA, \(\gamma\)-linolenic acid; STA, stearidonic acid; DGLA, dihomo-\(\gamma\)-linolenic acid; O3AA, omega-3 arachidonic acid; AA, arachidonic acid; EPA, eicosapentaenoic acid.

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**Author Summary**

A fundamental question in cell and developmental biology is how motile cells find their target destinations. One of the most important cell targeting mechanisms involves the sperm and oocyte, which unite during fertilization to produce the next generation of offspring. We have been using the nematode \(C.\) elegans to delineate these mechanisms. Our prior studies have shown that oocytes secrete F-series prostaglandins that stimulate sperm motility. Prostaglandins are widespread signaling molecules derived from polyunsaturated fatty acids or PUFA s. Mammals are not capable of synthesizing PUFA s and must receive them in the diet. \(C.\) elegans was not thought to synthesize prostaglandins because the genome lacks cyclooxygenases, enzymes that catalyze the rate-limiting step in mammalian prostaglandin synthesis. Here we show that \(C.\) elegans oocytes synthesize a heterogenous mixture of structurally related F-series prostaglandins derived from different PUFA classes, including the enantiomer of PGF\(_{2\alpha}\). These prostaglandins function collectively and redundantly to guide sperm to the fertilization site. Our results indicate that F-series prostaglandins can be synthesized independently of cyclooxygenase enzymes. This novel pathway may be evolutionarily conserved. Evidence is emerging that prostaglandins regulate sperm motility in the female reproductive tract of humans.
Results

PUFAs act collectively to promote sperm guidance

To explore the F-series prostaglandin biosynthetic mechanisms important for sperm guidance, we examined fat mutants deficient in the synthesis of specific PUFAs classes (Figure 1C). Two quantitative assays were used to evaluate sperm guidance. In both assays, wild-type males are stained with MitoTracker CMXRos dye, which brightly labels sperm [18]. Stained males are mated to unstained mutant or control hermaphrodites and separated. The first assay measures sperm distribution within the uterus one hour after mating, using three defined zones (Figure 1A). In controls, over 90% of sperm move around developing embryos to the fertilization site, called the spermatheca in an hour (Figure 2). The second assay uses time-lapse imaging to directly measure sperm velocity, directional velocity, and reversal frequency within the uterus immediately after mating. Sperm in control uteri move with an average velocity of 8.6 microns per minute, rarely reversing direction. Average directional velocity toward the spermatheca is less than average velocity as sperm migrate around embryos blocking their path (Table 1, line 1).

First, wild-type males were mated to fat-1(wa9) and fat-4(wa14) mutant hermaphrodites. fat-1(wa9) mutants have undetectable EPA (20:5n3) and omega-3 AA (20:4n6) levels relative to the wild type [32,33]. On the other hand, fat-4(wa14) mutants have undetectable EPA and AA, but have elevated omega-3 AA levels [32,33]. Minor sperm motility defects are evident from sperm distribution and time-lapse imaging assays, but a large percentage of sperm are still able to target the spermatheca (Figure 2 and Table 1, lines 2 and 3). These data raise the possibility that DGLA, which is still synthesized in both fat-1 and fat-4 mutants, is the key precursor for sperm guiding prostaglandins.

To test whether or not DGLA is essential, we compared sperm guidance values in fat-3(wa22) mutants to fat-1(wa9) fat-4(wa14) double mutants. fat-3(wa22) mutants contain abundant 18:3n3 (linolenic acid, ALA), but trace amounts of 20:3n3 and undetectable DGLA (20:3n6), AA, omega-3 AA, and EPA [32]. On the other hand, fat-1(wa9) fat-4(wa14) double mutants contain abundant DGLA with undetectable AA, omega-3 AA, and EPA [32,33]. Surprisingly, the sperm guidance values in both mutant strains were similar, albeit significantly less than controls (Figure 2 and Table 1, lines 1, 4, and 5; P<0.001). Sperm in both strains have similar directional velocity, although there are differences in velocity and reversal frequency (Table 1, lines 4 and 5). These data indicate that DGLA is not essential for sperm guidance, but it may have a modulatory or partially redundant role.

Finally, we measured sperm guidance in the healthiest 1–2 day old adult fat-2(wa17) mutants, avoiding those mutants with considerable pleiotropy. fat-2 encodes a Δ12 fatty-acyl desaturase that converts monounsaturated fatty acids into PUFAs. fat-2(wa17) mutants contain reduced levels of all PUFAs, although approximately 5% Δ12 desaturase activity persists in this strain that is sufficient for prostaglandin synthesis [20,32]. The sperm guidance defects in fat-2 mutants are more severe than fat-3(wa22) mutants and fat-1(wa9) fat-4(wa14) double mutants (Figure 2 and Table 1, lines 4, 5, and 6). Furthermore, the defects are similar to mex-2(h1060) mutants, which fail to transport PUFAs to oocytes and glp-4(bn2) mutants, which lack oocytes (Table 1, lines 6, 7, and 0) [18,34,35]. The trend from fat-2 mutant data is that sperm guidance defects inversely correlate with total PUFA content. We conclude that loss of single PUFAs classes does not have major effects on sperm guidance. Sperm guidance defects depend on the extent to which PUFA classes are collectively eliminated.

F-series prostaglandins are synthesized from distinct PUFAs

A collective requirement for PUFA synthesis could have multiple interpretations. One model is that F-series prostaglandins are synthesized from multiple PUFA classes. A non-mutually exclusive alternative is that worms compensate for loss of a PUFA class by up-regulating prostaglandin synthesis from another class. In both models, prostaglandins from different PUFAs must have largely redundant functions. These (and additional) models were tested through LC-MS/MS analyses of wild-type and fat mutant lipid extracts. Mammals synthesize F1, F2, and F3 class prostaglandins from DGLA, AA, and EPA, respectively (Figure 1B and Figure S1). To test whether wild-type worm extracts contained these classes, we used LC-MS/MS operated in multiple reaction monitoring (MRM) mode. In MRM, the parent ion to decomposition (product) ion mass transition is specified, providing excellent specificity and sensitivity. The F1 prostaglandins are detected with mass transition m/z 355/311, the F2 class with transition m/z 335/193, and the F3 class with transition m/z 351/193 or 351/191 [36–38]. We found that wild-type extracts contained multiple isomers of each class (Figure 3A). Comparing the collision-induced decomposition (CID or MS/MS) patterns of these prostaglandins to their corresponding chemically synthesized PGF1α, PGF2α, or PGF3α standards shows extensive similarities (Figure 4, Table 2, and Table 3). Key shared decomposition features include the loss of 44 Da (–C₂H₅O) from the parent ion.
followed by two 18 Da losses (~2 x H₂O), and the presence of the m/z 193 ion in the F2 and F3 series (Figure 1B) [36].

One of the two most abundant worm F1 isomers (CePGF1) has a chromatography retention time (RT) that is indistinguishable from PGF1a (Figure 3). Their CID patterns are nearly identical, with the exception of minor ions at m/z 157 and 223 (Figure 4). We have previously shown that CePGF2 and PGF2a have indistinguishable chromatography RTs and nearly identical CID patterns [20]. In contrast, the PGF3a RT is several seconds shorter than the closest worm F3 class prostaglandin at RT = 11.33 (Figure 3). While the CID patterns of PGF3a and this prostaglandin are similar, there are more differences than in F1 and F2 classes (Figure 4). One key difference is the abundance of product ion m/z 193, which is thought to derive from C14–C15 cleavage and methyl terminus side chain loss (Figure 1B) [36]. This product ion is less abundant in the closest eluting worm F3 prostaglandin compared to PGF3a. Another difference is the product ion m/z 167, which is not found in PGF3a. Therefore, C. elegans F3 class prostaglandins are likely to have a significant structural difference relative to PGF3a. These data indicate that worms synthesize an array of F1, F2, and F3 class prostaglandins, including CePGF1 and CePGF2.

Next, we examined F-series prostaglandins in fat-1(wa9) mutants, fat-4(wa14) mutants, and fat-1(wa9) fat-4(wa14) double mutants. There are several important features of the data. First, all three mutant strains lack EPA and F3 class prostaglandins (Figure 3A). Second, fat-1(wa9) mutants with elevated AA levels have elevated CePGF2 and a second PGF2 analog, called CePGF2b (Figure 3A–3B). Both CePGF2 and CePGF2b are absent in fat-4(wa14) mutants and fat-1(wa9) fat-4(wa14) double mutants that lack AA (Figure 3A–3B). Third, fat-4 mutants with elevated omega-3 AA levels have elevated m/z 193, which is thought to derive from C14–C15 cleavage and methyl terminus side chain loss (Figure 1B) [36]. This product ion is less abundant in the closest eluting worm F3 prostaglandin compared to PGF3a. Another difference is the product ion m/z 167, which is not found in PGF3a. Therefore, C. elegans F3 class prostaglandins are likely to have a significant structural difference relative to PGF3a. These data indicate that worms synthesize an array of F1, F2, and F3 class prostaglandins, including CePGF1 and CePGF2.

We previously showed that fat-3 is required for CePGF2 synthesis, but F1 and F3 class members were not analyzed [20]. A re-examination of fat-3(wa22) mutant extracts using LC-MS/MS shows that CePGF1, CePGF2, CePGF2b, and all F3 class prostaglandins are largely absent in fat-3 mutants (Figure S2A). However, several potential F-series prostaglandins are found, perhaps derived from trace levels of 20:3n3 found in fat-3(wa22) extracts [39]. Overall, the total 20-carbon prostaglandin level in fat-3 mutants is lower than the level in fat-1 fat-4 double mutants. As fat-3 mutants have approximately 12-fold higher 18:3n3 (ALA) than the wild type [39], we hypothesized that 18:3n3 might serve as a substrate for prostaglandin synthesis. The 18-carbon PGF2a metabolite 2,3-Dinor 11β PGF2a has a chromatography RT of 10.7 min, so we predicted a prostaglandin derived from 18:3n3 might have a similar RT (Table 3). An 18:3n3 F-series prostaglandin should have parent ion mass m/z 327, which is 28 Da lighter (-C₆H₄) than PGF2a. LC-MS/MS identified a putative F-series prostaglandin at RT = 10.9 min with considerable similarity to PGF1a, including loss of 44 Da from the parent ion followed by two 18 Da losses (Figure S2B). MRM using mass transition m/z 327/283 shows that this compound and at least one other compound are strongly up-regulated in fat-3 mutants relative to the wild type (Figure S2C). These data add further support to the model that multiple precursors can generate F-series prostaglandins and precursor abundance influences synthesis.

F-series prostaglandins have redundant functions in sperm guidance

The current data provide evidence that F-series prostaglandins have redundant functions. Consistent with this possibility, microinjecting nanomolar concentrations of PGF2a or PGF3a into the uteri of fat-2(wa17) or rme-2(b1008) mutants has an immediate and similar effect on sperm velocity [20]. To further test this model, we supplemented the diets of wild-type, fat-3(wa22) mutant, and fat-1(wa9) fat-4(wa14) double mutant hermaphrodites with AA or EPA. Whereas fat-2 and fat-3 mutants convert AA into EPA, fat-1(wa9) fat-4(wa14) double mutants cannot [18,32,33]. We found that AA or EPA supplementation rescued the sperm guidance defects of fat-2 bacteria, fat-3 mutants, and fat-1 fat-4 double mutants (Figure 2, Figure 5, and Figure 6; P < 0.001). In these experiments, the NA22 bacteria used as a food source do not accumulate PUFAs to high levels (<5% of total membrane lipids),

Table 1. Sperm motility values in wild-type and mutant hermaphrodite uteri.

| Description | Average Velocity (μm/min) | Average Directional Velocity (μm/min) | Reversal Frequency (rev/hr) | N |
|-------------|---------------------------|--------------------------------------|-----------------------------|---|
| 1. Wild type* | 8.62 ± 0.60 | 4.28 ± 0.78 | 1.8 | 88 |
| 2. fat-1(wa9) | 7.97 ± 0.51 | 2.50 ± 0.70 | 9.3 | 42 |
| 3. fat-4(wa14) | 6.50 ± 0.33 | 2.08 ± 0.54 | 5.9 | 60 |
| 4. fat-1(wa9) fat-4(wa14) | 5.68 ± 0.35 | 2.74 ± 0.48 | 5.7 | 55 |
| 5. fat-3(wa22)* | 3.84 ± 0.31 | 2.16 ± 0.42 | 2.5 | 49 |
| 6. fat-2(wa17)* | 3.85 ± 0.22 | 0.47 ± 0.30 | 8.1 | 94 |
| 7. rme-2(b1008)** | 3.68 ± 0.28 | 0.45 ± 0.32 | 9.5 | 75 |
| 8. gfp-4(bn2)** | 3.43 ± 0.25 | -0.23 ± 0.25 | 16.2 | 79 |
| 9. cyp-31A2(tm2711) | 7.63 ± 0.84 | 2.04 ± 1.07 | 3.5 | 26 |

The average velocity, average directional velocity toward the spermatheca, and average reversal frequency were determined for wild-type sperm within the uterus (Zone 2) of wild-type and mutant hermaphrodites. Value ± standard error of the mean is shown.

*Includes new data and published data from Kubagawa et al (2006) [18].

**Published data from Kubagawa et al (2006) for reference [18].

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resulting in modest dietary intake [18]. We have not yet developed methods to measure prostaglandins in supplementation experiments. The data, together with previous and published data, strongly support the hypothesis that F-series prostaglandins derived from different precursors function redundantly to promote sperm guidance.

F-series prostaglandins are not synthesized from D- or E-series intermediates

We next sought to investigate the mechanism by which F-series prostaglandins are synthesized from PUFA precursors. In the mammalian cyclooxygenase-dependent pathway for F2 class prostaglandin synthesis, PGH2, PGD2, and PGE2 are intermediates of PGF2α, 11β PGF2α, and 9β PGF2α (Figure S1) [22,29]. To test whether worm extracts contain these intermediates, we used MRM with mass transition m/z 351/189, which detects PGD2, PGE2, and PGH2 in the low picomolar range. Several peaks of low abundance were identified in wild-type extracts (Figure 3A). If these peaks represent PGD2, PGE2, and PGH2 isomers, then fat-1 loss should increase their levels because fat-1 mutants contain elevated AA and CePGF2. However, this was not the case. The minor peaks observed in wild-type extracts with mass transition m/z 351/189 are eliminated in fat-1 mutants, fat-4 mutants, and fat-1 fat-4 double mutants (Figure 3A). This pattern indicates that these compounds likely require EPA, but not AA for synthesis. In addition, CID of parent ions m/z 351 at RT = 11.4 min and

Figure 3. Prostaglandins in wild-type and fat mutant extracts. (A) MRM chromatograms of wild-type and fat mutant extracts. Liquid chromatography retention time (min) is shown on the X-axis and for selected prostaglandin isomers. Bolded retention times indicate prostaglandin isomers shown in Figure 4. Prostaglandin standard retention times are shown at the top. The dot diagram to the left indicates relative PUFA levels in the corresponding strains [33]. DGLA, dihomo-γ-linolenic acid; O3AA, omega-3 arachidonic acid; AA, arachidonic acid; EPA, eicosapentaenoic acid; cps, counts per second. (B) CePGF2 quantification in wild-type and fat mutant extracts. Error bars are SD. (C) Summary of MRM mass transitions and retention times for chemically synthesized standards. RT, retention time.

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12.6 min in wild-type extracts shows similarity to F3 class prostaglandins, with the exception of the minor product ion \( m/z \) 189 (data not shown). Thus, these compounds are not AA-derived PGD2, PGE2, or PGH2 analogs. These data indicate that worms either do not synthesize PGD2, PGE2, and PGH2 analogs or their concentrations are below the level of detection.

We have previously identified predicted prostaglandin D, E, and F synthases that are required for sperm guidance [20].

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### Table 2. LC-MS/MS data summary for chemically synthesized F-series prostaglandin standards.

| Name            | RT (min)* | [M-H] | \( m/z \) | Key product ions in CID (MS/MS) |
|-----------------|-----------|-------|------------|--------------------------------|
| PGD2            | 12.56     | 351   | 315, 271, 233, 203, 189 |
| PGE2            | 12.23     | 351   | 333, 315, 271, 235, 189, 175, 109 |
| PGH2            | 12.23     | 351   | 333, 315, 271, 235, 189, 175, 113, 109 |
| PGF1\(a\)       | 11.83*    | 355   | 337, 319, 311, 301, 293, 275, 265, 249, 237, 211, 195 |
| 8-iso PGF1\(a\) | 11.34     | 355   | 337, 319, 311, 293, 275, 265, 249, 237, 219, 211, 183 |
| 9\(\eta\)-PGF1\(a\) | 11.36     | 355   | 337, 319, 311, 301, 293, 275, 265, 237, 211, 183 |
| 8-iso 9\(\eta\)-PGF1\(a\) | 11.44     | 355   | 337, 319, 311, 293, 275, 265, 219, 211, 183 |
| 9\(\eta\), 11\(\jmath\)-PGF1\(a\) | 11.99     | 355   | 337, 319, 311, 301, 293, 275, 265, 237, 219, 211, 183 |
| PGF2\(a\)       | 11.73*    | 353   | 335, 317, 309, 291, 273, 263, 247, 209, 193, 171, 165 |
| ent-PGF2\(a\)   | 11.71     | 353   | 335, 317, 309, 291, 273, 247, 209, 193, 191, 171, 165 |
| 11\(\jmath\)-PGF2\(a\) | 11.48     | 353   | 335, 317, 309, 291, 273, 247, 209, 193, 173, 165 |
| 15\(\text{R}\)-PGF2\(a\) | 11.89     | 353   | 335, 317, 309, 291, 273, 247, 209, 193, 171, 165 |
| 8-iso PGF2\(a\) | 11.31     | 353   | 335, 307, 309, 291, 273, 247, 209, 193, 181, 171, 165 |
| 5-trans PGF2\(a\) | 11.60     | 353   | 335, 317, 309, 291, 273, 247, 209, 193, 171, 165, 111 |
| 8-iso 15\(\text{R}\)-PGF2\(a\) | 11.39     | 353   | 335, 317, 309, 291, 273, 263, 247, 209, 193, 171, 165 |
| 9\(\jmath\)-PGF2\(a\) | 11.22     | 353   | 335, 317, 309, 291, 273, 255, 247, 193, 173, 171 |
| PGF2\(b\)       | 11.26     | 351   | 333, 315, 307, 289, 271, 245, 219, 209, 193, 191, 165 |
| 8-iso PGF2\(b\) | 10.83     | 353   | 333, 315, 307, 289, 271, 245, 219, 209, 193, 191, 171 |
| 2,3-Dinor-11\(\jmath\)-PGF2\(a\) | 10.67     | 325   | 261, 245, 227, 219, 173, 163, 153, 145, 137, 113, 107 |
| 19\(\text{R}\)-hydroxy PGF2\(a\) | 9.19      | 369   | 351, 333, 325, 315, 307, 263, 235, 209, 193, 171, 165 |
| 20-hydroxy PGF2\(a\) | 9.13      | 369   | 351, 333, 325, 315, 307, 263, 209, 193, 181, 171, 165 |

Retention time (RT), parent ion mass ([M-H]−), and key product ion masses are shown for prostaglandin (PG) standards.

*Isomers within each prostaglandin class (i.e. PGF2\(a\), isomers) were run together and RTs are directly comparable. PGF1\(a\) and PGF2\(a\) classes were run on different days and a slight RT shift is observed. For example, the RTs for PGF1\(a\) and PGF2\(a\) are indistinguishable when run together.

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R11A8.5 encode glutathione S-transferase enzymes related to PGD and PGE synthases, respectively [20]. *gst*-4 mutants have more severe sperm guidance defects than the mild defects observed in R11A8.5 mutants [20]. To test whether these enzymes are essential for F-series prostaglandin synthesis, we conducted LC-MS/MS on *gst*-4 and R11A8.5 null mutant extracts. In four independent trials, we consistently observed a 25–75% reduction in prostaglandin levels in GST-4 null mutants compared to the wild type (Figure S3). R11A8.5 null mutants had a more complex pattern, with elevated levels of some prostaglandins and reduced levels of others (Figure S3). Null mutations are not currently available for the predicted PGF synthases Y39G8B.2 and C35D10.6. These data indicate that *gst* and R11A8.5 are not essential for F-series prostaglandin synthesis. Instead, they appear to have modulatory roles.

**CePGF2** consists predominantly of ent-PGF$_{2\alpha}$

Although *C. elegans* lacks cyclooxygenase enzymes, our data strongly support the hypothesis that CePGF2 is synthesized from AA. Furthermore, the CID of CePGF2 is almost identical to PGF$_{2\alpha}$ (Figure 4). The product ions $m/z$ 309 and 193 are characteristic of the F$_{2\alpha}$ cyclopentane ring [36]. The loss of 44 Da ($\Delta C_2H_4O$) from the parent ion followed by two 18 Da losses ($\Delta 2\times H_2O$), together with the presence of abundant ion $m/z$ 193, further indicates the compound has three hydroxyl groups at the C9, C11, and C15 positions (Figure 1B). There is no product ion $m/z$ 115 that is abundant in PGF$_{2\alpha}$ isomers with a hydroxyl group at the C5 position [40]. There are sometimes a few differences in minor product ions of unknown composition between PGF$_{2\alpha}$ and CePGF2 from wild-type extracts. However, the CePGF2 decomposition pattern from *fat*-1 mutant extracts, which have increased CePGF2 concentration, is almost identical to PGF$_{2\alpha}$ (Figure 4). Minor differences between CePGF2 and PGF$_{2\alpha}$ spectra could be due to low concentration and co-eluting compounds. We conclude that CePGF2 has chemical composition C$_{20}$H$_{34}$O$_5$ with an F$_{2\alpha}$ cyclopentane ring and hydroxyl groups at positions C9, C11, and C15.

Structural differences in similar compounds often cause changes in chromatographic retention. For example, the RTs of PGF$_{2\alpha}$ and PGF$_{3\alpha}$ differ by 34 seconds, although the only structural difference is an additional double bond at C17 in PGF$_{3\alpha}$ (Figure 1B). Given that CePGF2 and PGF$_{2\alpha}$ have indistinguishable retention times, we asked whether PGF$_{2\alpha}$ stereoisomers are resolved in our HPLC program. The COH bonds at C9, C11, and C15, the C-C bonds at C8 and C12, and the C=C bonds at C5 and C13 can be in different conformations. We analyzed by LC-MS/MS a series of chemically synthesized PGF$_{2\alpha}$ stereoisomers that included conformation changes at the C5, C8, C9, C11, and C15 positions. With the exception of the PGF$_{2\alpha}$ enantiomer (ent-PGF$_{2\alpha}$), all other stereoisomers were separated (Table 3). The RT differences relative to PGF$_{2\alpha}$ ranged from 7 seconds for 5-trans PGF$_{2\alpha}$ to 30 seconds for 9b PGF$_{2\alpha}$. Similar changes were observed with several PGF$_{1\alpha}$ and PGF$_{3\alpha}$ stereoisomers in 5-trans configuration.
reverse-phase chromatography (Table 3). It is noteworthy that CePGF1, CePGF2, PGF1, PGF2, and ent-PGF2 co-elute together and the CID patterns of ent-PGF2 and PGF2 are identical (Figure 3 and Table 3). These results indicate that CePGF2 is PGF2x, a co-eluting stereoisomer such as ent-PGF2x, or a racemic mixture. To distinguish between these possibilities, we developed a normal phase chiral LC-MS/MS method operated in MRM mode (mass transition 353/193) to separate PGF2x and ent-PGF2x. When fat-1(wa9) mutant extract, which contains abundant CePGF2 (Figure 3), was analyzed by chiral LC-MS/MS, the predominant isomer eluted with the same retention time as the ent-PGF2 standard (Figure S4). These results strongly support the hypothesis that CePGF2 consists predominantly of ent-PGF2x. Similarly, CePGF1 may be the enantiomer of PGF1x. Nuclear Magnetic Resonance spectroscopic studies will be necessary for absolute confirmation.

The adult germ line is essential for F-series prostaglandin synthesis

We have previously shown that CePGF2 synthesis requires the RME-2 low-density lipoprotein receptor, which transports PUFAs from the intestine to oocytes [18,20]. These experiments were conducted with extracts from mixed stage worm cultures. To evaluate the extent to which F-series prostaglandins are synthesized in the adult germ line, we synchronized mass worm cultures prior RNAi screen for these genes, we identified H02H12.8 (also called cpd-31A2) [18]. This gene and its paralog cpd-31A3 encode class 4 cytochrome P450 enzymes most closely related to CYP4V2 [41]. Given that cpd-31A2 and cpd-31A3 mRNAs share approximately 92% identity in their coding regions, RNAi might affect both gene products. Previous studies have shown that these two enzymes act redundantly to promote formation of the egg permeability barrier following fertilization [41,42]. To test whether cpd-31A2 and cpd-31A3 are required for sperm guidance, we analyzed their deletion mutants, cpd-31A2(tm2711) and cpd-31A3(tm3224) single mutants are both fertile with no obvious oogenesis defects. However, wild-type sperm fail to target the spermatheca efficiently in the mutants (Figure 8A; P=0.001). Similar results are observed in emb-8 RNAi hermaphrodites (P<0.001). emb-8 encodes a homolog of NADPH cytochrome P450 reductase, which provides electrons for cytochrome P450s. Sperm velocity in cpd-31A2(tm2711) mutants is not significantly different than velocity in the wild type (P=0.05), but directional velocity is decreased (Table 1, lines 1 and 9; P<0.05). Similar results were previously reported using RNAi [18]. These results indicate that cytochrome P450 function is required for sperm guidance.

To investigate the mechanism, we conducted LC-MS/MS analysis of cpd-31A2(tm2711) mutant extracts. MRM in three independent trials showed that F1, F2, and F3 class prostaglandins

Figure 6. Effect of EPA supplementation on sperm guidance in wild-type and fat mutant hermaphrodites. Sperm distribution 1 hr after mating EPA supplemented wild-type or mutant hermaphrodites to non-supplemented wild-type males. A representative image is shown to the left and average zone distribution ± standard deviation is to the right. See Figure 2 for sperm distribution in non-supplemented fat mutants. The spermatheca (S) is outlined in yellow. V, vulva; MT, MitoTracker channel; N, number of gonads scored. Scale bar, 20 μm.

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were consistently elevated in the mutant strain compared to the wild type (Figure 8B). LC-MS analysis in the mass range \( m/z \) 315–360 did not show global elevation of PUFA-derived compounds, demonstrating specificity for F-series prostaglandins (data not shown). Therefore, \( \text{cyp-31A2} \) is not essential for prostaglandin synthesis. A feature of class 4 cytochrome P450s is the ability to hydroxylate PUFAs and prostaglandins at the C19 and C20 positions [43]. We considered the possibility that \( \text{cyp-31A2} \) is required to form hydroxylated F-series prostaglandins. However, no evidence of these compounds can be found using LC-MS/MS (Figure S5). MRM with mass transition \( m/z \) 369/193, which detects 19-hydroxy PGF 2\( \alpha \) and 20-hydroxy PGF 2\( \alpha \) standards, fails to detect these metabolites in wild-type extracts or \( \text{fat-1(wa17)} \) mutant extracts with elevated CePGF2. Similar results were observed using MRM with mass transition \( m/z \) 367/193, which detects hydroxylated PGF 2\( \alpha \) (Figure S5). We conclude that \( \text{CYP-31A2} \) negatively regulates F-series prostaglandin metabolism, likely by affecting a process common to F1, F2, and F3 classes. The increased prostaglandin levels in \( \text{cyp-31A2} \) mutants could be due to continued synthesis in fertilized eggs (see Discussion).

**Discussion**

For fertilization to occur, sperm must sense the environment within the female reproductive tract and locate an oocyte that has recently completed meiotic maturation and ovulation. We have been using *C. elegans* to delineate the mechanisms that guide sperm to maturing oocytes [18,20,21,45]. Results from this study and

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**Figure 7. Prostaglandins in adult wild type and germline-deficient glp-4 mutants.** (A) MRM chromatograms of synchronized 1–2 day adult wild-type and glp-4(bn2) mutants. The F2 class was detected with mass transition \( m/z \) 353/193 and the F3 class was detected with mass transition \( m/z \) 351/193. Liquid chromatography retention time (min) is shown on the X-axis and for selected prostaglandin isomers. Cps, counts per second. (B) CePGF2 quantification in wild-type and glp-4 mutant extracts. Error bars are SD. doi:10.1371/journal.pgen.1003271.g007
previous studies support the following model. In the intestine, dietary fats are converted into PUFAs and incorporated into yolk lipoprotein complexes [18]. Yolk is secreted into the pseudocoelom and flows to the gonad, where it is endocytosed into oocytes by the RME-2 low-density lipoprotein receptor [34]. Yolk provides oocytes with 20-carbon PUFAs that are converted into at least 10 structurally related F-series prostaglandins, independent of cyclooxygenase enzymes. DGLA, AA, and EPA are major precursors for F1, F2, and F3 classes, respectively, that include PGF1\(\alpha\) and PGF2\(\alpha\) isomers. Conserved glutathione S-transferase and cytochrome P450 enzymes modulate oocyte F-series prostaglandin metabolism, although the biochemical mechanisms are not yet clear.

F-series prostaglandins secreted into the reproductive tract function collectively and partially redundantly to promote sperm velocity and directional motility toward the spermatheca. Prostaglandins provide positional information to sperm that allow them to locate and maintain their place in the spermatheca, despite the obstruction of passing fertilized eggs. As maturing oocytes enter the spermatheca during ovulation, they are immediately fertilized. The CYP-31A2 enzyme acts following fertilization to inhibit prostaglandin synthesis and secretion, ensuring that sperm are not attracted to developing embryos. Endocrine signals such as insulin, as well as signals from somatic gonadal cells transduced by gap junctions regulate oocyte prostaglandin metabolism [20,21,45]. Together, these mechanisms help couple nutritional status and...

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**Figure 8. Effect of cytochrome P450 enzymes on sperm guidance and prostaglandin metabolism.** (A) Sperm distribution 1 hr after mating wild-type or mutant hermaphrodites to wild-type males. A representative image is shown to the left and average zone distribution ± standard deviation is to the right. The spermatheca (S) is outlined in yellow. V, vulva; MT, MitoTracker channel; N, number of gonads scored. Scale bar, 20 \(\mu m\). (B) MRM chromatograms of wild-type and cyp-32A2(tm2711) mutants. The F1 class was detected with mass transition m/z 355/311, the F2 class was detected with mass transition m/z 353/193, and the F3 class was detected with mass transition m/z 351/191. Liquid chromatography retention time (min) is shown on the X-axis and for selected prostaglandin isomers. Cps, counts per second.

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oocyte development to reproductive output. New evidence supporting this model is discussed below.

F-series prostaglandin synthesis from multiple precursors

LC-MS/MS analyses of wild-type and fat mutant extracts provide strong evidence that F-series prostaglandins can be synthesized from DGLA, O3AA, AA, and EPA. The concentrations of CePGF1 and CePGF2 in wild-type extracts are similar, although absolute levels of CePGF1 were not determined. The concentrations of F3 class prostaglandins are difficult to estimate because appropriate standards are not available. Given the abundance of EPA in wild-type hermaphrodites and yolk [18,32], the F3 prostaglandins are likely the most abundant.

Precursor levels appear to have a profound impact on prostaglandin synthesis and signaling. For example, fat-1 mutants have approximately 13-fold higher AA levels relative to the wild type [32,33]. We found that CePGF2 levels increase by over three fold, helping to compensate for F3 class prostaglandin loss. LC-MS/MS analysis also detected a putative F-series prostaglandin(s) derived from 18:3n3 in fat-3 mutants. However, F-series prostaglandins derived from 18:3 or 18:4 were not identified in wild-type extracts. These results reveal unexpected complexity in the mechanisms that couple PUFA homeostasis to prostaglandin synthesis. A mutant strain completely deficient in prostaglandin synthesis has not been identified. Hence, the extent to which prostaglandins regulate development, immunity, and reproduction is not known.

A key feature of prostaglandin synthesis in C. elegans is the absence of cyclooxygenase enzymes [20]. The C. elegans genome does not encode cyclooxygenase homologs. Furthermore, drugs that inhibit cyclooxygenase activity do not affect sperm motility [20]. In this study, we show that C. elegans extracts contain PGI2x and PGF2x stereoisomers, but not key PGH2, PGD2, and PGE2 intermediates of cyclooxygenase-dependent pathways. These data provide compelling evidence that PGI2x and PGF2x are synthesized using a cyclooxygenase-independent mechanism.

Prostaglandins can be synthesized via a non-enzymatic mechanism mediated by free radical-initiated peroxidation [30]. However, free radical-initiated peroxidation generates a nonselective mixture of prostaglandin stereoisomers, including 8-iso PGF2 and 8-iso PGE2 that are undetectable in C. elegans. Free radicals could play a role in worm prostaglandin synthesis, but it seems likely that enzymes are involved. The enantiomeric form of PGF2x, which is not formed by cyclooxygenase and cannot be distinguished from PGF2x using standard LC-MS/MS or immunoassays, is the major F2 class prostaglandin in human urine [46] and C. elegans extracts. An attractive possibility is that F-series prostaglandins can be formed by a conserved, yet undiscovered mechanism that does not involve cyclooxygenases.

F-series prostaglandins function redundantly

We show that sperm guidance defects depend on the extent to which PUFA classes are collectively eliminated. As each PUFA class with at least three double bonds can serve as a precursor for prostaglandin synthesis, the data support the model F-series prostaglandins function collectively and partially redundantly. Indeed, supplementation of fat-2(wa17) mutants, fat-3(wa22)
mutants, and fat-1(wa9) fat-4(wa14) mutants with either AA or EPA largely rescues the sperm guidance defects. Similar results were observed when fat-2(wa17) mutants were supplemented with linoleic acid, AA, and EPA and sperm motility was analyzed by time-lapse video microscopy [10]. Microinjecting AA into fat-2 mutant gonads rescues the sperm distribution defect [18]. Finally, microinjection of nanomolar concentrations of PGF\textsubscript{2α} or PGE\textsubscript{2α} into the uterus of fat-2(wa17) or rmo-2(b1008) mutants has an immediate and similar effect on sperm velocity [20]. These results provide strong evidence that F2 or F3 class prostaglandins are sufficient for sperm guidance, yet another class is essential.

F1 class prostaglandins and prostaglandins derived from 18:3n3 are also likely to function redundantly with F2 and F3 classes. In fat-1 fat-4 double mutants, the F1 prostaglandins appear to partially compensate for the absence of F2 and F3 classes. Similarly, 18:3n3-derived prostaglandins appear to partially compensate for 20-carbon prostaglandin loss in fat-3 mutants. Thus, there appears to be tremendous flexibility in the mechanisms that synthesize prostaglandins and transduce their signals. Collective function explains the low abundance of individual prostaglandin isomers in wild-type extracts (relative to mammalian tissues). Why such flexibility and redundancy exist in an animal that generates its own omega-3 and omega-6PUFAs is not clear, but perhaps it provides a buffer against dietary or genetic changes.

The cytochrome P450 CYP-31A2 in sperm guidance

CYP-31A2 is a class 4 cytochrome P450 that is most similar to human CYP4V2, which is implicated in the retinal disease Bietti’s Crystalline Dystrophy [47]. This enzymatic class has a well-characterized function in hydroxylating fatty acids, particularly at the omega-1 and omega-2 positions [43]. Recombinant human CYP4V2 has been shown to hydroxylate EPA and docosahexaenoic acid [48]. In C. elegans, RNAi of CYP-31A1/A2/A3 together causes a reduction in hydroxylated EPA metabolites synthesized from microsomes [49]. We were unable to detect hydroxylated forms of F-series prostaglandins in mixed stage wild-type extracts. Thus, potential substrates for CYP-31A2 are EPA and other PUFAs. LC-MS/MS analysis of cyp-32A2 null mutants indicates that F1, F2, and F3 class prostaglandins are elevated by up to 2-fold. These data support the idea that CYP-31A2 down-regulates prostaglandin synthesis. Time-lapse imaging of sperm motility in cyp-32A2 null mutants and RNAi hermaphrodites is consistent with a gain of signaling activity, as sperm velocity is unaffected [18]. cyp-32A2 might act together with or in parallel to cyp-31A3, which is also required for sperm guidance.

An interesting possibility is that CYP-31A2 functions to down-regulate prostaglandin synthesis in fertilized eggs. Continued prostaglandin production in embryos might disturb positional information (i.e. the maintenance of a gradient in the uterus), a model supported by sperm guidance data. Consistent with this model, RNAi mosaic studies suggest that CYP-31A2 acts in the germ line, not the somatic gonad [18]. In addition to sperm guidance, CYP-31A2 promotes formation of the permeability barrier in fertilized eggs [42]. This barrier is thought to consist of long chain fatty acids conjugated to sugars and prevents the entry and exit of small molecules (<3,000 Da) [42]. Hence, CYP-31A2 might act after fertilization to divert PUFAs away from prostaglandin synthesis to permeability barrier formation. The permeability barrier could also prevent prostaglandins from being secreted into the extracellular environment.

Prostaglandins as conserved sperm guidance cues?

Outside of aquatic species, little is known regarding the mechanisms that guide sperm to oocytes. Human sperm are thought to receive multiple signals that guide them to cumulus-enclosed oocytes, which lie in the highly arborized lumen of the oviduct [9,11,50]. Recent studies have documented that progestosterone and prostaglandins bind to CatSper [9,10,12], a sperm Ca\textsuperscript{2+} channel implicated in hyperactivation and chemotaxis [51,52]. PGE\textsubscript{1} and PGF\textsubscript{2α} activate CatSper at physiological concentrations, suggesting that these interactions are functional [9,12]. Although the role of prostaglandins is not entirely clear, they are synthesized by cumulus-oocyte complexes and affect sperm motility in vitro [13,53,54]. Thus, an attractive scenario is that prostaglandins provide positional information to both human and worm sperm. While there are obvious differences between C. elegans and human reproductive systems, it is possible that some molecular features are shared. Delineating shared versus divergent mechanisms should lead to better understanding of those that are fundamental to animal reproduction.

In summary, we show that a heterogeneous mixture of structurally related F-series prostaglandins promotes sperm guidance in C. elegans. Prostaglandin synthesis does not involve conventional cyclooxygenase pathways, yet PGE\textsubscript{1α} and PGF\textsubscript{2α} stereoisomers are still generated. Furthermore, conserved enzymes such as cytochrome P450s regulates prostaglandin synthesis. The C. elegans model could provide unanticipated insight into the mechanisms and actions of prostaglandins, which are among the most widespread and important signaling molecules in metazoans.

Materials and Methods

C. elegans strains, culture, and RNA–mediated interference

C. elegans strains were maintained at 20°C, unless otherwise indicated, and fed with NA22 Es. coli bacteria, as previously described [18,20]. The fog-2(q71) strain was used for males. The following strains were used: N2 Bristol (wild type), BX24 [fat-1(wa9) II], BX26 [fat-2(wa17) II], BX30 [fat-3(wa22) II], BX17 [fat-4(wa14) IV], BX52 [fat-1(wa9) fat-4(wa14) IV], SS104 [gfp-4(bn2) I], cyp-31A2(tm2711) IV, cyp-31A3(tm3224) IV, and CB4108 [fog-2(q71) I]. The cyp-31A2(tm2711) and cyp-31A3(tm3224) mutants were provided by the Japanese National Bioresource Project. The cyp-31A2(tm2711) strain was backcrossed to the wild type three times. RNAi was performed by the feeding method [55]. HT115 bacterial strains were obtained from the feeding library [56] and sequenced for verification. Wild-type and gfp-4(bn2) mutant hermaphrodites were grown at 16°C and synchronized to the L1 stage using an egg preparation with minor modifications [57]. The larvae were shifted to 25°C and cultured for 2–3 days. For prostaglandin analyses, strains were grown on up to sixty-five 150 mm seeded plates, as previously described [20,21]. Cultures were supplemented with concentrated bacteria as needed to prevent starvation. Worms were washed off plates with M9 buffer, collected in polypropylene tubes, and stored at −80°C for lipid extraction. Tissue was weighed to ensure that equal amounts were analyzed.

Sperm guidance assays

MitoTracker Red CMXRos (Invitrogen) was used to stain fog-2(q71) males, as previously described [18,20]. Briefly, about 150 male worms were transferred to a watch glass with 300 μL M9 buffer. Three microliters of a 1 mM MitoTracker CMXRos solution in DMSO was added to the worm solution and mixed. Males were incubated in the dark for 2–3 hours and then transferred to a seeded plate. After 20 minutes, the males were transferred again to a fresh plate and allowed to recover overnight at 16°C. 10–20~2 day old adult hermaphrodites were anesthetized with 0.1% tricaine and 0.01% tetramisole hydrochloride in gluteraldehyde.
M9 buffer for 30 minutes [58]. The anesthetized hermaphrodites were transferred to a plate containing an ~1 cm drop of bacteria with 50–75 stained males. After 30 minutes of mating, the hermaphrodites were separated from the males and transferred to a fresh seeded plate. To directly observe sperm motility, mated hermaphrodites were mounted immediately onto a 2% agarose pad for time-lapse fluorescence microscopy. DIC and fluorescence images were taken every 30 seconds. Directional velocity toward the spermatheca was measured by creating a straight line through the uterus from the vulva to the spermatheca. The distance traveled along this line from the beginning of a sperm trace to the end was divided by time. Positive values indicate movement toward the spermatheca relative to the starting point. A change in migration direction of greater than 90° within 3 consecutive frames was classified as a reversal. Sperm traces range from a minimum of 2.5 minutes to a maximum of 21 minutes. At least 3 videos from different animals were used for quantification.

To assess sperm distribution, mated hermaphrodites were incubated in the dark at room temperature for 3 days. NA22 L. Lipid extraction

To assess sperm distribution, mated hermaphrodites were incubated in the dark at room temperature for 3 days. NA22

A two sample T-test was used to test for significance. Zone 3 was defined as the region in half. AxioVision software was used to measure distances. A two sample T-test was used to test for significance.

PUFA supplementation

AA or EPA containing plates were prepared as previously described [18]. Briefly, PUFAs were added slowly to cooled media containing 0.1% NP-40. The final PUFA concentration was 200 μM. Plates were kept in the dark at room temperature for 24 hours and then seeded with NA22 E. coli bacteria. Plates were incubated in the dark at room temperature for 3 days. NA22 bacteria accumulate PUFAs in their lipids at ranges from 1–5% [18].

Lipid extraction

For lipid extraction, 2.0 grams of synchronized adult worms or 6.0 grams of mixed-staged worms were used. Hydrophilic lipids were extracted from frozen worm pellets using a liquid-liquid extraction technique [59]. Briefly, frozen worms were suspended with 12 ml of ice cold acetone/saline containing 0.005% butylated hydroxytoluene (BHT) to prevent oxidation. Worms were evenly dispersed into twelve 5 ml self-standing plastic tubes for use in the Bullet Blender 5 homogenizer (Next Advance). 0.7–0.8 ml of 0.5 mm diameter Ceria stabilized zirconium oxide beads were added to each tube and the Bullet Blender was set to 2000 rpm for 5 min) were used for homogenization. Liquid/liquid extraction was performed as described above with the following exception: 0.8 ml of 2.0 mm diameter Ceria stabilized zirconium oxide beads and Bullet Blender speed 9 (5 min) were used for homogenization.

Chromatography and mass spectrometry

LC-MS/MS analyses of commercial standards and tissue extracts were performed as previously described [20,21] using a system consisting of a Shimadzu Prominance HPLC with a refrigerated auto sampler (Shimadzu Scientific Instruments, Inc., Columbia, MD) and an API 4000 (Applied Biosystems/MDS Sciex, Concord, Ontario, Canada) triple quadrupole mass spectrometer. The chromatographic separation was performed on a Synergy hydro RP-C18 column pre-equilibrated with 0.1% formic acid. The mobile phase consists of 0.1% formic acid [A] and acetonitrile containing 0.1% formic acid [B] and was pumped at a flow rate of 0.2 ml/min. The gradient started with 10% B and went up to 80% B from 0–11 min, 80–100% B from 11–14 min and returned back to 10% B at 16 min. The column effluent was introduced into the mass spectrometer using an ESI interface operating in negative ion mode. Nitrogen was used as a nebulizer and curtain gas (CUR = 10). The collision gas, collision energy and temperature were set at 10, −55 eV and 600°C, respectively. The LC-MS/MS system was controlled by BioAnalyist 1.4.2 software. For comparative analysis of different strains, the extracts and standards were run consecutively.

To determine CePGF2 concentration, a stock solution of PGF2a (1 μg/ml in 80% MeOH) was serially diluted with the same solvent to obtain 1000, 100, 10, 1, 0.1 ng/ml concentrations. The samples were analyzed by the MRM method. The standard curve exhibited excellent linearity in the range of concentration 0.1–1000 ng/ml with a correlation coefficient >0.99. Average CePGF2 concentration was calculated from three MRM analyses using two independent sample extractions.

Separation of PGF2a stereoisomers that co-elute in reverse-phase liquid chromatography was achieved by following a previously published method with modification [46]. Separation was carried out on a ChiralPak AD-H column (4.6 mm×250 mm, 5 μm, Chiral Technologies, Exton, PA) with 12.5% ethyl alcohol and 12.5% isopropyl alcohol in hexane containing 0.1% formic acid at 1 ml/min. MRM was carried out in atmospheric pressure chemical ionization (APCI) negative ion mode with mass transition m/z 353/193. The instrument conditions were optimized with the collision gas, collision energy and temperature were set at 10, −35 eV and 450°C, respectively.
Supporting Information

Figure S1 Mammalian F2 class prostaglandin synthesis. (A) Cyclooxygenase-dependent pathways. (B) Structures of arachidonic acid and PGF2a.

Figure S2 Prostaglandins in fat-3 mutant. (A) MRM chromatograms of wild-type and fat-3(wa22) mutant extracts. The F1 class was detected with mass transition m/z 355/311, the F2 class was detected with mass transition m/z 353/193, and the F3 class was detected with mass transition m/z 351/193. Liquid chromatography retention time (min) is shown on the X-axis and for major prostaglandin isomers, Cps, counts per second. (B) LC-MS/MS of chemically synthesized PGF13 compared to a putative F-series prostaglandin derived from 18:3n3 in fat-3 mutant extracts. Red color indicates ions shared by the standard and the unknown prostaglandin, after subtracting the mass difference between DGLA and 18:3n3 (28 Da). Blue color indicates ions that are not shared. m/z is on the X-axis. (C) MRM chromatograms using the mass transition m/z 327/283. 2,3-Dinor-11β-PGF2a is an 18-carbon PGF2a metabolite.

Figure S3 F2 class prostaglandins in gst-4(ok2358) and R11A8.5(ok3316) mutant extracts. MRM chromatograms using the mass transition m/z 353/193, gst-4 and R11A8.5 encode glutathione S-transferases with sequence similarities to PGD and PGE synthases, respectively. Liquid chromatography retention time (min) is shown on the X-axis and for major prostaglandin isomers. Cps, counts per second.

Figure S4 The PGF2a, enantiomer co-clutes with the predominant F2 class prostaglandin in fat-1 mutant extracts using chiral chromatographic separation. Normal phase chiral LC-APCI-MS/MS chromatograms operated in MRM with mass transition m/z 353/193. Chromatograms of chemically synthesized standards (top) and mixed staged fat-1(wa9) mutant extract (bottom) are shown.

Figure S5 Absence of hydroxylated F-series prostaglandins in wild-type worm extracts. (A) MRM chromatograms of mixed staged fat-1(wa9) mutant extracts. The mass transition m/z 353/193 was used to detect F2 class prostaglandins and mass transition m/z 369/193 was used to detect hydroxylated forms, such as 20-hydroxy PGF2a. Liquid chromatography retention time (min) is shown on the X-axis and for major prostaglandin isomers. The retention times for 20-hydroxy PGF2a and 19-hydroxy PGF2a are 9.13 min and 9.19 min, respectively. Cps, counts per second. (B) MRM chromatograms of mixed staged wild-type extracts. The mass transition m/z 351/193 was used to detect F3 class prostaglandins and mass transition m/z 367/193 was used to detect hydroxylated forms. Cps, counts per second.

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Author Contributions

Conceived and designed the experiments: MAM JKP. Performed the experiments: HDH DD JKP. Analyzed the data: HDH JKP MAM. Wrote the paper: MAM HDH.

References

1. Evans JP, Florman HM (2002) The state of the union: the cell biology of fertilization. Nat Cell Biol 4 Suppl: e57–63.
2. Marcello MR, Singaravelu G, Singerson A (2013) Fertilization. Adv Exp Med Biol 737: 321–350.
3. Gadella BM, Evans JP (2011) Membrane fusions during mammalian fertilization. Adv Exp Med Biol 713: 65–80.
4. Han SM, Cottee PA, Miller MA (2010) Sperm and oocyte communication mechanisms controlling C. elegans fertility. Dev Dyn 239: 1263–1281.
5. Kaupp UB, Kashikar ND, Weyand I (2008) Mechanisms of sperm chemotaxis. Biochem Soc Trans 38: 1270–1274.
6. Guerrero A, Wood CD, Nishigaki T, Carneiro J, Darszon A (2010) Tuning sperm chemotaxis. Biochem Soc Trans 38: 1270–1274.
7. Chang H, Suarez SS (2010) Rethinking the relationship between hyperactivation and chemotaxis in mammalian sperm. Biol Reprod 83: 507–513.
8. Eisenbach M, Giojalas LC (2006) Sperm guidance in mammals - an unpaved road to the egg. Nat Rev Mol Cell Biol 7: 276–285.
9. Edmonds JW, McKinney SL, Prasain JK, Miller MA (2011) The gap junctional protein INX-1 functions in oocyte precursors to promote C. elegans sperm guidance. Dev Biol 359: 47–58.
10. Evans JP, Florman HM (2002) Fertilization. Adv Exp Med Biol 737: 321–350.
11. Linkho PV, Botchkina IL, Kirichok Y (2011) Progesterone activates the principal Ca2+ channel of human sperm. Nature 471: 307–311.
12. Strunk T, Goodwin N, Brenker C, Kashikar ND, Weyand I, et al. (2011) The CatSper channel mediates progesterone-induced Ca2+ influx in human sperm. Nature 471: 307–316.
13. Teves ME, Guidobaldi HA, Unates DR, Sanchez R, Miska W, et al. (2009) Molecular mechanism for human sperm chemotaxis mediated by progesterone. PLoS ONE 4: e82031.
14. Brenker C, Goodwin N, Weyand I, Kashikar ND, Naruse M, et al. (2012) The CatSper channel: a polymodal chemosensor in human sperm. EMBO J 31: 1654–1665.
15. Schuetz AW, Dubin NH (1981) Progesterone and prostaglandin secretion by ovulated rat cumulus cell-oocyte complexes. Endocrinology 108: 457–463.
16. L’Hernautil SW (2009) The genetics and cell biology of spermatogenesis in the nematode C. elegans. Mol Cell Endocrinol 306: 59–65.
17. Kim S, Spike C, Greenstein D (2013) Control of Oocyte Growth and Meiotic Maturation in Caenorhabditis elegans. Adv Exp Med Biol 757: 277–320.
18. Hill KL, L’Hernautil SW (2001) Analyses of reproductive interactions that occur after heterospecific matings within the genus Caenorhabditis. Dev Biol 232: 105–114.
19. Kubagawa HM, Watts JL, Corrigan C, Edmond JW, Snul E, et al. (2006) Oocyte signals derived from polysaturated fatty acids control sperm recruitment in vivo. Nat Cell Biol 8: 1143–1148.
20. Ward S, Carrell JS (1979) Fertilization and sperm competition in the nematode Caenorhabditis elegans. Dev Biol 73: 304–321.
21. Edmonds JW, Prasain JK, Dorand D, Yang Y, Huang HD, et al. (2010) Insulin/FOXO signaling regulates ovarian prostaglandins critical for reproduction. Dev Cell 19: 839–851.
22. Edmonds JW, McKinney SL, Prasain JK, Miller MA (2011) The gap junctional protein INX-1 functions in oocyte precursors to promote C. elegans sperm guidance. Dev Biol 359: 47–58.
23. Funk CD (2001) Prostaglandins and leukotrienes: advances in eicosanoid biology. Science 294: 1871–1875.
24. Burr GO, Burr MM (1929) A new deficiency disease produced by the rigid exclusion of fat from the diet. J Biol Chem 82: 345–367.
25. Burr GO, Burr MM (1930) On the nature and role of the fatty acids essential for nutrition. J Biol Chem 86: 587–621.
26. Van Domburg RT, Bakhle YS, Potting RM (1998) Cyclooxygenases 1 and 2. Annu Rev Pharmacol Toxicol 38: 97–120.
27. Wathes DC, Aabyasekara DR, Aitken RJ (2007) Polyunsaturated fatty acids in pregnancy and chemotaxis in mammalian sperm. Biol Reprod 76: 507–513.
32. Watts JL, Browse J (2002) Genetic dissection of polyunsaturated fatty acid synthesis in Caenorhabditis elegans. Proc Natl Acad Sci U S A 99: 5854–5859.
33. Kahn-Kirby AH, Dantzker JL, Apicella AJ, Schafer WR, Browse J, et al. (2004) Specific polyunsaturated fatty acids drive TRPV-dependent sensory signaling in vivo. Cell 119: 889–900.
34. Beanan MJ, Strome S (1992) Characterization of a germ-line proliferation mutation in C. elegans. Development 116: 753–768.
35. Murphy RC, Barkley RM, Zemski Berry K, Hankin J, Harrison K, et al. (2005) Electrospray ionization and tandem mass spectrometry of eicosanoids. Anal Biochem 346: 1–42.
36. Murphy RC, Barkley RM, Zemski Berry K, Hankin J, Harrison K, et al. (2005) Electrospray ionization and tandem mass spectrometry of eicosanoids. Anal Biochem 346: 1–42.
37. Masoodi M, Nicolaou A (2006) Lipidomic analysis of twenty-seven prostanoids and isoprostanes by liquid chromatography/electrospray tandem mass spectrometry. Rapid Commun Mass Spectrom 20: 3023–3029.
38. Kulas J, Schmidt C, Rothe M, Schunck WH, Menzel R (2008) Cytochrome P450-specific delta9 fatty acid desaturase from Caenorhabditis elegans. Biochem Biophys Res Commun 372: 263–269.
39. Waugh RJ, Morrow JD, Roberts LJ, 2nd, Murphy RC (1997) Identification and relative quantitation of F2-isoprostane regioisomers formed in vivo in the rat. Free Radic Biol Med 23: 943–954.
40. Benenati G, Penkov S, Muller-Reichert T, Entchev EV, Kurzchalia TV (2009) Two cytochrome P450s in Caenorhabditis elegans are essential for the organization of eggshell, correct execution of meiosis and the polarization of embryos. Mech Dev 126: 382–393.
41. Olson SK, Greenan G, Desai A, Muller-Reichert T, Oegema K (2012) Hierarchical assembly of the eggshell and permeability barrier in C. elegans. J Cell Biol 198: 731–748.
42. Hsu MH, Savas U, Griffin KJ, Johnson EF (2007) Human cytochrome p450 family 4 enzymes: function, genetic variation and regulation. Drug Metab Rev 39: 515–538.
43. Hsu MH, Savas U, Griffin KJ, Johnson EF (2007) Human cytochrome p450 family 4 enzymes: function, genetic variation and regulation. Drug Metab Rev 39: 515–538.
44. Yin H, Gao L, Tai HH, Murphy LJ, Porter NA, et al. (2007) Urinary prostaglandin P2alpha is generated from the isoprostane pathway and not the cyclooxygenase in humans. J Biol Chem 282: 329–336.
45. McCarter J, Bartlett B, Dang T, Schell T (1999) On the control of oocyte meiotic maturation and ovulation in Caenorhabditis elegans. Dev Biol 205: 111–128.
46. Golovko MY, Murphy LJ (2008) An improved LC-MS/MS procedure for brain prostanooid analysis using brain fixation with head-focused microwave irradiation and liquid-liquid extraction. J Lipid Res 49: 893–902.