Dephosphorylation and inactivation of NPR2 guanylyl cyclase in granulosa cells contributes to the LH-induced decrease in cGMP that causes resumption of meiosis in rat oocytes

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ABSTRACT

In mammals, the meiotic cell cycle of oocytes starts during embryogenesis and then pauses. Much later, in preparation for fertilization, oocytes within preovulatory follicles resume meiosis in response to luteinizing hormone (LH). Before LH stimulation, the arrest is maintained by diffusion of cyclic (c)GMP into the oocyte from the surrounding granulosa cells, where it is produced by the guanylyl cyclase natriuretic peptide receptor 2 (NPR2). LH rapidly reduces the production of cGMP, but how this occurs is unknown. Here, using rat follicles, we show that within 10 min, LH signaling causes dephosphorylation and inactivation of NPR2 through a process that requires the activity of phosphoprotein phosphatase (PPP)-family members. The rapid dephosphorylation of NPR2 is accompanied by a rapid phosphorylation of the cGMP phosphodiesterase PDE5, an enzyme whose activity is increased upon phosphorylation. Later, levels of the NPR2 agonist C-type natriuretic peptide decrease in the follicle, and these sequential events contribute to the decrease in cGMP that causes meiosis to resume in the oocyte.

KEY WORDS: Guanylyl cyclase, Luteinizing hormone, Meiosis, Natriuretic peptide receptor, Ovarian follicle, Phosphorylation

INTRODUCTION

Oocytes in mammalian preovulatory follicles are held in meiotic prophase arrest by cyclic (c)GMP that is produced in the granulosa cells surrounding the oocyte, which then diffuses into the oocyte through gap junctions (Norris et al., 2009; Vaccari et al., 2009). Luteinizing hormone (LH), which is released from the anterior pituitary during each reproductive cycle, acts on a G-protein coupled receptor in the outer granulosa cells of the follicle (Bortolussi et al., 1977, Rajagopalan-Gupta et al., 1998) (see Fig. 1A), to relieve the inhibition of the prophase-to-metaphase transition. LH acts by lowering the levels of cGMP in the granulosa cells, thus reducing cGMP in the oocyte. Although this regulatory system is best understood in mice (Norris et al., 2009; Vaccari et al., 2009; Zhang et al., 2010; Kawamura et al., 2011; Robinson et al., 2012; Tsuji et al., 2012; Geister et al., 2013), there is evidence that similar mechanisms operate in other mammals (Törnell et al., 1991; Peng et al., 2013; Hiradate et al., 2014; Zhang et al., 2014), including women (Kawamura et al., 2011; Liu et al., 2014). The LH-induced resumption of the meiotic cell cycle leads into a series of events by which the oocyte matures to become a fertilizable egg (Conti et al., 2012; Clift and Schuh, 2013; Holt et al., 2013; Mehlmann, 2013; Hunzicker-Dunn and Mayo, 2014).

As a consequence of the initial cGMP decrease in the mural granulosa cells, cGMP diffuses out of the oocyte into the surrounding granulosa cells, and the decrease in cGMP in the oocyte re-initiates the meiotic cell cycle (Norris et al., 2009; Vaccari et al., 2009). The direct effect of the cGMP decrease in the oocyte is to relieve the inhibition of the cAMP phosphodiesterase PDE3A, which is competitively inhibited by cGMP. Thus, the cGMP reduction causes the levels of cAMP to decrease. cAMP, which is produced in the oocyte (Mehlmann et al., 2002, 2004; Horner et al., 2003; Hinckley et al., 2005; Ledent et al., 2005), maintains meiotic prophase arrest through activation of protein kinase A (PKA) (Bornslaeger et al., 1986; Kovo et al., 2006). PKA activity stimulates the CDC25B phosphatase and stimulates the WEE1B and MYT1 kinases, and this keeps the CDK1 kinase that controls the prophase-to-metaphase transition phosphorylated and inactive (Conti et al., 2012; Holt et al., 2013; Mehlmann, 2013). When LH signaling reduces oocyte levels of cGMP, and in turn cAMP, meiosis resumes.

cGMP is produced in the granulosa cells of mice by natriuretic peptide receptor 2 (NPR2), also known as guanylyl cyclase-B (Zhang et al., 2010; Kawamura et al., 2011; Robinson et al., 2012; Tsuji et al., 2012; Geister et al., 2013). NPR2 appears to be the only relevant guanylyl cyclase in mouse follicles because follicle-enclosed oocytes in the ovaries of mice with non-functional NPR2 resume meiosis spontaneously (Zhang et al., 2010; Tsuji et al., 2012; Geister et al., 2013) and an inactivating mutation in NPR2 reduces the cGMP content of the follicle to an undetectable level (Geister et al., 2013). Gene expression analysis and enzyme assays have indicated that other guanylyl cyclases are present at levels much lower than those of NPR2 (Robinson et al., 2012). NPR2 is located throughout the granulosa cells of the follicle, in both the mural granulosa and the cumulus cells directly surrounding the oocyte; it is not expressed in the oocyte (Gutkowski et al., 1999; Zhang et al., 2010). The agonist of NPR2, C-type natriuretic peptide (CNP), released from a precursor protein that is encoded by the Nppc gene is produced only in the mural granulosa cells (Zhang et al., 2010).

NPR2 is a single transmembrane-spanning enzyme that is activated by the binding of CNP to its extracellular domain (Potter et al., 2006; Potter, 2011). In order for the CNP activation signal to be transmitted to the catalytic domain, the juxtamembrane intracellular region of NPR2 must be phosphorylated on some
combination of five serine residues and two threonine residues that have been identified as regulatory (Potter, 1998; Potter and Hunter, 1998; Yoder et al., 2010, 2012). However, unlike many growth factor receptors, NPR2 phosphorylation is not increased upon binding to its agonist CNP (Potter, 1998). Thus, there are at least two separate mechanisms by which signaling pathways could increase or decrease the guanylyl cyclase activity of NPR2 — changing the amount of CNP or changing the level of receptor phosphorylation.

LH signaling is known to decrease the amount of CNP in rat and mouse ovaries (Jankowski et al., 1997; Kawamura et al., 2011; Robinson et al., 2012; Liu et al., 2014) and in human and porcine follicular fluid (Kawamura et al., 2011; Zhang et al., 2014); the decrease in the levels of CNP is associated with a decrease in Nppc mRNA (Kawamura et al., 2011; Tsuji et al., 2012; Liu et al., 2014). However, in the mouse ovary, where the kinetics are best characterized, the CNP decrease is first detected at 2 h (Robinson et al., 2012; Liu et al., 2014), whereas the decrease in cGMP is detected at 15 to 20 min (Norris et al., 2010; Liu et al., 2014). Guanylyl cyclase activity in mouse follicle membranes decreases to approximately half of the basal level at 20 min after LH application, and this is independent of any change in CNP (Robinson et al., 2012; Liu et al., 2014). Cultured human granulosa cells also show a rapid decrease in cGMP production, measured in the presence of a constant concentration of CNP (Liu et al., 2014).

The mechanism underlying this early decrease in guanylyl cyclase activity is unknown. Here, we show that the rapid reduction in NPR2 activity in rat follicles in response to LH signaling is caused by the dephosphorylation of NPR2, which is mediated by a process that requires the activity of the protein phosphatases of the phosphoprotein phosphatase (PPP) family, the most likely candidates being PPP1, PPP2 and/or PPP6. The rapid dephosphorylation of NPR2 is accompanied by a rapid phosphorylation of the cGMP phosphodiesterase PDE5 (also known as PDE5A), an enzyme whose activity is increased upon phosphorylation. Later, CNP levels decrease in the follicle, and these sequential events contribute to the decrease in cGMP that causes meiosis to resume in the oocyte.

RESULTS

LH signaling reduces NPR2 activity and cGMP content in rat ovarian follicles

Previous studies demonstrating an LH-induced decrease in guanylyl cyclase activity in ovarian follicles have been conducted using mice (Robinson et al., 2012), but the amount of protein that can be obtained from mouse follicles is small. We therefore tested whether a similar regulatory system operates in rats, from which an order of magnitude more follicle protein per animal can be obtained, making analysis of changes in phosphorylation feasible.

To test whether LH causes a decrease in NPR2 guanylyl cyclase activity in rat follicles, and to investigate the time course of the decrease as a basis for subsequent mechanistic studies, isolated preovulatory rat follicles were incubated for various times with or without LH. Because NPR2 is located in the plasma membrane,
the follicles were then homogenized to obtain a crude membrane fraction. The membranes were assayed for guanylyl cyclase activity, with and without the NPR2 agonist CNP; CNP-dependent activity indicates the activity of NPR2 (Fig. 1B,C). By 30 min after LH exposure, the CNP-dependent guanylyl cyclase activity decreased to 75% of the initial level and stayed at this reduced level for at least 4 h, with no additional change (Fig. 1B,C). Approximately 40% of the decrease to the plateau level had occurred by 10 min (Fig. 1C). No change in CNP-dependent guanylyl cyclase activity was seen in follicles incubated for 4 h without LH (Fig. 1C). The cGMP content of the follicle also decreased rapidly in response to LH—most of the change had occurred by 10 min (Fig. 1D).

The LH-induced decrease in CNP-dependent guanylyl cyclase activity was not explained by a reduction in the affinity of NPR2 for CNP because the fraction by which the activity decreased in response to LH, when it was measured in the presence of 50 nM CNP, was similar to that measured in the presence of 1 μM CNP (Fig. 1B). LH had no effect on guanylyl cyclase activity when it was measured in the presence of 1% Triton X-100 and 5 mM MnCl₂ (Fig. 1B), a condition that activates guanylyl cyclase independently of natriuretic peptide and phosphorylation and that is indicative of guanylyl cyclase protein levels (Potter and Hunter, 1999; Abbey and Potter, 2003). Because the predominant guanylyl cyclase in ovarian follicles is NPR2, this indicates that the LH-induced decrease in CNP-stimulated activity is not due to a decrease in the amount of NPR2 protein.

LH signaling causes rapid dephosphorylation of NPR2

To investigate whether the rapid LH-induced decrease in NPR2 activity correlated with a decrease in NPR2 phosphorylation, we first considered the use of ³²PO₄ metabolic labeling. However, because the expression level of NPR2 in native tissues is low, this approach has only been feasible for overexpressing cells (Potter, 1998). The NPR2 fraction of the total cell protein in ovarian follicles is 5% of that in the 3T3 cells which had been transfected with NPR2 and used in previous ³²PO₄ labeling studies, based on the relative levels of enzyme activity (Potter, 1998). To obtain the same amount of NPR2 protein used in the transfected 3T3 cell studies, this would have required ~10 mg of follicle membrane protein, corresponding to dissection of follicles from ~30 rats, for a single gel lane. Likewise, detection of changes in phosphorylation by mass spectrometry (e.g. Cargnello et al., 2012) would require much more NPR2 protein than is practical to obtain from rat follicles.

We then investigated whether an LH-induced shift in NPR2 migration could be detected using standard SDS-PAGE, as a previous study showed that dephosphorylation results in a small shift in the electrophoretic migration of the closely related guanylyl cyclase natriuretic peptide receptor 1 (NPR1) (Potter and Garbers, 1992). In standard SDS-PAGE gels, NPR2 from rat follicles migrated as a predominant band at the same position as rat NPR2 from stably transfected HEK-293T cells, at the typically observed size of ~150 kDa (Potter, 1998). The fuzziness of this band is primarily due to multiple glycosylation sites (Koller et al., 1993). There was also a minor band at ~115 kDa, which is thought to represent the polypeptide chain that has not been post-translationally modified (Koller et al., 1993; Potter, 1998) (Fig. 2A). However, treatment with LH did not result in a consistent shift in the electrophoretic migration of NPR2 under the conditions that we used (Fig. 2A).

As an alternative approach to investigate whether LH exposure led to dephosphorylation of NPR2, we used gels including Phos-tag acrylamide and MnCl₂. In these gels, phosphorylated proteins transiently interact with the Mn²⁺-Phos-tag complex, which retards their migration relative to less-phosphorylated or non-phosphorylated forms (Kinoshita et al., 2006, 2012; McTague et al., 2012; Yu et al., 2013). NPR2 was purified from the membranes by using sequential immunoprecipitation and fractionation on gels containing Phos-tag acrylamide and was visualized by western blotting. In these gels, NPR2 from untreated follicles migrated as multiple bands, indicating the presence of multiple phosphorylated forms of the enzyme (Fig. 2B; see also Fig. 3A, Fig. 4A). Although molecular weight standards are not accurate indicators of relative molecular mass on a Phos-tag gel, they are useful as descriptive markers; the NPR2 bands migrated in a region that extended from above the 150 kDa marker to approximately the 250 kDa marker.

Controls that were performed using pre-immune serum (supplementary material Fig. S1), and co-migration of the immunoreactive bands from follicles with those from a HEK cell line that stably expressed NPR2 (Fig. 2A, Fig. 3A), validated the specificity of the antibody. The relative amount of NPR2 staining in a sample of follicle membranes was 2.7% of that in a sample from overexpressing HEK cells (normalized to the amount of membrane protein loaded per lane), and this corresponded closely to the relative amount of NPR2 activity (2-4% of that reported for this cell line, Robinson and Potter, 2011; Robinson and Potter, 2012), further confirming the antibody specificity. The use of Phos-tag gel migration as an indicator of the phosphorylation state of NPR2 was validated by showing that incubation of follicle membranes under conditions that promoted or inhibited phosphatase activity resulted in faster or slower migrating forms of NPR2 (Fig. 2E-G).

Exposure of the follicles to LH dramatically compressed the majority of the NPR2 into a predominant lower band, indicating dephosphorylation (Fig. 2B; see also Fig. 3A, Fig. 4A). The shift in migration was quantified by measuring the total amount of NPR2 immunostaining in the upper region relative to that in the region of the lower band (Fig. 2C,D). Most of the decrease in the relative amount of immunoreactive material in the upper region had occurred by 10 min after LH exposure, and a minimum level was reached at 30 min (Fig. 3A, B). The dephosphorylation persisted for at least 4 h (Fig. 3A, B), as did the reduction of NPR2 activity (Fig. 1C).

No decrease in NPR2 phosphorylation occurred in follicles that had been incubated for 4 h in the absence of LH (Fig. 3A, B). When LH was applied for 30 min and then washed out, the NPR2 dephosphorylation and activity measured 4 h later were the same as when LH was present continuously (two independent experiments, data not shown). Based on measurements of the total immunoreactive protein, no change in the total amount of NPR2 protein was detected when comparing membranes from follicles with or without treatment with LH (Fig. 3C). This conclusion is also supported by our earlier finding that LH did not affect the amount of guanylyl cyclase activity measured in the presence of Triton X-100 and MnCl₂ (Fig. 1B).

Most of the decrease in NPR2 phosphorylation had occurred by 10 min after application of LH (Fig. 3B), whereas only approximately 40% of the decrease in guanylyl cyclase activity had occurred by this time, compared with that measured at 30 min (Fig. 1C). The cause of the delay between the decreases in NPR2 phosphorylation and enzyme activity is unknown, but one possibility is that dephosphorylation of a particular serine or threonine residue that is crucial for the activity decrease occurs late in a series of dephosphorylation events, all of which are detected by the Phos-tag method. Consistent with this hypothesis, the individual phosphorylation sites in NPR2 have varying effects on the activity of the enzyme (Potter and Hunter, 1998; Yoder et al., 2012).
The LH-induced decrease in NPR2 activity is prevented by inhibiting NPR2 dephosphorylation with PPP-family protein phosphatase inhibitors

To test whether preventing the LH-induced dephosphorylation of NPR2 inhibits the decrease in NPR2 activity, and to begin to distinguish which of the approximately 30 known serine-threonine phosphatase catalytic subunits are required for dephosphorylation of NPR2, we treated the follicles with specific phosphatase inhibitors. Serine-threonine phosphatases belong to two main families – PPP or PPM (protein phosphatase Mg2+ or Mn2+ dependent); in addition, there is a smaller family (FCP) whose only known function is to dephosphorylate RNA polymerase II (Cohen, 2004; Swingle et al., 2007). Among these, only the PPP family is inhibited by the natural toxins cantharidin and okadaic acid (Swingle et al., 2007; Pereira et al., 2011). PPP family phosphatases (also known as PP2C) are not inhibited by 100 μM cantharidin (Li et al., 1993) or by 10 μM okadaic acid (Wang et al., 1995).

Pre-incubation of follicles for 1 h with cantharidin (100 μM) or okadaic acid (10 μM) prevented the LH-induced dephosphorylation and decrease in guanylyl cyclase activity of NPR2 (Fig. 4; supplementary material Fig. S2). These results indicate that the activity of PPP family phosphatases is required for the dephosphorylation of NPR2 in response to LH and that dephosphorylation is required for the LH-induced decrease in NPR2 activity. Among the PPP family phosphatases, our results with cantharidin argue against an important function for PPP3 (also known as PP2B or calcineurin) because PPP3 is insensitive to 100 μM cantharidin (Honkanen, 1993; Pereira et al., 2011).

These toxin results do not distinguish between other PPP family phosphatases because PPP1, PPP2, PPP4, PPP5 and PPP6 are all inhibited by the concentrations of cantharidin and okadaic acid that we applied to the follicles (Swingle et al., 2007; Pereira et al., 2011). Other toxins with greater specificities for particular PPP-family phosphatases (fostriecin, cytotatin and rubratoxin) were also tested and found to have no consistent effects under the conditions used.
However, these results were not definitive because we could not determine whether significant amounts of these toxins penetrated into the cytoplasm of the follicle cells.

**The LH-induced decrease in follicle cGMP content is attenuated by inhibiting the decrease in NPR2 guanylyl cyclase activity**

To determine the effect of preventing the LH-induced decrease in NPR2 guanylyl cyclase activity on the LH-induced cGMP decrease, we measured the cGMP content of follicles that had been pre-treated with 100 μM cantharidin for 1 h and then treated for 30 min with LH. Without cantharidin pre-treatment, LH caused cGMP content to decrease to 7% of the control level (Fig. 1D and Fig. 4D). Cantharidin pre-treatment attenuated the LH-induced decrease – cGMP content only decreased to 56% of the control level (Fig. 4D).
The effect of phosphatase inhibition on the LH-induced decrease in cGMP content is consistent with the conclusion that the LH-induced decrease in NPR2 phosphorylation and guanylyl cyclase activity contributes to the cGMP decrease. However, the finding that phosphatase inhibition did not completely prevent the LH-induced cGMP decrease, whereas it did completely prevent the LH-induced decrease in NPR2 phosphorylation and activity (Fig. 4A-C), indicates that LH must also induce another change in the follicle, such as an increase in cGMP phosphodiesterase activity. The finding that most of the cGMP decrease had occurred by the 10 min time point after treatment with LH (Fig. 1D), whereas only 40% of the NPR2 activity decrease had occurred by this time point (Fig. 1C), also suggests that LH induces an increase in cGMP phosphodiesterase activity in parallel with the decrease in NPR2 guanylyl cyclase activity.

**LH signaling causes phosphorylation of the cGMP phosphodiesterase PDE5**

An important component of the cGMP phosphodiesterase activity in mouse granulosa cells is contributed by PDE5 (Vaccari et al., 2009). PDE5 activity is stimulated by phosphorylation (Corbin et al., 2000; Rybalkin et al., 2002); therefore, we examined whether LH signaling increased PDE5 phosphorylation in rat follicles. In gels without Phos-tag, PDE5 migrated as a band at ~90 kDa, and treating follicles with LH did not change this migration (Fig. 5A). However, in Phos-tag-containing gels, PDE5 migrated as two separate bands, and treatment of the follicles with LH for 10 or 30 min increased the fraction of PDE5 in the upper band, indicating phosphorylation (Fig. 5B,C). PDE5 remained phosphorylated until at least 4 h after stimulation with LH (data not shown). In *in vitro* phosphorylation of PDE5 in follicle lysates with the catalytic subunit of PKA also caused a shift of PDE5 to the upper band, confirming that the LH-induced shift was due to phosphorylation (Fig. 5D).

Although the effect of LH on the cGMP phosphodiesterase activity of PDE5 in rat follicles is unknown, the rapid phosphorylation of PDE5 in response to LH is expected to increase PDE5 activity based on *in vitro* studies (Corbin et al., 2000; Rybalkin et al., 2002). An increase in PDE5 activity could contribute to the LH-induced cGMP decrease in the follicle, acting in parallel with the decrease in the production of cGMP that results from the dephosphorylation of the NPR2 guanylyl cyclase.

**Analysis of PPP family phosphatase gene expression in granulosa cells**

To distinguish the PPP-family phosphatases that might contribute to the dephosphorylation of NPR2, we investigated which of these are expressed in granulosa cells. The rat genome contains 13 PPP-family genes, encoding seven subfamilies of PPP-family phosphatases (Pereira et al., 2011). We detected mRNA encoding all of these phosphatases, although the fractions of those encoding PPP4, PPP5 and PPP7 were each <2% of the total (Fig. 6). mRNA encoding PPP1, PPP2, PPP3 and PPP6 constituted 61%, 11%, 19% and 6% of the total, respectively. Although the amount of mRNA is not directly proportional to the amount of the protein they encode, and although the localization of a particular phosphatase could affect its functional significance, these measurements point to a role for PPP1, PPP2, PPP3 and/or PPP6, rather than PPP4, PPP5 or PPP7. Because our earlier findings with cantharidin argue against a function of PPP3 in the LH-induced decrease in NPR2 activity, PPP1, PPP2 and/or PPP6 are the phosphatases that are most likely to be important in this signaling cascade.

**LH signaling gradually reduces the follicle content of the NPR2 agonist CNP, contributing to the decrease in cGMP production that triggers resumption of meiosis**

The guanylyl cyclase assays described above were performed using membranes in the presence of a saturating concentration of CNP (1 μM), so the observed decreases in NPR2 activity occurred independently of any changes in CNP. However, previous studies in
several species have shown that on a time scale of hours after hormone injection to activate LH receptors in vivo, CNP levels decrease (see Introduction). To integrate our findings regarding the dephosphorylation of NPR2 with this previous knowledge of another component of the regulatory system, we directly compared the kinetics of the decrease of CNP levels in the same experimental system that we had used for the phosphorylation studies. CNP content, as measured using ELISA, did not change during the initial 2 h after the application of LH to follicles, but by 4 h, CNP levels had decreased to 56% of the baseline level (Fig. 7A). CNP content in the follicles did not change during parallel incubations without LH (Fig. 7B). The LH-induced CNP decrease would further decrease NPR2 activity, beginning 2-4 h after the initial dephosphorylation. However, the magnitude of this subsequent activity decrease is unknown because the ELISA measurements detect the CNP peptide, as well as its biologically inactive precursor protein.

Because the significance of the decrease in guanylyl cyclase activity is to reduce cGMP and re-initiate meiosis in the oocyte, we also examined the time course of nuclear envelope breakdown (the first morphologically detectable event in the resumption of meiosis) following the addition of LH to isolated follicles in our culture system. The time by which 50% of the oocytes had undergone nuclear envelope breakdown was ~4 h, with a marginally significant decrease observed at 2 h; almost all oocytes had resumed meiosis 12-24 h after LH exposure (Fig. 7C). This time course is similar to that previously observed using a slightly different follicle culture system or after injection of LH into rats (Tsafiriri, 1985). The effect of phosphatase inhibitors on LH-induced resumption of meiosis could not be determined because these inhibitors also act on phosphatases in the oocyte, causing meiosis to resume independently of LH (Rime and Ozon, 1990).
possibly change the conformation of NPR2 such that it is a better or worse substrate for either enzyme. Although the kinases involved have not been identified, our results indicate that the phosphatases that are important for this dephosphorylation belong to the PPP family, and not to the PPM or FCP families. Among the PPP family, our findings using cantharidin argue against an important function for PPP2 or PPP6, which are important for this dephosphorylation. Because PPP3 is activated by Ca²⁺, this conclusion is consistent with the finding that deletion of Gα₁₁ in mouse granulosa cells, and the resulting prevention of LH-induced inositol phosphate accumulation, does not inhibit meiotic progression in response to LH (Breen et al., 2013). Taken together with our gene expression analysis, PPP1, PPP2 and/or PPP6 emerge as the most likely candidates for mediating the LH-induced dephosphorylation of NPR2.

Studies of rat granulosa cells in culture have shown that LH activation of Gαq leads to the PKA-dependent activation of PPP2 and that PPP2 is associated with the MAP2D A-kinase anchoring protein (Flynn et al., 2008). One possible scenario, although not the only one, is that NPR2 could also be associated with MAP2D, such that it might be dephosphorylated by PPP2 in response to LH-Gα₁₁-PKA signaling. Because NPR2 contains multiple phosphorylation sites, and the sequences surrounding these sites differ, the regulation of other phosphatases (and kinases) might also be important in initiating or maintaining the dephosphorylation and inactivation of NPR2 in response to LH.

In particular, EGF receptor signaling is required for part of the LH-induced decrease in the cGMP content and the resumption of meiosis in mouse follicles; how much of the cGMP decrease depends on EGF receptor kinase activity is variable in different studies (Park et al., 2004; Vaccari et al., 2009; Norris et al., 2010; Hsieh et al., 2011; Liu et al., 2014). The release of EGF receptor ligands from the outer mural granulosa cell layers where the LH receptors are located provides a mechanism for paracrine signaling to cells in the interior of the follicle. In rats, the EGF receptor dependence of the LH-induced cGMP decrease has not been determined, but EGF receptor kinase activity is required over a period of hours for LH-induced resumption of meiosis (Ashkenazi et al., 2005; Reizel et al., 2010).

Signaling through various other hormones and growth factors also reduces the guanylyl cyclase activity of NPR2 in other cells, for example, vasopressin (Abbey and Potter, 2002), PDGF (Chrisman and Garbers, 1999; Abbey and Potter, 2003), lysophosphatidic acid (Abbey and Potter, 2003; Potthast et al., 2004), sphingosine-1-phosphate (Abbey-Hosch et al., 2005) and thyrotropin-releasing hormone (Thompson et al., 2014). In studies of sphingosine-1-phosphate acting on cultured fibroblasts that overexpress NPR2 (Abbey-Hosch et al., 2005), there is a correlation between dephosphorylation and the decrease in guanylyl cyclase activity. However, LH signaling in the ovarian follicle is the first example of a physiological pathway that is mediated by such a mechanism.

Other developmental processes that are regulated by the activity of NPR2 and the closely related natriuretic peptide receptor NPR1 could be controlled similarly. In particular, bone growth is affected by mutations in Npr2 or the Nppc gene that encodes CNP; increased NPR2 activity results in longer bones, whereas decreased activity results in shorter bones (Chusho et al., 2001; Tamura et al., 2004; Yasoda et al., 2004; Olney, 2006; Miura et al., 2012; Geister et al., 2013). Natriuretic peptide receptors also function in the development of the nervous system (Ter-Avetisyan et al., 2014) and heart (Becker et al., 2014). Some of the actions of growth factors and hormones that affect chondrocyte differentiation, axon bifurcation and cardiomyocyte proliferation might involve the regulation of natriuretic peptide receptor phosphorylation and/or levels of natriuretic peptides, as seen for LH-mediated regulation of meiosis in the ovary.

MATERIALS AND METHODS

Isolation and culture of rat ovarian follicles

Ovaries were obtained from 25- to 26-day-old CD-Sprague-Dawley rats (Charles River Laboratories); procedures were approved by the animal care committee of the University of Connecticut Health Center. Rats were injected with 12 U of equine chorionic gonadotropin 48 h before ovary

**Fig. 8. LH signaling in rat ovarian follicles decreases NPR2 guanylyl cyclase activity by way of a rapid dephosphorylation of regulatory sites followed by a slower decrease of the levels of the agonist CNP.** The functional domains of the homodimeric transmembrane protein NPR2 are shown in gray. Binding of CNP (green) to the extracellular domain and phosphorylation of seven juxtamembrane regulatory sites (red) both increase the catalytic activity of the enzyme. LH signaling acts by way of a PPP-family phosphatase to dephosphorylate some of these sites; dephosphorylation occurs by 10 min and persists for at least 4 h. By 4 h after LH exposure, the CNP content of the follicle decreases. Both of these changes result in decrease guanylyl cyclase activity, contributing to the decrease in cGMP that restarts meiosis.
collection to stimulate follicle growth and LH receptor expression. Preovulatory follicles, 700-900 μm in diameter, were dissected from the ovaries and cultured as previously described for mouse follicles (Norris et al., 2008) with some modifications (see supplementary Materials and Methods).

Cantharidin was obtained from Tocris Bioscience (R&D Systems) and prepared as a 50 mM stock in dimethylsulfoxide (DMSO). Okadaic acid was obtained from LC Laboratories and was prepared as a 1 mM stock in DMSO.

Preparation of crude membranes from rat follicles and measurement of guanylyl cyclase activity

Crude membranes were prepared from rat follicles, and guanylyl cyclase assays were conducted as previously described for mouse follicles (Robinson and Potter, 2011; Robinson et al., 2012). cGMP production was measured after a 9-min assay period and was approximately linear over this period (Robinson et al., 2012). Basal activity without CNP was subtracted from the activity in the presence of CNP to obtain the CNP-dependent activity. Additional details are described in the supplementary Materials and Methods.

Immunoprecipitation, Phos-tag acrylamide gel electrophoresis and western blotting

Immunoprecipitation was used to purify the low-abundance NPR2 protein from rat follicle membranes. NPR2 was immunoprecipitated by incubation with a rabbit polyclonal antisera (6328) made against the ten C-terminal amino acids of NPR2 (Abbey and Potter, 2002). Phosphorylated forms of NPR2 were separated by electrophoresis on SDS-PAGE gels made with 6% acrylamide that had been co-polymerized with 25 μM Phos-tag-acrylamide (WAKO Chemicals) and 100 μM MnCl₂. Blots were probed with the 6328 antisera. Phosphorylated forms of PDE5 were separated from follicle lysates, using Phos-tag gels as described for NPR2. The blots were probed with an affinity-purified rabbit polyclonal antibody made against a C-terminal sequence from human PDE5 (catalog no. 2395, Cell Signaling Technology). Additional details, as well as protocols for the generation of dephosphorylated NPR2 and phosphorylated PDE5 controls, are described in the supplementary Materials and Methods.

Images were analyzed using ImageJ software. As an indicator of changes in NPR2 phosphorylation, the intensity of the immunostaining was measured within boxes surrounding the upper region and the lower band (see Fig. 2C; supplementary material Fig. S1). The background intensity of each lane was collected from a box below the NPR2 signal and subtracted. The background-corrected intensity of the upper region was then divided by that of the lower band. This ratio method corrects for variability in the amount of protein that was immunoprecipitated and loaded in each lane. As a measure of total immunoreactive protein, we added the intensity of the immunostaining was measured after a 9-min assay period and was approximately linear over this period (Robinson et al., 2012). Basal activity without CNP was subtracted from the activity in the presence of CNP to obtain the CNP-dependent activity. Additional details are described in the supplementary Materials and Methods.

ELISA measurements of cGMP and CNP in follicles

The cGMP and CNP contents of rat follicles were measured as previously described (Norris et al., 2009; Robinson et al., 2012) by using ten follicles per sample and ELISA kits from Enzo Life Sciences (no. ADI-900-014 for cGMP) and Phoenix Pharmaceuticals (no. FEK-012-03 for CNP). Data were analyzed using Prism software (GraphPad).

Measurement of relative amounts of phosphatase mRNAs in granulosa cells

RNA was extracted from mural granulosa cells, mRNAs were reverse transcribed using random hexamers, and quantitative TaqMan analysis was performed as previously described (Robinson et al., 2012). Primer sequences are listed in supplementary material Table S1.

Statistics

Differences between a single treatment and control were analyzed by paired t-test using Prism software. Other data were analyzed by either repeated measures ANOVA using Prism (where sample sizes between groups were equal) or by repeated measures mixed models in IBM SPSS (v. 21.0). Post-hoc t-tests were corrected for multiple comparisons by the Holm–Bonferroni method (Holm, 1979). P-values<0.05 were considered to indicate a significant difference.

Acknowledgements

We thank Marco Conti, Jackie Corbin, Nava Dekel, John Eppig, Michael Goldberg, Mary Hunzicker-Dunn, Eiji Kinoshita, Lisa Mehlmann, Matthew Movsesian, Dieter Müller, Viacheslav Nikolaev and Rachael Norris for helpful discussions.

Competing interests

The authors declare no competing financial interests.

Author contributions

All authors performed experiments or data analysis and contributed to the development of the concepts. J.R.E., L.R.P. and L.A.J. prepared the manuscript, and all authors edited the manuscript prior to submission.

Funding

This work was supported by National Institutes of Health grants [R37HD014939, R01GM098309 and T32AR050938]; and by the Fund for Science. Deposited in PMC for release after 12 months.

Supplementary material

Supplementary material available online at http://dev.biologists.org/lookup/suppl/doi:10.1242/dev.112219/-/DC1

References


Table S1. Primers and fluorescent probes used for qRT-PCR analysis of relative expression levels of PPP family phosphatases.

<table>
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<tr>
<th>gene</th>
<th>forward primer, 5’-3’</th>
<th>probe, 5’-3’</th>
<th>reverse primer, 5’-3’</th>
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<tr>
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Probes were labeled with FAM (5’) and TAMRA (3’).
Figure S1. Validation of the specificity of the NPR2 antibody. Blot showing the lack of immunoreactivity when the immunoprecipitation and western blot of follicle membranes were done with preimmune serum instead of the 6328 antibody against a C-terminal peptide from NPR2. The follicles were treated with or without LH for 30 minutes.
Figure S2. Inhibition of the LH-induced dephosphorylation and inactivation of NPR2 by treatment of follicles with the PPP family phosphatase inhibitor okadaic acid. (A) Follicles were incubated with or without 10 µM okadaic acid for one hour, then with or without LH for 30 minutes. Crude membranes were isolated and used for immunoprecipitation, Phos-tag gel electrophoresis, and immunoblotting for NPR2. In the presence of okadaic acid, basal phosphorylation of NPR2 increased, but LH did not change the ratio of NPR2 in the upper region and lower band, indicating that okadaic acid inhibited the LH-induced dephosphorylation of NPR2. (B) Graph showing the results of 4 experiments like that shown in A (mean ± s.e.m.). (C) Membranes from follicles treated with or without okadaic acid followed by LH, as described above, were assayed for NPR2 guanylyl cyclase activity (4 experiments). Values not indicated by the same letter are significantly different. We also attempted to determine the effect of okadaic acid on the LH-induced decrease in cGMP, but these experiments were not interpretable because okadaic acid alone caused cGMP to decrease in some of the trials.
Figure S3. Blot images for figures 3A and 4A, with red boxes indicating the upper region (more phosphorylated) and lower band (less phosphorylated) for which immunostaining intensity was measured. (A) Figure 3A. (B) Figure 4A.
SUPPLEMENTARY MATERIALS AND METHODS

Isolation and culture of rat ovarian follicles

Preovulatory follicles, 700 - 900 µm in diameter, were dissected from the ovaries of rats that had been injected 48 hours previously with equine chorionic gonadotropin. Approximately 30 follicles were obtained per rat. The follicles were placed on Millicell culture inserts (PICMORG50, Millipore, Billerica, MA; 10-30 follicles per insert). MEMα medium (Invitrogen, Carlsbad, CA) was supplemented with 25 mM NaHCO₃, 3 mg/ml BSA, 5 µg/ml insulin, 5 µg/ml transferrin, 5 ng/ml selenium, 50 µg/ml streptomycin, and 75 µg/ml penicillin G. The follicles were cultured at 37°C in 5% CO₂ in air, and experimental procedures were started one to 4 hours after isolation. LH and other reagents were applied to the medium under the culture membrane, and 200-400 µl was also added to the top of the membrane to ensure rapid exposure of the follicles. Ovine LH and equine chorionic gonadotropin, purified from biological sources, were obtained from A.F. Parlow (National Hormone and Peptide Program, Torrance, CA). LH was used at a saturating concentration (10 µg/ml; approximately 350 nM). The kinetics of nuclear envelope breakdown in response to LH were determined by incubating isolated follicles with LH, and at various times afterwards, opening them with 30 gauge needles to release the cumulus-oocyte complex for observation of the presence or absence of the nucleus and nucleolus.

Preparation of crude membranes from rat follicles

To prepare crude membranes from rat follicles, 20-100 follicles were washed in PBS and then lysed in phosphatase inhibitor buffer (buffer A) containing 25 mM Hepes (pH 7.4), 50 mM NaCl, 50 mM NaF, 2 mM EDTA, 20% glycerol, 1 µM microcystin-LR (Cayman Chemical Co., Ann Arbor, MI), and protease inhibitors (complete Mini, EDTA-free; Roche Applied Science), in a 0.1 ml glass homogenizer (Wheaton, Millville, NJ) on ice. The follicle wash procedure was done at room temperature and was started 3 minutes before the homogenizer was placed on ice. LH incubation times refer to the times at which the homogenizer was chilled. The homogenate (200-400 µl volume) was centrifuged at 10,000xg for 20
minutes at 4°C; the pellet was resuspended in buffer A using a probe sonicator. Protein content was
determined by a BCA assay (Thermo Fisher Scientific, Rockford, IL). The crude membrane fraction
contained approximately 10 µg of protein per follicle. Aliquots were frozen in liquid N₂ and stored at -
80°C. Crude membranes were also prepared from HEK-293T cells stably expressing NPR2, from plates
of cells at 70-80% confluency that had been serum starved for >2 hours.

**Measurement of guanylyl cyclase activity in follicle membranes**

Guanylyl cyclase assays were conducted as previously described (Robinson and Potter, 2011). Assays
were performed at 37°C using 3-20 µg of follicle protein per assay tube, in the presence or absence of
CNP (1 µM except as indicated). The reaction mixture contained 25 mM Hepes (pH 7.4), 50 mM NaCl,
0.1% BSA, 1 mM EDTA, 0.5 µM microcystin, 5 mM MgCl₂, as well as 1 mM of the allosteric activator,
ATP, and 1 mM of the substrate, GTP. 5 mM creatine phosphate and 0.1 mg/ml creatine kinase were
included in the reaction mixture to maintain ATP and GTP concentrations. 0.5 mM
isobutylmethylxanthine was included to inhibit cGMP phosphodiesterase activity.

**Immunoprecipitation of NPR2 from rat follicle membranes**

Crude membrane samples (130-230 µg protein) were diluted to 400 µl in immunoprecipitation buffer
containing 50 mM Tris-HCl (pH 7.5), 100 mM NaCl, 50 mM NaF, 10 mM NaH₂PO₄, 2 mM EDTA, 1%
NP-40, 0.1% SDS, 0.5% sodium deoxycholate, 1 µM microcystin-LR, and protease inhibitors. NPR2 was
immunoprecipitated by incubation with 0.6 µl of 6328 rabbit polyclonal antiserum made against the 10 C-
terminal amino acids of NPR2 (Abbey and Potter, 2002) for one hour at 4°C, and then with 25 µl of
washed protein AG magnetic beads (Thermo Fisher Scientific, Rockford, IL) overnight at 4°C. The beads
were washed and resuspended in Laemmli sample buffer with 75 mM dithiothreitol. Protein was eluted
by heating at 70°C for 10 minutes. Approximately 50% of the membrane NPR2 was recovered after
immunoprecipitation.
Phos-tag acrylamide gel electrophoresis and Western blotting

Phosphorylated forms of NPR2 or PDE5 were separated by electrophoresis on SDS-PAGE gels made with 6% acrylamide copolymerized with 25 μM Phos-tag-acrylamide (WAKO Chemicals USA, Richmond, VA) and 100 μM MnCl₂ (145 x 160 x 1.5 mm gel dimensions). Gels containing 6% acrylamide, but without Phos-tag-acrylamide or MnCl₂, were used for comparison. Immunoprecipitated NPR2, or lysates of follicles that had been washed in PBS and then sonicated in Laemmli sample buffer with 75 mM dithiothreitol, were electrophoresed at 25 mA/gel for 6-8 hours at 4°C. The gels were then incubated for 20 minutes in 400 ml transfer buffer (100 mM Tris, 192 mM glycine, no SDS or methanol) containing 1 mM EDTA to chelate Mn²⁺, and then washed for 20 minutes in 400 ml transfer buffer alone to remove Mn²⁺-EDTA.

Protein was transferred to a nitrocellulose membrane for 17-20 hours with 500 mA constant current at 4°C. The membrane was stained with Ponceau-S, and blocked with 0.1% Tween and 2% milk. Blots for NPR2 were probed overnight at 4°C with a 1:50,000 dilution of the 6328 antiserum (see main text), and then with a 1:500 dilution of Clean-Blot IP Detection Reagent coupled to HRP (Thermo Fisher Scientific, Rockford, IL). Blots for PDE5 were probed overnight with a 1:500 dilution of the 2395 antibody from Cell Signaling Technology (see main text) and then with a 1:5000 dilution of a goat-anti-rabbit antibody coupled to HRP (catalog # sc-2004; Santa Cruz Biotechnology Inc., Dallas, TX). Blots were developed using ECL Prime (GE Healthcare Bio-Sciences, Piscataway, NJ), and imaged using a charge-coupled device camera (G:Box Chemi XT4, Syngene, Frederick, MD).

In vitro dephosphorylation of NPR2 in follicle membranes

To confirm that the LH-induced acceleration of NPR2 migration in a Phos-tag gel was due to dephosphorylation (Fig. 2E,F), follicle membranes were incubated at 30°C for 30 min, either with phosphatase inhibitors (50 mM NaF + 2 mM EDTA + 1 μM microcystin-LR), or under conditions that
promoted phosphatase activity (no phosphatase inhibitors, and 2 mM MgCl₂; see Bryan and Potter, 2002). NPR2 was then immunoprecipitated and separated on a gel containing Phos-tag acrylamide; NPR2 was visualized by western blotting. To prepare membranes for these assays, follicles were homogenized in buffer A without microcystin-LR, to avoid irreversible modification of phosphatases. After centrifugation, the membranes were suspended in a buffer containing 25 mM Hepes, 50 mM NaCl, 20% glycerol, and protease inhibitors. After incubation under the indicated conditions, aliquots were frozen for immunoprecipitation and western blotting (Fig. 2 E,F), and for guanylyl cyclase assays (Fig. 2G).


**In vitro phosphorylation of PDE5 in follicle lysates**

To confirm that the LH-induced retardation of PDE5 migration in a Phos-tag gel was due to phosphorylation (Fig. 5D), a lysate of follicles was incubated with the catalytic subunit of protein kinase A (PKA c, kindly provided by Jackie Corbin, Vanderbilt University), following the procedure described by Rybalkin et al., 2002. The follicles were lysed in a glass homogenizer in a buffer containing 50 mM Tris, pH 7.5, 1.5 mM EDTA, 25 mM NaF, 0.2 mM Na vanadate, and protease inhibitors (Roche complete Mini) followed by sonication. Aliquots containing 100 µg protein were then incubated at 30°C for 30 minutes with 4 µM PKA c, or with the buffer in which the PKA c was dissolved. Reactions were performed with or without 10 µM cGMP, which is required for phosphorylation of PDE5 by PKA c (Corbin et al., 2000). The samples were then spin-dialyzed into a buffer compatible with Phos-tag gel electrophoresis (50 mM Tris, pH 7.5, 25 mM NaF, 0.2 mM Na vanadate, and protease inhibitors), using a 0.5 ml, 10K Amicon Ultra centrifugal filter (EMD Millipore, Billerica, MA).