GPR126 is a member of the adhesion G protein-coupled receptors (aGPCRs) that is essential for the normal development of diverse tissues, and its mutations are implicated in various pathological processes. Here, through screening 34 steroid hormones and their derivatives for cAMP production, we found that progesterone (P4) and 17-hydroxyprogesterone (17OHP) could specifically activate GPR126 and trigger its downstream Gi signaling by binding to the ligand pocket in the seven-transmembrane domain of the C-terminal fragment of GPR126. A detailed mutagenesis screening according to a computational simulated structure model indicated that K1001ECL2 and F1012ECL2 are key residues that specifically recognize 17OHP but not progesterone. Finally, functional analysis revealed that progesterone-triggered GPR126 activation promoted cell growth in vitro and tumorigenesis in vivo, which involved Gi-SRC pathways in a triple-negative breast cancer model. Collectively, our work identified a membrane receptor for progesterone/17OHP and delineated the mechanisms by which GPR126 participated in potential tumor progression in triple-negative breast cancer, which will enrich our understanding of the functions and working mechanisms of both the aGPCR member GPR126 and the steroid hormone progesterone.

Significance

The steroid hormone progesterone is highly involved in different physiological-pathophysiological processes, including bone formation and cancer progression. Understanding the working mechanisms, especially identifying the receptors of progesterone hormones, is of great value. In the present study, we identified GPR126 as a membrane receptor for both progesterone and 17-hydroxyprogesterone and triggered its downstream G protein signaling. We further characterized the residues of GPR126 that interact with these two ligands and found that progesterone promoted the progression of a triple-negative breast cancer model through GPR126-dependent Gi-SRC signaling. Therefore, developing antagonists targeting GPR126-Gi may provide an alternative therapeutic option for patients with triple-negative breast cancer.
aGPCRs, GPR126 can be cleaved at a GPCR proteolytic site (GPS) in the GAIN domain to produce the N-terminal fragment (NTF) and C-terminal fragment (CTF) (14). In addition to the GAIN domain, the NTF of GPR126 contains a complement, Uegf, Bmp1 (CUB) domain, a pentraxin (PTX) domain, a hormone binding domain, and 27 putative N-glycosylation sites. Even after cleavage, the NTF and CTF of aGPCRs are still associated with each other in many cellular conditions (23–26). Whereas the NTF of GPR126 is known to form docking sites for extracellular matrix proteins, the CTF of GPR126 forms a 7TM bundle, and its cellular part can engage with G proteins to transduce extracellular stimuli to intracellular signaling (27–29). Recently, laminin-211, type IV collagen, and the prion protein (PrPc) were identified as extracellular endogenous ligands for GPR126 and triggered cAMP signaling to induce biological effects in Schwann cells through Gs coupling (30–32). Biochemical characterization suggested that both laminin-211 and type IV collagen bind to GPR126 through the NTF. However, whether other endogenous ligands directly interact with the CTF of GPR126 and regulate its function is unclear.

Recently, we identified hydrocortisone, a steroid hormone or glucocorticoid, as the endogenous ligand of GPR97 (also called ADGRG3, another aGPCR) (33). At the residues contacting glucocorticoids in the ligand binding pocket of GPR97 are quite conserved among the aGPCR-G subfamily, we were interested in whether certain steroid hormones similar to hydrocortisone may act as ligands of GPR126 and trigger its downstream signaling. In the present study, through screening 34 steroid hormones or their derivatives for cAMP production, we found that progestrone and 17-hydroxyprogesterone (17OHP) could specifically activate GPR126 and trigger its downstream Gi signaling. Furthermore, we identified key interaction residues and pivotal conformational changes of GPR126 in response to these two steroid hormones by computer modeling, alanine scanning, and fluorescent arsenical hairpin bioluminescence resonance energy transfer (FiaSH-BRET) methods. Finally, we explored the biological consequences of GPR126 activation in response to progestrone in triple-negative breast cancer (TNBC) cells and found that progesterone-triggered GPR126 activation promoted cell growth in vitro and tumorigenesis in vivo at least partially through Gi-tyrosine-protein kinase SRC (SRC) pathways.

Results

Screening Steroid Hormone Ligands of GPR126. Our recent studies found that GPR97 (also called ADGRG3) was able to recognize glucocorticoids, one type of steroid hormone, via its ligand pocket localized in its 7TM bundle and transduce signals via Go coupling (33). Importantly, the residues in contact with glucocorticoids in the ligand binding pocket of GPR97 are quite conserved among the aGPCR-G subfamily (33). In particular, among 13 residues of GPR97 that directly interact with cortisol, 10 residues were homologous with the other aGPCR-G subfamily member GPR126 (also called ADGRG6) (SI Appendix, Fig. S1). The glucocorticoids shared four ring steroid cores with other steroid hormones. We therefore suspected that certain steroid hormones may be able to bind to GPR126 and induce its downstream signaling.

GPR126 was reported to primarily couple to Gs in response to agonist binding (30–32). We therefore screened 34 steroid hormones and their derivatives for Gs activation using GloSensor-cAMP assays. Despite that positive control GPR126-Stachel-mimicking peptide (126-SMP) was able to stimulate approximately ninefold of intracellular cAMP increase more than control vehicle in GPR126-overexpressing HEK293 cells, no steroid hormone showed an effect by increasing intracellular cAMP levels more than twofold. Notably, 17OHP, 11-deoxycortisol, and testosterone induced weak but detectable increase on intracellular cAMP accumulation. (Fig. 1A and SI Appendix, Fig. S2 A–F) (29). The EC50 of 17OHP, 11-deoxycortisol, and testosterone are 2.7 ± 0.5 μM, 1.4 ± 0.14 μM, and 0.8 ± 0.06 μM, respectively. The Emax of these three steroid hormones and a previously reported PrPc are less than one-sixth compared with 126-SMP, and thus are weak partial agonists of GPR126 for eliciting cAMP accumulation.

In addition to Gs activity, previous reports indicated that GPR126 could stimulate Gi activity (7). We therefore stimulated HEK293 cells overexpressing GPR126 with Forskolin (Fsk) and determined whether the application of steroid hormones could dampen the cellular cAMP level by stimulating Gi activity. Notably, progesterone and 17OHP, but no other steroid hormones and the reported ligands (Stachel-mimicking peptide, collagen IV, PrPc) were able to inhibit Fsk-induced cAMP accumulation in HEK293 cells overexpressing GPR126 but not in HEK293 cells transfected with control vectors (Fig. 1A and B and SI Appendix, Fig. S2 G–M). The EC50 values for full-length GPR126 activated by progesterone or 17OHP were 539.6 ± 3.1 or 95 ± 1.8 nM, respectively (SI Appendix, Fig. S2M). In addition, these signals were blocked by PTX, a selective Gi/0 subtype inhibitor (Fig. 1 C and D and SI Appendix, Fig. S3 A and B). To investigate whether progesterone and 17OHP were able to activate GPR126 in native cells, we measured cAMP levels in GPR126 stable knocking down MDA-MB-231 cells or its control cells after administration of different concentrations of progesterone or 17OHP and Fsk (5 μM) for 10 min. The intracellular cAMP levels were detected using a cAMP enzyme linked immunosorbet assay (ELISA) kit. The results showed that Fsk induced cAMP up-regulation by approximately three times compared to cells treated with control vehicle. Both progesterone and 17OHP inhibited cAMP levels induced by Fsk in GPR126-expressing cells in a concentration-dependent manner. In the condition of GPR126 knocking down, progesterone and 17OHP showed no significant effects on cAMP levels, indicating that progesterone and 17OHP could inhibit cAMP levels via GPR126 (SI Appendix, Fig. S3 C and D).

Previous studies also suggested that several membrane-associated receptors, such as membrane bound progestrone receptors (also called progestin and adipoQ receptors, PAQRs), PGRMC1 (progestin receptor membrane component 1), and GABA-A, have been involved in the rapid, cell surface-initiated progestrone actions (34). Among these membrane receptors, GABA-A is an ion channel and PGRMC1 is a single transmembrane protein, which have not been reported to couple to G proteins (35, 36). Despite distinct trans-7TM helical topology, the membrane progestrone receptors (PAQR5/6/7/8/9) have been reported to activate G protein signaling in response to progestrone stimulation (37–41). PAQR 5/7/8 were reported to couple to Gi, whereas PAQR 6/9 coupled to Gs (42). We therefore compared the potency and efficacy of progesterone in activation of GPR126 and PAQRs. The progesterone activated PAQR7 and PAQR8 ∼100-fold stronger than GPR126, but with significant lower efficacy once these membrane receptors have similar plasma membrane expression levels (SI Appendix, Fig. S3 E and F). Considering that progesterone concentration varied across a great range, especially during pregnancy and before a baby is newly born, these data indicated that GPR126 and PAQR 5/7/8 may mediate progesterone responses in different pathophysiological contexts.
Nascent aGPCRs in general underwent autocleavage at their GPs, which produced the NTF (α-subunit) and a CTF (β-subunit), harboring the 7TM bundle (23, 24, 26, 43, 44). The Stachel sequence that lies in the N terminus of aGPCRs, such as that in GPR126, is known to mediate the self-activation of these receptors and may interfere with steroid hormone binding. We therefore generated constructs of GPR126-β and GPR126-β-ΔGPS (whose Stachel sequence was deleted) and examined their activation in response to progesterone or 17OHP stimulation. Importantly, the EC₅₀ for GPR126-β activated by progesterone or 17OHP was ∼10-fold lower than that of full-length GPR126 in cAMP inhibition assays, suggesting that the α-subunit of GPR126 may block steroid-induced GPR126 activation (SI Appendix, Fig. S4). Moreover, further deletion of the Stachel sequence by GPR126-β-ΔGPS increased the EC₅₀ of 17OHP toward cAMP inhibition by approximately twofold, suggesting a positive regulatory role of the Stachel sequence for GPR126 activated by 17OHP (SI Appendix, Fig. S4). We next examined whether GPR126 could couple to Gq or induce β-arrestin-2 recruitment upon progesterone or 17OHP administration. The results showed no obvious Gq activity or β-arrestin-2 recruitment of GPR126 in response to progesterone or 17OHP stimulation, using angiotensin II and ATIR as controls (SI Appendix, Fig. S5 G and H) (45, 46). Collectively, these results indicated that progesterone selectively induces the Gi coupling of GPR126 but is not detectable with Gs, Gq, or β-arrestin-2, whereas 17OHP induces both Gs and Gi coupling of GPR126 but not Gq or β-arrestin-2 coupling.

Conformational Changes within GPR126 Extracellular Domains as Ligand Binding. Our recent study showed that the binding of steroid hormones to the orthosteric site was able to induce selective conformational changes in the extracellular domain of GPR97 (33). We therefore investigated the extracellular conformational changes of GPR126 by the FIAsH-BRET method (33, 47). Specific positions of each extracellular loop (ECL) were screened for FIAsH motif incorporation (Fig. 2A and SI Appendix, Fig. S5A). The Nluc and FIAsH fusions used in our assays didn’t significantly alter their responses to progesterone stimulation in the cAMP assay, suggesting their functional integrity (SI Appendix, Fig. S5 B–D). Importantly, the binding of both progesterone and 17OHP induced a significant increase in the BRET signal between the Nluc-tagged N terminus and FIAsH-labeled ECL2 in a concentration-dependent manner, suggesting the movement of ECL2 close to the N terminus in response to stimulation with these two agonists (Fig. 2B and SI Appendix, Fig. S5 E–G). The FIAsH-BRET experiments also revealed different conformational changes of GPR126 in response to these two steroid hormones. Whereas progesterone induced an increase in the BRET signal between the Nluc-N terminus and FIAsH-labeled S1 site of ECL1, 17OHP showed no significant difference. Instead, 17OHP promoted an increase in the BRET signal between the FIAsH-labeled S2 site of ECL1 and the S6 site of ECL3 (Fig. 2B and SI Appendix, Fig. S5 E–G).

Structural Model of the Interaction of Progesterone and 17OHP with GPR126. The binding mode of progesterone and 17OHP in the GPR126 ligand pocket was probed by computational simulation using the Lamarckian Genetic Algorithm and Auto Dock 4 (Fig. 3 A–D and SI Appendix, Fig. S6 A and B) (33, 47–49). GPR126 was modeled according to the Swiss model using the structure of GPR97 as a template (33, 50). The steroid core of progesterone sits flat and in a direction perpendicular to TM5, and at an angle of ∼60° from the central TM3, which was similar to the binding mode of cortisol bound.
with GPR97 in our recently solved complex structure (Fig. 3 A and B). Compared to progesterone, the modeled 17OHP assumed a conformation perpendicular to the plane of the plasma membrane, rotating by ∼90° compared with that of progesterone (Fig. 3 C and D).

In the modeled ligand binding pocket of GPR126, nine hydrophobic residues and two polar residues from TM1-TM3, TM6-TM7, and ECL2 contacted progesterone in the simulated model (Fig. 3 B and G and SI Appendix, Fig. S6 C). The model of 17OHP shared nine common contacting residues of GPR126, compared with those of progesterone (Fig. 3 D and G and SI Appendix, Fig. S6 D). Consistent with these two simulated models, mutations of eight common contacting residues of both steroid ligands, including F9152, L9412, W1014ECL2, W1081ECL2, L1091ECL1, F1099ECL2, and N1103ECL2 to Ala, but not several other randomly selected surrounding residues, such as L9162A or L9373, significantly impaired GPR126 activities in response to progesterone or 17OHP stimulation in cAMP inhibition assays (Fig. 3 E and F and SI Appendix, Figs. S7 A–F and S8 A–D).

In particular, mutation of K1001ECL2 or F1012ECL2 to Ala impaired only the extracellular conformational change of GPR126 in response to 17OHP, but showed no significant effects in response to progesterone stimulation. Conversely, mutation of F10856–57 impaired only the GPR126 response stimulated by progesterone but not by 17OHP (Fig. 3 F and SI Appendix, Figs. S6 F, S7 G and K, and S8 F–I). The basal BRET value of mutations of K1001ECL2, F1012ECL2, or F10856–57 to Ala showed no significant changes (SI Appendix, Fig. S7D). These results indicated that these interactions, in particular the potential H-bond between K1001ECL2 and the 17-hydroxyl group of 17OHP, as indicated by the simulated model, are the governor of the differential recognition of steroid hormones by GPR126.

It’s worth noting that genome-wide association studies analysis has discovered dozens of SNPs in the GPR126 (ADGRG6) gene tightly associated with particular disorders or diseases, such as shortened height and AIS. Of the reported 22 SNPs in GPR126, rs17280293 (S123G), rs11155242 (K230Q), and rs2143390 (D373E) caused missense mutations (2, 51, 52). Importantly, one spontaneous mutation in GPR126 (V769E) rs2143390 (D373E) caused missense mutations (2, 51, 52). Another mutation in GPR126 (R1057Q) was reported to be associated with aggressive periodontitis (AgP) in a Japanese population (18, 53). The S123G mutation is located in the CUB domain and K230Q mutation is located in the PTX domain, while the V769E mutation is located in the GAIN domain and R1057Q mutation is located in the intracellular loop (SI Appendix, Fig. S7 M). Because only the positions of V769E and R1057Q mutations are located close to the 7TM domain, which senses the progesterone binding and couples to G proteins, we investigated the effects of these two mutations in response to progesterone stimulation in cAMP inhibition assays. Notably, the R1057Q mutant, but not the V769E, decreased the potency of progesterone (Fig. 3 H and SI Appendix, Fig. S7 N and O).

**GPR126 Expression Is Elevated in BC Tissues and Is Associated with a Poor Prognosis.** Progesterone is a key female hormone and plays essential roles in driving BC (54–57). We therefore investigated whether the activation of GPR126 by progesterone contributed to BC and how GPR126 correlates with BC progression. We examined the expression of GPR126 in 14 fresh BC tissues paired with adjacent noncancerous tissues (ANTs) by Western blot. Strikingly, 11 of 14 (78.6%) of the BC specimens had higher GPR126 protein levels than the ANT specimens (Fig. 4 A). Quantification analysis showed that the mean GPR126 level was ∼1.4-fold higher in BC tissues than that in ANTs (Fig. 4 B). Next, we performed immunohistochemistry (IHC) on another 42 BC tissue samples with paired ANT tissues to analyze GPR126 expression. Consistently, GPR126 expression was highly expressed in BC tissues (38 of 42: 90.48%, average IHC score 6.97) compared to ANT tissues (average IHC score 1.71) (P < 0.001) (Fig. 4 C and D). The aforementioned findings were further verified by analyzing the public clinical gene expression data of BC patients from The Cancer Genome Atlas (TCGA; https://portal.gdc.cancer.gov/) database (Fig. 4 E). Importantly, we found that high GPR126 expression was associated with shorter overall survival by evaluating the prognostic value of GPR126 expression using the same database (Fig. 4 F). Together, these results indicate the involvement of GPR126 in the occurrence and development of BC.

**Progesterone Promotes BC Cell Growth In Vitro and Tumorigenesis In Vivo by Activating GPR126.** To elucidate the biological roles of progesterone-induced GPR126 activation in BC development, the TNBC MDA-MB-231 cell line, which...
expresses a relatively high level of GPR126 but lacks expression of nuclear progesterone receptor (PR), was exploited (58). Two independent GPR126 short-hairpin RNAs (shRNAs) (specific for GPR126, designated Sh-GPR126-1 and Sh-GPR126-2) and one control shRNA (nonspecific sequence, designated Sh-CTRL) were stably introduced into MDA-MB-231 cells by lentivirus infection and subjected to puromycin selection. GPR126 protein levels were reduced by ∼80% in both Sh-GPR126-1 and Sh-GPR126-2 cells compared with Sh-CTRL cells (Fig. 5A and SI Appendix, Fig. S9A). Then, these cells were treated with or without progesterone (P4, 10 nM) for the indicated time (SI Appendix, Fig. S9B). The MTT [3-(4,5-dimethylthiazol-2-yl)-2, 5-diphenyltetrazolium bromide] assay showed that progesterone treatment led to an approximately twofold increase in the growth rate of Sh-CTRL cells, whereas such proliferation-promoting effects of progesterone were attenuated in GPR126-silenced cells (Fig. 5B). In accordance with this, colony-formation analysis revealed that progesterone treatment resulted in an increase in colony number and size in Sh-CTRL cells, whereas GPR126 knockdown largely abolished the increase in colony number and size in response to progesterone stimulation (Fig. 5C and SI Appendix, Fig. S9C). These results suggest that progesterone promoted the proliferation of BC cells by binding to GPR126 in vitro. At the same time, we detected the effects of the reported ligands (Stachel-mimicking peptide, collagen IV, PrPc) on the growth of MDA-MB-231 cells. Strikingly, different from progesterone, an MTT assay showed that the reported ligands had no significant effect on cell proliferation, indicating the different regulatory roles of GPR126 activation triggered by different ligands (SI Appendix, Fig. S9D).

Our biochemical and computational simulation analysis indicated that the F10856.57 is the key residue to mediate progesterone or 17OHP binding to GPR126. Detailed interactions between progesterone and GPR126 (B) as well as 17OHP and GPR126 (D) ligand binding pocket residues according to computational simulation. (E and F) Alanine mutagenesis scanning of putative residues in GPR126 ligand binding pocket on progesterone induced cAMP inhibition using GloSensor assay (E) and 17OHP (F) ligand binding pocket residues according to computational simulation. (G) The residues in GPR126 that contact with progesterone or 17OHP. Data information: (E and F) ΔpEC_{50} values of GPR126 SNPs (V769E and R1057Q) in cAMP inhibition assay were analyzed upon progesterone stimulation. Data information: (E and F) Values are the mean ± SEM of three independent experiments. **P < 0.01; ***P < 0.001; ns, no significant difference; GPR126-WT transfected cells were compared to GPR126 mutant transfected cells. All data were analyzed by one-way ANOVA with Tukey’s test. The ΔpEC_{50} values were obtained according to the data in SI Appendix, Figs. S7 B–E and H–J and S8 A–D and F–H.
overexpression of wild-type GPR126 or L937A mutant, but these promotions were significantly attenuated in cell overexpression with W1081A or F1085A mutants (Fig. 5D and SI Appendix, Fig. S9 F and G). Consistently, only exogenous overexpression of the W1081A mutant, but not L937A or F1085A mutants, inhibited 17OHP-induced cell growth in GPR126-silenced MDA-MB-231 cells (SI Appendix, Fig. S9 E, F, and H). Compared with wild-type GPR126, both K1001A and F1012A mutants alleviated cell growth promotion in response to 17OHP, but not progesterone (SI Appendix, Fig. S9 I and J–M). We also detected the effects of GPR126 SNPs on cell growth induced by progesterone. MTT assay showed that exogenous overexpression of R1057Q, but not V769E, inhibited progesterone-induced cell growth in GPR126-silenced MDA-MB-231 cells (SI Appendix, Fig. S10). Collectively, these data suggest that recognition of progesterone and 17OHP by specific residues in GPR126, or natural GPR126 missense mutation, contributed to its function in promotion of cell growth and colony formation.

To extend our in vitro observations to a more pathological context, we investigated whether progesterone-induced GPR126 activation regulated the tumorigenic capacity of MDA-MB-231 cells using an athymic nude mouse xenograft model. MDA-MB-231-Sh-GPR126-1 and Sh-CTRL cells were subcutaneously inoculated into the axillae of 12 BALB/c-nude mice, 6 for each group injected intraperitoneally with either physiological saline or progesterone (15 mg/kg, every 3 d; 32 d). The mouse weight and subcutaneous tumor size were measured every 2 d. To determine the plasma concentration of progesterone delivered through intraperitoneal injection, we measured plasma progesterone levels after administration. The results showed that the plasma concentration of progesterone initially increased rapidly, reached the maximum (151 nM) ∼1 h after injection, and declined thereafter until 8 h to maintain a steady level of the reported physiological levels during pregnancy from below 6.36 nM (2 ng/mL) to 477 nM (150 ng/mL) (59).

Notably, although the rates of mouse weight gain were not significantly different between each group (SI Appendix, Fig. S7O), the tumors originating from MDA-MB-231-Sh-GPR126-1 cells grew more slowly than those originating from...
MDA-MB-231-Sh-CTRL cells at the implantation sites (Fig. 5E). Moreover, progesterone treatment significantly enhanced tumor growth in the Sh-CTRL group but had minimal effects on tumor growth in the Sh-GPR126-1 group (Fig. 5E). After 32 d, the mice were killed. We found that the volume and weight of xenografted tumors originating from MDA-MB-231-Sh-GPR126-1 cells were much smaller and lighter than those originating from MDA-MB-231-Sh-CTRL cells (Fig. 5F and SI Appendix, Fig. S7F). In addition, progesterone treatment significantly increased the size and weight of tumors in the Sh-CTRL group but had no significant effects on those in the Sh-GPR126-1 group, suggesting that progesterone promotes tumorgenesis through GPR126 in vivo (Fig. 5E and F). Taken together, our results demonstrated that GPR126 activated by progesterone stimulated the proliferation of BC cells in vitro and tumor formation in vivo.

**Gi-Dependent SRC Activation Contributes to Progesterone/GPR126-Mediated Cellular Function.** Increasing evidence has highlighted the importance of SRC in tumorigenesis and in Gi-coupled receptor signaling (60–66). We therefore sought to determine the involvement of SRC signaling in progesterone-stimulated cell proliferation through GPR126-Gi activation in BC cells. Phosphorylation of SRC at Y416 (p-SrcY416) acts as the known active form to trigger the cascade phosphorylation of a series of downstream effectors, including the phosphorylation of AKT at S473 (p-AKTY473), and phosphorylation of ERK at T202 and Y204 (p-ERKY202/Y204) (67, 68). Then the phosphorylation status of these proteins was detected in MDA-MB-231 cells treated with progesterone for different times. Strikingly, in MDA-MB-231 cells, progesterone promoted the phosphorylation of SRC at Y416 as well as p-AKTY473 and p-ERKY202/Y204 in a time-dependent manner and reached a maximum 5 min after progesterone treatment (SI Appendix, Fig. S10 A and B). Importantly, either GPR126 silencing or Gi inhibition abrogated increase of p-SRCY416, p-AKTY473, and p-ERKY202/Y204 induced by progesterone, indicating the essential role of the GPR126-Gi axis in SRC signaling activation in response to progesterone stimulation (SI Appendix, Fig. S11 C and E and SI Appendix, Fig. S11 D and F).

We next investigated GPR126 ligand pocket mutations on the p-SRCY416 level in response to progesterone or 17OHP stimulation. Progesterone treatment was able to promote the pY416 SRC phosphorylation in GPR126 wild-type or L937A mutant-overexpressed cells with GPR126 silencing, but failed to increase the p-SRCY416 level in GPR126-silenced MDA-MB-231 cells with ectopic overexpression of F1085A or W1018A mutants (SI Appendix, Fig. S12 A and B). Specifically, 17OHP treatment was able to promote the pY416 SRC phosphorylation in GPR126 wild-type, L937A, or F1085A mutants overexpressed cells with GPR126 silencing, but failed to increase the p-SRCY416 level in GPR126-silenced MDA-MB-231 cells with ectopic overexpression of W1018A mutant (SI Appendix, Fig. S12 C and D). These results support the correlation between the specific progesterone/17OHP recognition and the downstream SRC activation. Further MTT analysis revealed that both the SRC inhibitor (Src-I1, 2 μM) and Gi inhibitor (PTX, 100 ng/mL) significantly suppressed progesterone-induced cell growth (Fig. 6A). Collectively, these data suggested that progesterone binding to GPR126 promoted Gi coupling and downstream SRC activation, thus promoting cancer cell growth (Fig. 6B).

**Discussion**

GPR126 has been reported to play critical roles in the normal development of diverse tissues and is implicated in various pathological processes, such as scoliosis and carcinogenesis, through involvement in many different signaling networks (1–14). The identification of specific ligands for GPR126 in distinct physiological or pathophysiological processes is of great importance for elucidating its precise functions and corresponding mechanisms. Previous studies have identified at least three proteins or protein fragment—laminin-211, type IV collagen, and PrPc—as ligands for GPR126 (30–32). Laminin-211 and type IV collagen are major components of the extracellular matrix of Schwann cells and axonal neurons. Laminin-211 binds to a specific domain of GPR126, ranging across residue D446 to residue C807, whereas type IV collagen interacts with the CUB and PTX domains in GPR126-NTF. Similar to the GPRGKPG motif of type IV collagen, PrPc has a conserved sequence (KKRPKPG) in the amino-terminal flexible tail, which is necessary for the activation of GPR126 (30–32). All three protein ligands contribute to Schwann cell development and peripheral nervous system myelination by activating GPR126 signaling. Although it is known that both type IV collagen and laminin-211 require the extracellular domain of GPR126 for their efficient interactions, whether these three protein ligands of GPR126 directly interact with the 7TM domain of the CTF remains elusive.

In the present study, we found that progesterone and 17OHP are ligands of GPR126, both in vitro and particularly in a tumor progression model (Fig. 6B). Progesterone and 17OHP are canonical steroid hormones that play important roles in the development and functional maintenance of the female reproductive system. Our results suggest that progesterone and 17OHP bind to the ligand pocket in the 7TM domain of the CTF rather than the NTF of GPR126. Moreover, a detailed mutagenesis screening according to a computational simulated structure model indicates that K1001ECL2 and F1012 ECL2 are key residues that specifically recognize 17OHP but not progesterone, probably by interaction with the hydroxyl group. Recently, we identified that glucocorticoids, a group of steroid hormones, including hydrocortisone and cortisone, are able to activate GPR97 (33). Both GPR126 and GPR97 belong to the aGPCR subfamily (69, 70). Our work on GPR126, together with this previous study showing that GPR126 and GPR97 share the conserved ligand binding pocket, raises the possibility that multiple steroid hormones may function as ligands for different aGPCRs, especially members of the ADGRG subfamily.

Our data show that progesterone activated GPR126 with an EC₅₀ of ~539.6 nM, which is significantly different from the reported EC₅₀ of progesterone to activate its nuclear receptor (~1 nM), suggesting these two receptors may respond to progesterone in different physiological contexts. It is known that concentrations of progesterone vary across a wide range during the menstrual period. In human, the concentration of progesterone in the premenstrual and follicular phase keeps below 0.636 nM (0.2 ng/mL), and then goes up to 63.6 nM (20 ng/mL) in the luteal phase (71). Notably, the concentrations of progesterone are significantly elevated during pregnancy. The highest concentration of progesterone could reach to ~1.09 nM (342 ng/mL, covering its EC₅₀ to activate GPR126) before a baby is newly born (71). Moreover, GPR126 expression profile analysis indicated the enrichment of Gpr126 in mouse uterus (SI Appendix, Fig. S12E) and mammary gland (72). These data indicate that progesterone-
GPR126 signaling may play important roles in modulation of pregnancy and mammary gland development, which deserve future investigation.

Compared with progesterone, 17OHP has been much less studied. Notably, patients with classic congenital adrenal hyperplasia had markedly higher plasma 17OHP concentration (mean concentration of 280 nM) compared to normal people (mean concentration of 4.7 nM) (73). Moreover, recent reports have indicated that the hepatic levels of 17OHP were significantly increased in diabetic mice with a maximum concentration up to 1.8 μM and the accumulated 17OHP contributed to the pathogenesis of hyperglycemia and insulin resistance through activating glucocorticoid receptor (EC_{50} = 7.14 μM) (74). Given that our data show the EC_{50} values of 17OHP to activate full-length GPR126 and its β-subunit GPR126-β were 94 ± 0.63 nM and 10.0 ± 1.23 nM, respectively, the indicated concentrations of 17OHP in congenital adrenal hyperplasia patients and diabetic mice are sufficient to activate full-length GPR126, implying the possible involvement of 17OHP-mediated GPR126 activation in such pathological processes. Recent studies have also indicated the β-subunit of aGPCRs, such as GPR133, functions independently (75). Therefore, it doesn’t exclude the possibility that the 17OHP may regulate function of GPR126 in normal physiological processes.

Although all previously reported agonists activate Gs signaling downstream of GPR126 and subsequently stimulate the production of cAMP (30–32), our studies demonstrate that GPR126 was able to stimulate Gi activity in response to progesterone or 17OHP stimulation. These results suggest that different ligands for the same aGPCR member, for example...
GPR126, may have distinct biased signaling properties that constitute the fine-tuning signaling network for these receptors with diverse functions. Recently, different biased properties of the G protein subtype or arrestins of GPCR ligands have been suggested to have great therapeutic potential, such as that for angiotensin type I receptors, opioid receptors, and adrenergic receptors (66, 76–90). Therefore, understanding the preferences of GPR126 may facilitate future therapeutic development targeting this particular receptor.

Given the critical roles of progesterone in regulating BC (54–56), we determined the involvement of GPR126 in BC progression and the functional consequence of progesterone-induced GPR126 activation. By examining the GPR126 expression levels in clinical samples and analyzing the public clinical gene-expression data of BC patients from the TCGA database, we found that GPR126 was highly expressed in the tumor tissue of BC patients. In addition, prognostic analysis revealed that high GPR126 expression correlates with unfavorable overall survival in BC patients, implicating GPR126 in the occurrence and progression of BC. Functionally, progesterone-triggered GPR126 activation significantly promoted the cell growth and tumorigenesis of TNBC cells both in vitro and in vivo. A preliminary mechanistic investigation demonstrated that Gi-dependent SRC activation downstream of the progesterone-GPR126 interaction contributes to these effects. TNBC is more aggressive and has a poorer prognosis than other types of BC because of the lack of expression of nuclear receptors for estrogen and progesterone, thus leading to refractoriness to hormone therapy (91). Our study showed that despite the absence of progesterone nuclear receptor, progesterone is still able to promote the development of BC by activating the membrane receptor GPR126 and its downstream Gi signaling. Therefore, developing antagonists targeting GPR126-Gi may provide an alternative therapeutic option for patients with TNBC, which requires further investigation.

GPR126 is expressed in a wide range of tissues and functions in a variety of physiological and pathological processes, such as the development of bone, ear, and heart, myelination of the peripheral nervous system, and neuromuscular diseases (2, 5, 7, 9, 51, 92, 93). In addition to the contributions of progesterone-GPR126 signaling to BC, further elucidation is needed to examine whether these widely distributed steroid hormones regulate other pathophysiological processes through GPR126 and whether the mechanism of GPR126 signaling activation by progesterone deciphered here has broader applications.

In summary, our present study found that the steroid hormones progesterone and 17OHP can activate the Gi signaling of GPR126 by binding to its 7TM domain, and the interaction between progesterone and GPR126 has important roles in a tumor progression model (Fig. 6B). The identification of a membrane receptor for progesterone other than classic nuclear receptors and the definition of the mechanisms of GPR126 involvement in potential tumor progression in TNBC will enrich our understanding of the functions and working mechanisms of both the αGPCR member GPR126 and progesterone.

Materials and Methods

Cell Lines. HEK293 cells were obtained from the Cell Resource Center of Shanghai Institute for Biological Sciences (Chinese Academy of Science) and grown in Dulbecco’s Modified Eagle Medium (DMEM) culture with 10% FBS (Gibco). MDA-MB-231 cells were originally obtained from the American Type Culture Collection (ATCC) and grown in monolayer culture in Roswell Park Memorial Institute (RPMI) 1640 with 10% fetal bovine serum (FBS) (Gibco) and 1% penicillin/streptomycin. Both kinds of cells were maintained at 37 °C in a humidified atmosphere consisting of 5% CO2 and 95% air.

Chemicals, Reagents, and Peptides. Cortisol, corticosterone, calcifiediol, calcitriol, 11-deoxycorticisol, deoxycorticosterone, DHEA (dihydroepiandrosterone), estradiol, 2-methoxyestrone, pregnenolone, progesterone, vitamin D3, 16a-hydroxyprogesterone, 11β-hydroxyprogesterone, vitamin D2, prednisone, betamethasone were purchased from Med Chem Express. Androstane, 20α-hydroxycholesterol, dehydrotestosterone, 11α-hydroxyprogesterone, vitamin D2, prednisone, betamethasone were purchased from Med Chem Express. Androstane, 20α-hydroxycholesterol, dehydrotestosterone, 11α-hydroxyprogesterone, vitamin D2, prednisone, betamethasone were purchased from Med Chem Express.
Cell Signaling Technology. ERK2 antibody (sc-1647) was from Santa Cruz Biotechnology.

The high purity (above 95%) 126-SMP [VHF(4-Me)GVLMDLPRSASQI] were chemically synthesized at Chinapeptides (Shanghai, China, http://www.chinapeptides.com). All the steroid chemicals or peptides were dissolved in dimethyl sulfoxide (DMSO) to make stock solutions at final concentrations of 100 mM, which was further diluted to working concentrations with phosphate buffer solution (PBS).

Constructs. The human GPR126 gene (ADGRG6) was cloned into the pcDNA3.1 vector with an N-terminal Flag epitope tag (DYKDDDDK). The CTF of GPR126 (GPR126-β) and GPR126-β-ΔGPS (whose Stachel sequence was deleted) were subcloned into pcDNA3.1 vector with an N-terminal Flag tag. The GPR126 mutations S868A, F915A, L937A, L951A, Y999A, K1001A, F1012A, W1014A, F1085A, L1091A, F1099A, N1103A, S123G, K230Q, D373E, V769E, and R1057Q were generated using the QuikChange mutagenesis kit (Stratagene). The human PAQR5, PAQR6, PAQR7, PAQR8, and PAQR9 genes were cloned into the pcDNA3.1 vector with an N-terminal Flag epitope tag (DYKDDDDK). For detection of conformational changes of extracellular region using FlAsH-BRET assay, NanoLuc (Nluc) and a four-amino acid linker (Gly-Ser-Ser-Gly) were fused to the N-terminal of GPR126. FlAsH sequence (Cys-Pro-Gly-Cys-Cys) was inserted into the ECL1, ECL2 or ECL3 region of Nluc-GPR126, as indicated in SI Appendix, Fig. S54. All primers used above are listed in SI Appendix, Tables S1 and S2. The GloSensor plasmid was obtained from Promega. All constructs were generated using Quikchange site-directed mutagenesis kit (Stratagene) and verified by DNA sequencing.

Measurement of Receptor Expression at Cell Surface by ELISA. To evaluate the expression level of wild-type GPR126 and its mutants, HEK293 cells were transfected with wild-type and mutant GPR126 or vehicle (pcDNA3.1) using PEI reagent in six-well plates, according to the manufacturer’s instructions (33, 47). After incubation at 37 °C for 16 h, cells were then seeded into 24-well plates at a density of 10^5 cells per well for another 16-h incubation at the same temperature. Next, cells were fixed with 4% (wt/vol) paraformaldehyde for 15 min and blocked with 5% (wt/vol) BSA for 1 h at room temperature. Cells were then incubated with the monoclonal anti-FLAG primary antibody (Cat #F1804, Sigma Aldrich) overnight at 4 °C and followed by incubation of a secondary goat anti-mouse antibody (Cat #A-21235, Thermo Fisher) conjugated to horseradish peroxide for 1 h at room temperature. After washing, TMB (3, 3,5 ,5 -tetramethylbenzidine) solution was added into each well for color reaction. An absorbance value was measured at 450 nm using the Tecan (Infinite M200 Pro NANO Quant) luminescence counter for calculating and analyzing the relative expression levels of corresponding plasmids.

cAMP Accumulation and Inhibition Assay. To detect the cAMP accumulation of GPR126 and the inhibitory effects on Fsk-induced cAMP accumulation of different GPR126 constructs or mutants in response to different ligands, GloSensor cAMP assay (Promega) was performed according to previous publications (33). HEK293 cells were transfected with wild-type or mutant GPR126 or vehicle (pcDNA3.1) and GloSensor plasmid using PEI reagent in six-well plates according to the manufacturer’s instructions. After incubation at 37 °C for 24 h, cells were then seeded into 96-well plates at a density of 4 × 10^4 cells per well for another 24-h incubation at the same temperature. The cells were preincubated with serum-free medium (Gibco) containing 5% (vol/vol) dilution of the GloSensor cAMP reagent stock solution (Promega) for 2 h. After that, Fsk (5 μM, using in cAMP inhibition assay) and different concentrations of ligands were added into each well. Then, the luminescence intensity was examined immediately using an EnVision multilabel microplate detector (Perkin-Elmer). Data were analyzed using the sigmoidal dose-response function in GraphPad Prism 7.0.

FlAsH-BRET Assay. To monitor the extracellular conformational changes of GPR126 deduced by progesterone and 17OHP binding, a FlAsH-BRET assay was performed as previously described (33, 45, 54). In brief, HEK293 cells were seeded in six-well plate and transfected with GPR126 FlAsH-BRET sensors (SI to S6, as indicated in SI Appendix, Fig. S54) or relative mutants based on sensor S4 and incubated for 48 h at 37 °C in 5% CO2. Then the cells were labeled with 2.5 μM FlAsH EDT2 (Thermo Fisher Scientific) in accordance with the manufacturer’s specification. After that, the cells expressing the labeled GPR126 FlAsH-BRET sensors were seeded into a 96-well plate at a density of 5 × 10^3 cells per well. The indicated concentrations of progesterone and 17OHP were added into each well together with luciferase substrate coelenterazine-h (5 μM). The BRET signal was calculated as the ratio of FlAsH to Nluc emission. The change in BRET signal due to ligand addition was recorded as ΔBRET.

Molecular Dynamics Simulation of Progesterone-GPR126 Complex and 17OHP-GPR126 Complex. The initial binding pose of progesterone or 17OHP with GPR126 was determined by Auto Dock4 (48). The size of docking box was set to 60 × 60 × 60 Å, the number of GA runs was set to 200 with the 0.375 Å grid spacing, and the center of mass of progesterone or 17OHP was set to the docking center. We analyzed the molecular docking results of progesterone or 17OHP with GPR126, the conformation with the lowest binding energy was selected for the next molecular dynamics (MD) simulation. All the MD simulation input files of progesterone or 17OHP with GPR126 complex were generated by the CHARMM-GUI website (95). We embedded the progesterone-GPR126 complex or 17OHP-GPR126 complex into the double-layer POPC membrane, adding with 0.15 M NaCl to balance the system charge, and incorporation the ions into the system using the Monte Carlo method. The TIP3P model was used for hexagonal type water box. The steepest descent method was used to minimize the progesterone or 17OHP with GPR126 complex simulation system with a series of restraints for the GPR126, 17OHP, or progesterone and lipid atoms to ensure that the energy of the entire system converges to 500 kJ/(mol·nm). The simulation system was heated from 0 K to 310 K in the NVT ensemble (number of particles, volume and temperature are conserved) for 1,000 ps. Following this, the NPT ensemble simulations were run at 1 atm for 1,000 ps with 10.0 kcal mol⁻¹ Å⁻² harmonic restraints. The 200-ns MD simulation was carried out by Gromacs2019.5 (96) with a Charmm36m all-atom force field (97). In the MD simulation, the LINCS algorithm was used to constrain all the bonds involving hydrogen. The cutoff value for nonbonding interactions was 12 Å. A time step of 2 fs was used and the trajectories were saved every 10 ps.

IHC and Scoring. Samples of breast tumor and adjacent tissues were obtained from 14 patients undergoing surgical excision of tumors at Shandong Provincial Hospital (Jinan, China). The use of pathological specimens and the review of all pertinent patient records were approved by the Shandong Provincial Hospital Ethical Review Board. Informed consent was obtained from the patients. Frozen tissues were subjected to Western blot analysis. Breast tissue microarray was purchased from Wuhan Servicebio Technology, which contains 42 cases of breast adenocarcinoma with paired paraneoplastic tissues. The clinic pathological features of the following samples were available on the company’s website. IHC was performed following standard protocol which was described in detail in SI Appendix files.

Establishment of GPR126 Stable Knockdown Cell Lines. Two shRNA oligos targeting human GPR126 including 5’-GCTCACAGTAGTAGCTAACTCTAATGAATAT-3’ (Sh1) and 5’-GCTCAACTCTAATGAATAT-3’ (Sh2) were synthesized and fused into pLKO.1 vector (Addgene) as instructions and a nonspecific sequence (5’-TTTCCGAACGGTT-CAGT-3’) was used as negative control. Lentivirus expressing GPR126 shRNAs or control shRNA was prepared as described previously (98). MDA-MB-231 cells were infected with the indicated lentiviral supernatant along with polybrene (4 μg/ml; Millipore/Chemicon), followed by selection with puromycin (2 μg/ml; Solarbio). Finally, the monoclonal cell lines stably expressing GPR126 shRNAs or control shRNAs were obtained. The expression levels of GPR126 in GPR126 knockdown cells (designated as sh-GPR126-1, and sh-GPR126-2) and their controls (designated as sh-CTRL) were confirmed by Western blot analysis.

Western Blot Analysis. For protein phosphorylation detection, MDA-MB-231 cells were starved in serum free medium for 8 h and then treated with progesterone (10 nM) for 5 min. Then cells were harvested and lysed using lysis buffer (50 mM Tris, pH 8.0; 150 mM NaCl; 1 mM NaF; 1% Nonidet P-40; 2 mM EDTA; 10% glycerol; 0.25% sodium deoxycholate; 1 mM Na3VO4; 0.3 μM aprotinin; 130 μM Bectasin; 1 μM leupeptin; 1 μM pepstatin; 5 mM iodoacetate; 10 mM pyrophosphate; 1 mM PMSF) on ice for 30 min. Then, 30 μg whole-cell protein lysates were separated by sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS/PAGE) and transferred to polyvinylidene fluoride (PVDF) membrane by electrobobbling. The membrane was immunooblotted with the indicated manufacturer.
primary and secondary antibodies. Protein bands from the Western blot were quantified using ImageJ software (NIH). For GPR126 expression detection, breast tumor tissues and adjacent nontumorous tissues were grinded in the cold on ice for 1 h. Then the samples were further analyzed using Western blot analysis.

cAMP Level Detection by ELISA Kit. cAMP levels in MDA-MB-231 cells were detected using the cAMP Parameter Assay Kit (R&D Systems) according to the manufacturer’s instruction. Detailed protocol was described in SI Appendix files.

Cell Proliferation Assay. Cell proliferation were detected using MITI assay which was described in detail in SI Appendix files.

Colony Formation Assays. Colony formation assays were held followed by protocol described in SI Appendix files.

Nude Mice Xenograft Model. Nude mice xenograft experiments were carried on followed by protocol described in SI Appendix files.

Statistical Analysis. All data are representative of at least three independent experiments. All statistical analyses were performed using GraphPad Prism 7.0. The data are presented as the mean ± SEM. Student’s t-test or one-way ANOVA with Tukey’s test was used for comparisons among groups. Survival data were analyzed using Kaplan-Meier statistical method. P < 0.05 was considered statistically significant. In the graphed data, *P < 0.05, **P < 0.01, and ***P < 0.001, and #P < 0.05, ##P < 0.01, and ###P < 0.001.

Data Availability. All study data are included in the main text and SI Appendix.

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PNAS 2022 Vol. 119 No. 15 e2117004119 https://doi.org/10.1073/pnas.2117004119 11 of 12
