

R-spondins are BMP receptor antagonists in early embryonic development

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ABSTRACT

BMP signalling plays key roles in development, stem cells, adult tissue homeostasis, and disease. How BMP receptors are extracellularly modulated and in which physiological context, is therefore of prime importance. R-spondins (RSPOs) are a small family of secreted proteins that co-activate WNT signalling and function as potent stem cell effectors and oncogenes. Evidence is mounting that RSPOs act WNT-independently but how and in which physiological processes remains enigmatic. Here we show that RSPO2 and RSPO3 also act as BMP antagonists. RSPO2 is a high affinity ligand for the type I BMP receptor BMPRI1A/ALK3, and it engages ZNRF3 to trigger internalization and degradation of BMPRI1A. In early *Xenopus* embryos, Rspo2 is a negative feedback inhibitor in the BMP4 synexpression group and regulates dorsoventral axis formation. We conclude that R-Spondins are bifunctional ligands, which activate WNT- and inhibit BMP signalling via ZNRF3, with implications for development and cancer.

1 INTRODUCTION

2 Bone Morphogenetic Proteins (BMPs) are a subfamily of TGF β growth factors that exert a
3 plethora of crucial functions in embryonic development, adult tissue homeostasis as well as
4 regeneration, and they underlie human pathology such as skeletal disorders, cancer, and fibrosis
5 in multiple organs¹⁻⁵. Due to their accessibility, extracellular components of the BMP pathway
6 are of particular interest as therapeutic targets⁶ and mechanistic understanding of receptor
7 modulation should improve the ability to manipulate BMP-dependent processes.

8 BMPs signal through a tetrameric receptor kinase complex composed of type I (BMPRI1A/
9 ALK3, BMPRI1B/ALK6, ACVR1/ALK2, or ACVRL1/ALK1) and type II receptors (BMPRI2,
10 ACVR2A, ACVR2B)⁷. Ligands and receptors combine in a combinatorial fashion⁸ and
11 phosphorylate SMAD1, 5, and 8, which enter the nucleus with SMAD4 to regulate target gene
12 expression^{9, 10}. There exists a multitude of extracellular modulators of TGF β signaling, either
13 soluble or membrane-associated proteins that control ligand availability, processing, ligand–
14 receptor interaction, and receptor activation¹¹. However, only two BMP receptor antagonists are
15 known, which directly bind and inhibit receptor function, the TGF β -family proteins BMP3 and
16 Inhibin^{12, 13}.

17 R-Spondins (RSPO1-4) are a family of four secreted ~30kDa proteins implicated in development
18 and cancer¹⁴⁻²⁰. RSPOs are a key ingredient to maintain organoid cultures where they stimulate
19 stem cell growth^{21, 22}. They amplify WNT signaling by preventing Frizzled/LRP5/6 receptor
20 ubiquitination and degradation via transmembrane E3 ubiquitin ligases ring finger 43 (RNF43)
21 and zinc and ring finger 3 (ZNRK3), thereby sensitizing cells to WNT ligands^{14, 23-25}. RSPOs
22 bind to ZNRK3/RNF43 and to the stem cell marker Leucine-rich repeat containing G protein-

23 coupled receptor 5 (LGR5), and two related proteins, LGR4 and LGR6, leading to the
24 internalization of the RSPO-LGR-ZNRF3/RNF43 complex and lysosomal degradation^{14, 17, 26}.
25 RSPOs harbor two furin-like repeats (FU1, FU2) domains that bind to ZNRF3/RNF43 and
26 LGRs, respectively²⁷. In addition, they contain a thrombospondin 1 (TSP1) domain, which
27 possess about 40% overall sequence homology^{24, 28}. The TSP1 domain is not essential for
28 WNT/LRP6 signaling but it binds to HSPGs (Heparan Sulfate Proteoglycans) and thereby
29 promotes WNT5A/PCP (planar cell polarity) signaling^{24, 29}.

30 Unexpectedly, recent studies showed that RSPO2 and RSPO3 can potentiate WNT signaling in
31 the absence of all three LGRs *in vitro* and *in vivo*^{27, 30}. Moreover, WNT and RSPO ligands are
32 functionally non-equivalent since e.g. WNT ligand overexpression cannot induce crypt
33 expansion in contrast to RSPO2 or RSPO3³¹ and RSPO2 and WNT1 have distinct effects on
34 mammary epithelial cell growth³² and cochlea development³³. Hence, these inconsistencies in
35 our current understanding raise the questions: Do RSPOs possess WNT-independent functions?
36 Do they engage other receptors? If so, in which physiological processes is this relevant?

37 Here we show that RSPO2 and RSPO3 are high affinity ligands for the BMP receptor
38 BMPR1A/ALK3. RSPO2 forms a ternary complex between BMPR1A and the E3 ligase ZNRF3,
39 which triggers endocytosis and degradation of the BMP receptor. We show that Rspo2
40 antagonizes BMP signaling during embryonic axis formation in *Xenopus*. By gain- and loss-of-
41 function experiments *rspo2* cooperates with Spemann organizer effectors to regulate the BMP
42 morphogen gradient, which controls dorsoventral axis formation. Our study reveals R-spondins
43 as a novel class of BMP receptor antagonists in development, inviting re-interpretation of the
44 mode of action of R-Spondins and ZNRF3 in stem cell- and cancer biology.

45 RESULTS

46 RSPO2 and -3 antagonize BMP4 signaling independently of WNT

47 In considering possible WNT-independent functions of RSPOs, we revisited our early
48 observation that *rspo2* overexpression affected BMP signaling in *Xenopus* embryos²⁰. We tested
49 if RSPO2 could suppress BMP signaling in human cells. To this end, we utilized human
50 hepatocellular carcinoma (HEPG2) cells, which express very low levels of RSPOs
51 ([Supplementary Fig. 1a](#)). Intriguingly, treatment with RSPO2 and RSPO3 but not RSPO1 and
52 RSPO4 decreased BMP4 signaling, while all RSPOs showed similar ability to amplify WNT
53 signaling ([Fig. 1a](#), [Supplementary Fig. 1b](#)). Importantly, inhibition of BMP signaling by RSPO2
54 and -3 was independent of WNT/ β -catenin signaling, since it remained unaffected by siRNA
55 knockdown of *β -catenin* ([Fig. 1b](#), [Supplementary Fig. 1c-d](#)). RSPO2 and -3, but not RSPO1 and
56 RSPO4 treatment decreased phosphorylation of Smad1, which is a hallmark of BMP signaling
57 activation ([Fig. 1c-d](#), [Supplementary Fig. 1e-f](#)). Focusing on RSPO2, we confirmed that RSPO2
58 overexpression decreased Smad1 phosphorylation and treatment with RSPO2 protein decreased
59 BMP target *ID1* expression ([Supplementary Fig. 1g](#), [Fig. 1e](#)). Inhibition of BMP signaling by
60 RSPO2 was unaffected by siRNA knockdown of *LGR4/5*, *LRP5/6*, *DVLI1/2/3* and *ROR1/2* ([Fig.](#)
61 [1f-g](#), [Supplementary Fig. 1h-j](#)), suggesting independence of WNT/LRP and WNT/PCP signaling.
62 Moreover, different from RSPO2, treatment with WNT3A, WNT3A surrogate³⁴, or the WNT
63 antagonist DKK1 had no effect on BMP signaling ([Fig. 1h](#), [Supplementary Fig. 1k-l](#)),
64 corroborating WNT-independent RSPO2 function.

65 To delineate the domains required for BMP inhibition, we analyzed deletion mutants of RSPO2
66 and found both the TSP1- and FU-domains to be important for signaling inhibition ([Fig. 1i-j](#))²⁴.

67 We next investigated RSPO2 deficiency in H1581 cells, a human large cell lung carcinoma cell
68 line that expresses high levels of *RSPO2* (Supplementary Fig. 1a). Knockdown of *RSPO2* but not
69 *LRP5/6* sensitized H1581 cells to BMP stimulation (Fig. 1k-l, Supplementary Fig. 1m-n). We
70 conclude that RSPO2 and -3 antagonize BMP signaling independently of WNT signaling.

71

72 **Rspo2 antagonizes BMP signaling during *Xenopus* embryonic axis development**

73 To analyze if Rspo2 inhibits BMP signaling *in vivo*, we turned to early *Xenopus* development. In
74 the early amphibian embryo, the Spemann organizer is a small evolutionary conserved signaling
75 center, which plays an eminent role in regulating embryonic axis formation and neural induction.
76 One essential molecular mechanism underlying Spemann organizer function resides in its
77 secretion of BMP antagonists, which create a BMP morphogen gradient that patterns the
78 embryo³⁵⁻³⁷. Since *rspo2* is expressed and functions in WNT-mediated myogenesis of early
79 *Xenopus* embryos²⁰, we analyzed if it may have an additional role as BMP antagonist in axial
80 patterning.

81 *bmp4* overexpression ventralizes *Xenopus* embryos, resulting in small heads and enlarged ventral
82 structures³⁸. Injection of wild-type *rspo2* mRNA, but neither its Δ FU1/2 nor Δ TSP1 deletion
83 mutants rescued these *bmp4*-induced malformations (Fig. 2a-b). This domain requirement is
84 different from that for WNT signaling activation, where only FU1 and FU2 but not the TSP1
85 domain are essential²⁰. Conversely, injection of a previously characterized *rspo2* antisense
86 Morpholino (Mo)²⁰ increased endogenous BMP signaling, and this was unaffected by *lrp6* Mo
87 (Fig. 2c)³⁹. Strikingly, coinjection of *bmp4* Mo and *rspo2* Mo neutralized each other in BMP

88 signaling reporter assay (Fig. 2d), BMP target gene expression (*vent1*, *sizzled*) (Fig. 2e-f), as well
89 as defects in dorsoventral axis development (Supplementary Fig. 2a-b). Typically,
90 overexpression of common BMP antagonists such as *noggin* or *chordin* that sequester BMP
91 ligands, leads to strongly dosalized *Xenopus* embryos, with enlarged heads and cement glands³⁵⁻
92 ³⁷. In contrast, overexpression of *rspo2* failed to induce enlarged heads but instead induced *spina*
93 *bifida* with reduced head structures, yielding the first indication that *rspo2* does not act by the
94 common mode of sequestering BMP ligands (Supplementary Fig. 2c).

95 To confirm the *rspo2* morpholino data, we used a previously established guide RNA (gRNA)²⁷
96 to generate Crispr-Cas9-mediated *Xenopus rspo2* knockout (KO) embryos (Supplementary Fig.
97 3a-e). We then established gRNAs to generate Crispr-Cas9 mediated knockouts of the BMP
98 antagonists *chordin* (*chd*) and *noggin* (*nog*) (Supplementary Fig. 3a-e), whose microinjection
99 with Cas9 protein yielded mildly ventralized embryos, which were rescued by *chordin* or *noggin*
100 DNA, validating the specificity of the gRNAs (Supplementary Fig. 3f-i). Injection of *rspo2*
101 gRNA with Cas9 protein resulted in mildly ventralized embryos (Fig. 2g-h, Supplementary Fig.
102 4a-b) and increased BMP target gene (*sizzled*, *vent1*) expression, similar to knockouts of *chordin*
103 or *noggin* (Supplementary Fig. 4c-f). Importantly, combined injection of *rspo2* gRNA with either
104 *chordin* or *noggin* gRNAs yielded strongly ventralized embryos (Fig. 2g-h, Supplementary Fig.
105 4a-b) and hyperactivated BMP signaling (Supplementary Fig. 4c-f). Moreover, injection of *rspo3*
106 mRNA rescued *bmp4*-mediated increase of *sizzled* expression, suggesting that overexpressed
107 *rspo3* is also able to antagonize BMP signaling in *Xenopus* (Supplementary Fig. 4g-h), as in
108 HEPG2 cells (Fig. 1a). We conclude that *rspo2* is required to antagonize BMP signaling and acts
109 in concert with BMP antagonists for proper axial patterning during *Xenopus* embryogenesis.

110

111 **Rspo2 is a negative feedback regulator in the *Xenopus* BMP4 synexpression group**

112 In early vertebrate embryos, genes belonging to certain signaling networks form characteristic
113 synexpression groups, i.e. genetic modules composed of genes that show tight spatio-temporal
114 RNA coexpression and that function in the respective signaling pathway⁴⁰. A well-characterized
115 example is the BMP4 synexpression group, members of which are expressed like this growth
116 factor—dorsally in the eye, heart and proctodeum of tailbud stage *Xenopus* embryos (Fig. 3a).
117 This group consists of at least eight members, which all encode positive or negative feedback
118 components of the BMP signaling cascade as studied in early development, including ligands,
119 receptors and downstream components of the pathway⁴¹. Interestingly, we found that *rspo2* is
120 part of the BMP4 synexpression group, being coexpressed with *bmp4* from gastrula to tadpole
121 stages (Fig. 3a), suggesting that its expression depends on BMP signaling as for other
122 synexpressed genes. To test this idea, we employed *Xenopus* animal cap explants, which express
123 low levels of *rspo2* and *bmp4* to monitor *rspo2* induction upon *bmp4* overexpression (Fig. 3b).
124 Indeed, *bmp4* induced *rspo2* expression by qRT-PCR (Fig. 3c) and *in situ* hybridization (Fig. 3d-
125 e), similar to *bmp4* direct targets *sizzled* (Fig. 3c-e) and *vent1* (Fig. 3c). To test whether *rspo2* is
126 an immediate early target of BMP4, we blocked protein synthesis with cycloheximide (CHX)⁴¹.
127 Interestingly, while induction of the direct BMP4 targets *sizzled* and *vent1* by *bmp4* was
128 unaffected by CHX, *rspo2* induction was inhibited (Fig. 3b-e). We conclude that *rspo2* is a
129 negative feedback inhibitor within the BMP4 synexpression group and that it is an indirect BMP
130 target gene, whose expression may depend on transcription factors of the e.g. Vent or Msx
131 families^{41, 42} (Fig. 3f).

132

133 **RSPO2 and -3 bind BMPR1A via the TSP1 domain to antagonize BMP signaling**

134 Given that RSPOs act by promoting receptor endocytosis^{14, 17}, we postulated that RSPO2 might
135 regulate BMP signaling through its receptors: ACVR1, BMPR1A and BMPR1B. To test this
136 hypothesis, we analyzed the effect of RSPO1-4 treatment on BMP signaling induced by
137 constitutively active ACVR1/BMPR1A/BMPR1B (ACVR1/BMPR1A/BMPR1B^{QD}).

138 Interestingly, RSPO2 and -3 treatment specifically inhibited BMPR1A^{QD} but not ACVR1^{QD} or
139 BMPR1B^{QD}, while RSPO1 and -4 had no effect to any of the constitutively active receptors (Fig.
140 4a-c).

141 Indeed, cell surface binding assay and *in vitro* binding assay revealed that RSPO2 and -3, but not
142 RSPO1 and -4, bound the extracellular domain (ECD) of BMPR1A (Fig. 4d-e, Supplementary
143 Fig. 5a). RSPO2 showed high affinity with BMPR1A ECD ($K_d \approx 4.8$ nM) (Fig. 4f), comparable
144 to the RSPO-LGR interaction²⁴. To further delineate the domains required for BMPR1A
145 binding, we analyzed deletion mutants of RSPO2 in cell surface binding assays with BMPR1A
146 ECD, and found BMPR1A binding required the TSP1- but not the FU domains of RSPO2, while,
147 conversely, LGR binding required the FU domains but not TSP1 (Supplementary Fig. 5b-c). The
148 importance of the TSP1 domain was confirmed by *in vitro* binding assay showing that the
149 isolated TSP1 domain of RSPO2, but not RSPO1, was sufficient to interact directly with
150 BMPR1A ECD (Fig. 4g-h). Similarly, BMPR1A binding required the TSP1 domain also in
151 RSPO3, suggesting that an analogous mode of binding applies to RSPO2 and -3 (Supplementary
152 Fig. 5d-e). Our results indicate that the specificity for the RSPO-BMPR1A interaction resides in
153 the TSP1 domain of RSPOs. Consistently, the RSPO1 TSP1 domain shows only 43% and 50%

154 sequence similarity to RSPO2 and RSPO3 respectively²⁸. We next asked whether TSP1-domain
155 swapping could convey BMP signaling inhibition to RSPO1. To this end, we generated a RSPO1
156 chimera (R1-TSP^{R2}) possessing the TSP1 domain of RSPO2 (Fig. 4i). R1-TSP^{R2} activated WNT
157 signaling (Fig. 4j) and interacted with LGR4 (Supplementary Fig. 5f). However, unlike wild-type
158 RSPO1, R1-TSP^{R2} bound to BMPR1A (Supplementary Fig. 5f) and antagonized BMP signaling,
159 mimicking the effects of RSPO2 (Fig. 4k).

160 The importance of the TSP1 domain in BMP inhibition was further corroborated in *Xenopus*,
161 where we took advantage of the fact that the TSP1-domain is encoded by a distinct exon in the
162 3'-end of the *rspo2* gene. We generated a *rspo2* Mo (*rspo2*^{ΔTSP} Mo), which specifically abolished
163 TSP1-domain splicing, yielding 3' truncated *rspo2* mRNA lacking the TSP1 domain but
164 retaining the FU domains (Fig. 5a). Microinjection of *rspo2*^{ΔTSP} Mo resulted in ventralized
165 tadpoles with shorter axis and reduced heads compared to control tadpoles, which was partially
166 rescued by introducing a non-targeted *rspo2* mRNA (Supplementary Fig. 6a-b). *rspo2*^{ΔTSP}
167 Morphants had no effect on WNT signaling (Fig. 5b), confirming that it does not interfere with
168 Rspo2 FU domains that are essential for WNT activation. However, *rspo2*^{ΔTSP} Mo increased
169 BMP signaling (Fig. 5c). Similar to *chordin* and *rspo2* Morphants, *rspo2*^{ΔTSP} Morphants showed
170 expanded expression of the BMP target genes *vent1* and *sizzled* in gastrulae (Fig. 5d-e,
171 Supplementary Fig. 6c-d)³⁸, and corresponding tadpoles were ventralized, displaying decreased
172 *bfl* and *myoD*- and increased *sizzled* expression (Fig. 5f-g)³⁸. Coexpression of dominant negative
173 *bmpr1a* (*bmpr1a*^{DN}) rescued these defects (Fig. 5d-g, Supplementary Fig. 6c-d). Taken together,
174 these results emphasize that the TSP1 domain is a key element in providing target specificity to
175 RSPOs, both *in vitro* and *in vivo*, and that it dictates their BMP-inhibitory function.

176

177 **RSPO2 destabilizes the BMP receptor BMPR1A**

178 To investigate the consequence of RSPO-BMPR1A binding, we monitored BMPR1A protein
179 levels upon *RSPO2* knockdown in H1581 cells and found that si*RSPO2* treatment increased
180 BMPR1A protein levels (Fig. 6a). Similarly in *Xenopus* whole embryos, microinjection of
181 mRNA encoding *rspo2* but not *rspo2*^{ΔFU1/2} or *rspo2*^{ΔTSP} decreased protein levels from coinjected
182 *bmpr1a*-EYFP mRNA (Fig. 6b). Immunofluorescence microscopy (IF) of *Xenopus* animal cap
183 explants showed that *Bmpr1a*-EYFP localizes to the plasma membrane, where it was once again
184 reduced by *rspo2* but not by *rspo2*^{ΔFU1/2} or *rspo2*^{ΔTSP} mRNA (Fig. 6c-e). Focusing on *Xenopus*
185 ventrolateral marginal zone (VLMZ) explants, where endogenous *rspo2*, *bmpr1a* and *bmp4* are
186 coexpressed, showed that ablation of *rspo2* by Mo injection results in significant increase of
187 *Bmpr1a*-EYFP plasma membrane levels (Fig. 6f-h). Moreover, in VLMZ from *rspo2*^{ΔTSP}
188 Morphants, *Bmpr1a* levels were also increased (Fig. 6f-h), which was confirmed by western blot
189 analysis (Fig. 6i). Altogether, our results suggest that RSPO2 destabilizes BMPR1A.

190

191 **RSPO2 requires ZNRF3 to antagonize BMP receptor signaling**

192 We next turned to the role of the FU domains in RSPO2, which are also required for inhibition of
193 BMP signaling (Fig. 1j, Fig. 2a-b and Fig. 6b-e). FU1 and FU2 domains confer RSPO binding to
194 ZNRF3/RNF43 and LGRs, respectively²⁷. Since our results demonstrated an LGR-independent
195 mode of action (Fig. 1f), and since *rspo2* destabilized *Bmpr1a* (Fig. 6b), we hypothesized that
196 RSPO2 acts via ZNRF3/RNF43 E3 ligases to interfere with BMPR1A. ZNRF3 and RNF43 were

197 both expressed in HEPG2 and H1581 cells, and could be significantly knocked down by siRNA
198 (Supplementary Fig. 7a). Knockdown of *ZNRF3/RNF43* (Fig. 7a) or expression of a dominant
199 negative ZNRF3 (*ZNRF3^{ΔR}*)²⁶ (Fig. 7b) prevented inhibition of BMP signaling by RSPO2 in
200 HEPG2 cells, supporting that RSPO2 requires ZNRF3/RNF43 to antagonize BMP signaling. In
201 *Xenopus*, *znrf3* was broadly expressed from gastrula stages onwards, like *bmpr1a*
202 (Supplementary Fig. 7b-c). *znrf3* ablation by Mo elicited head and axis defects that were rescued
203 by coinjection of human *ZNRF3* mRNA, as previously described⁴³ (Supplementary Fig. 7d-e).
204 Interestingly, *znrf3* Morphants at neurula showed increased BMP signaling by BMP-reporter
205 assay and *rspo2* mRNA coinjection could not reduce it (Fig. 7c). Moreover, IF in *Xenopus*
206 animal cap explants showed that *rspo2*-induced destabilization of Bmpr1a protein levels was
207 prevented by *ZNRF3^{ΔR}* (Fig. 7d-e). Altogether, these results support that to function as BMP
208 antagonist, RSPO2 requires ZNRF3.

209

210 **RSPO2 requires the FU1 but not FU2 domain to antagonize BMP signaling**

211 To corroborate that to function as BMP antagonist, RSPO2 depends on ZNRF3/RNF43, but not
212 on LGRs, we next generated deletion mutants of the FU1 and FU2 domains in human RSPO2,
213 which mediate binding to ZNRF3/RNF43 and LGRs, respectively²⁷ (Supplementary Fig. 8a).
214 RSPO2^{ΔFU1} lost ZNRF3 binding (Supplementary Fig. 8b), yet it bound LGR4 (Supplementary
215 Fig. 8c), but did not inhibit BMP4 signaling (Fig. 7f). Conversely, RSPO2^{ΔFU2} bound ZNRF3 but
216 not to LGR4 (Supplementary Fig. 8b-c), yet it still antagonized BMP4 signaling (Fig. 7g). To
217 corroborate LGR-independent function *in vivo*, we generated *Xenopus* Rspo2^{ΔFU1} and FU2 point
218 mutant Rspo2^{F107E} (Supplementary Fig. 8d)¹⁷, which displayed ZNRF3 and LGR4 binding

219 characteristics like human RSPO2 mutants ([Supplementary Fig. 8e-f](#)). IF in *Xenopus* animal cap
220 explants injected with *bmpr1a*-EYFP and either *rspo2* wildtype or *rspo2* mutants confirmed that
221 FU1 but not FU2 deletion eliminates the ability of Rspo2 to remove plasma membrane Bmpr1a
222 ([Fig. 7h-i](#)). Taken together, our results clearly indicate that the FU1 mediated ZNRF3/RNF43
223 binding is crucial while FU2 mediated LGR binding is dispensable for RSPO2 to antagonize
224 BMP receptor signaling.

225

226 **RSPO2 bridges BMPR1A and ZNRF3 and triggers BMP receptor clearance from the cell** 227 **surface**

228 The interaction of RSPO2 and RSPO3 with BMPR1A as well as ZNRF3, suggested that R-
229 spondins bridge both transmembrane proteins. *In vitro* binding assays ([Fig. 8a-b](#)) and
230 colocalization by IF ([Fig. 8c-d](#), [Supplementary Fig. 9a-b](#)), confirmed that ZNRF3 interacted with
231 BMPR1A in the presence of RSPO2 or RSPO3 but not of RSPO1. Emphasizing once again the
232 importance of the FU1- and TSP1 domains for this interaction, *in vitro* ZNRF3-BMPR1A-
233 RSPO2 ternary complex formation was prevented by TSP1-, FU1/2-, or FU1 deletion
234 ([Supplementary Fig. 9c-g](#)), whereas it remained intact upon FU2 deletion ([Supplementary Fig.](#)
235 [9h](#)).

236 Since ZNRF3/RNF43 eliminate WNT receptors from the cell surface by co-internalization and
237 lysosomal degradation^{25, 26}, we considered an analogous function in BMPR1A turnover. We
238 monitored BMPR1A localization by IF in H1581 cells and found that it was absent from the
239 plasma membrane but abundantly colocalized with ZNRF3 in cytoplasmic vesicles ([Fig. 8e, i](#)),

240 suggesting that it may be internalized by endogenous RSPO2. Indeed, upon knockdown of
241 *RSPO2*, but not *LRP6* or *LGR4/5*, BMPR1A accumulated at the plasma membrane (Fig. 8f-i).
242 Importantly, IF (Fig. 8j-m) and cell surface biotinylation assays (Fig. 8n) showed that upon
243 *ZNRF3/RNF43* siRNA treatment, BMPR1A also accumulated at the plasma membrane.

244 To test if RSPO2/ZNRF3 target BMPR1A for endocytosis and lysosomal degradation, we treated
245 cells with the clathrin inhibitor monodansylcadaverin (MDC), which eliminated inhibition of
246 BMP signaling by RSPO2 (Fig. 8o). In addition, si*RSPO2* abolished the colocalization of
247 BMPR1A with the early endosome marker EEA1 (Fig. 8p-q) and lysosomal marker Lamp1 (Fig.
248 8r-s), suggesting that RSPO2 binding promotes BMPR1A internalization and degradation via
249 ZNRF3 ternary complex formation. Consistently, 20 min exposure to RSPO2 increased
250 internalized BMPR1A in cell surface biotinylation assays in H1581 cells (Supplementary Fig.
251 10a) and induced vesicular Bmpr1a-EYFP in *Xenopus* animal caps (Supplementary Fig. 10b-c).
252 Taken together, our results support a model (Supplementary Fig. 10d) wherein RSPO2 bridges
253 ZNRF3 and BMPR1A and routes the ternary complex towards clathrin-mediated endocytosis for
254 lysosomal degradation, thereby antagonizing BMP signaling. We suggest that a similar
255 mechanism applies to RSPO3 but not RSPO1 and -4.

256

257 **DISCUSSION**

258 The three main findings of our study are i) the discovery R-spondins as a novel class of BMP
259 receptor antagonists, ii) that RSPO2 depletes BMPR1A/ALK3 by engaging ZNRF3 for
260 internalization and lysosomal degradation, and iii) that in *Xenopus*, *rspo2* is a negative feedback

261 inhibitor of the BMP4 synexpression group, which cooperates with Spemann organizer effectors
262 to inhibit BMP signaling during axis formation. Given the importance of RSPOs and BMPs as
263 developmental regulators, as well as growth factors of normal and malignant stem cells, these
264 conclusions have implications for development and cancer.

265 With regard to stem cells, R-spondins are a key ingredient of the culture media, which have
266 made the organoid revolution possible^{21, 22} and their rational use requires an understanding of
267 their mechanism of action. For example, the fact that R-Spondins inhibit BMP signalling may
268 explain the reported non-equivalence of WNT and RSPO ligands in stem cells and
269 development³¹⁻³³. It may also explain their potency as stem cell growth factors, as e.g. intestinal
270 stem cells requires both, WNT activation and BMP inhibition^{21, 22}.

271 TGF β growth factors play an eminent role in biology and medicine, and their receptor signalling
272 is exquisitely regulated extracellularly with over 20 TGF β antagonists, most of which antagonize
273 signaling by ligand sequestration (e.g. Cerberus, Chordin, Follistatin, Gremlin, Noggin, and Sost)
274 ^{1, 11}. Two extracellular BMP receptor antagonists are known, BMP3 and Inhibin^{12, 13}. Both are
275 TGF β family members, whose unproductive binding to type II receptors prevents signal
276 transmission. Relatedly, the BMP antagonist BAMBI is a BMP pseudoreceptor lacking kinase
277 activity, which also leads to formation of a dead-end complex with BMP receptors⁴⁴. In contrast,
278 RSPO2 and -3 share no sequence homology with TGF β family members, they inhibit type I
279 instead of type II BMP receptors, and they do so by a novel mechanism, which engages the
280 ZNRF3 E3 transmembrane ubiquitin ligase to internalize BMPR1A. RSPO2 thereby routes
281 BMPR1A to clathrin-mediated endocytosis for lysosomal degradation. This mode of action

282 resembles the function of the Spastic Paraplegia related gene NIPA1, a transmembrane
283 antagonist, which promotes BMP receptor type II endocytosis and lysosomal degradation⁴⁵.

284 Other type I BMP receptors besides BMPR1A include ACVRL1, ACVR1 and BMPR1B⁷.
285 However, we found that RSPO2 specifically binds to BMPR1A but not to ACVR1 or BMPR1B
286 (data not shown), which explains why ACVR1 and BMPR1B signaling were not antagonized by
287 RSPO2 in human cells (Fig. 4a-c). Consistently, BMPR1A and e.g. BMPR1B only show 42%
288 identity in their extracellular domain⁴⁶.

289 BMPR1A engages not only various BMPs but also GDFs¹, and hence RSPO-mediated inhibition
290 may potentially affect signalling in multiple contexts. On the other hand, the specificity of
291 RSPO2 for BMPR1A may provide therapeutic opportunities on the background of pleiotropic
292 BMP ligands effects.

293 RSPO2 engages ZNRF3 to antagonize BMP signaling, implying that ZNRF3 is also a negative
294 regulator not only of WNT, but also BMP signaling. Consistently, our results indicate that loss of
295 *ZNRF3* increases BMP signaling. Moreover, *ZNRF3* overexpression induces expression of
296 Spemann organizer genes in *Xenopus* embryos, which is characteristic not only for WNT but
297 also BMP inhibition⁴³. In WNT signalling, the role of RSPO2 is to protect WNT receptors from
298 ubiquitination and internalisation by ZNRF3, by forming a ternary complex with LGR4-6 and
299 triggering endocytosis. In contrast, during BMP signalling, RSPO2 directly forms a ternary
300 complex with ZNRF3-BMPR1A to internalize and degrade the type I receptor. Our data also
301 imply a possible function of RNF43 in antagonizing BMP signalling, inviting a closer inspection
302 of its loss-of-function phenotypes^{25, 26}.

303 A number of studies emphasized the importance of the Furin domains in RSPOs, which are
304 necessary and sufficient for activation of WNT signaling^{17, 20, 26, 28}, however, the role of the TSP1
305 domain has received less attention. We found that the specificity of RSPOs for BMP signaling is
306 dictated by the TSP1 domain, which binds directly to BMPR1A. Unlike RSPO2 and -3, RSPO1
307 and -4 do not inhibit BMP signaling, the key difference residing in the TSP1 domain, as domain
308 swapping of the TSP1 domain is sufficient to confer BMP inhibition upon RSPO1. The
309 physiological role *in vivo* is highlighted by *rspo2* Morphants specifically lacking the TSP1
310 domain, which displayed phenotypic defects due to BMP hyperactivation (Fig. 5). The TSP1
311 domain also binds to heparin sulfate proteoglycans (HSPG) e.g. syndecans (SDC)^{27, 29, 30}, which
312 raises the possibility of cooperation between RSPOs and SDC in BMP receptor regulation.
313 Indeed, SDC1 and SDC3 have been implicated as negative regulators in BMP signaling, but the
314 underlying mechanisms remained unclear^{47, 48}. Hence, it will be interesting to investigate the role
315 of SDCs in BMPR1A-RSPO interactions. HSPGs are also coreceptors in FGF signaling, which
316 may explain why misexpressed *rspo2* can inhibit FGF signaling in *Xenopus* animal cap
317 explants⁴⁹.

318 We established that *Xenopus* Rspo2 cooperates with Noggin and Chordin released by the
319 Spemann organizer in repressing BMP signaling to modulate the BMP morphogen gradient,
320 which controls axial patterning. Yet, overexpression of *rspo2* unlike of *noggin* and *chordin*, does
321 not strongly dorsalize early embryos. The reason is that instead of sequestering BMP ligands,
322 RSPO2 specifically targets the BMP receptor BMPR1A in early *Xenopus* embryos and that
323 BMPR1A and BMPR1B play overlapping roles in dorsoventral patterning^{50, 51}. Also unlike
324 *noggin* and *chordin*, *rspo2* is not expressed in the organizer but is a negative feedback inhibitor
325 of the BMP4 synexpression group, similar to the BMP pseudoreceptor *bambi*^{44, 52}. Like *bambi*,

326 *rspo2* is an indirect BMP4 target gene, which may require Vent- or Msx transcription factors for
327 expression. Negative feedback in BMP signaling expands the dynamic BMP signaling range
328 essential for proper embryonic patterning and reduce inter-individual phenotypic and molecular
329 variability in *Xenopus* embryos⁴². Indeed, *rspo2* deficiency by itself has only mild effects on axis
330 formation and dorsoventral marker gene expression, while defects manifest upon misbalance of
331 BMP signaling (*bmp4*-overexpression, *noggin/chordin* knockdown). Functional redundancy
332 between BMP antagonists is a characteristic feature observed in fish, frog, and mouse embryos⁵³⁻
333 ⁵⁶. We note that the mouse *Rspo2* expression pattern at E9.5 mimics that of mouse *Bmp4*,
334 including forebrain, midbrain/hindbrain junction, branchial arches and limb apical ectodermal
335 ridge^{57, 58}. Thus, although *Rspo2* deficient mouse embryos gastrulate normally⁵⁹, it may be
336 fruitful to analyze compound mutants between *Rspo2* and BMP antagonists for axial defects. The
337 fact that R-Spondins are bifunctional ligands, which activate WNT- and inhibit BMP signalling
338 has implications for development, stem cell biology, and cancer. Mechanistically, the general
339 picture emerging is that R-Spondins function as adapters, which escort client extracellular
340 proteins for ZNRF3/RNF43-mediated degradation, e.g. LGR4-6 and BMPRI1A. Our results
341 assign a key role to the largely ignored TSP1 domain of R-Spondins in providing target
342 specificity. The substantial sequence variability between TSP1 domains of RSPO1-4 invites
343 screening for additional RSPO receptor targets beyond BMPRI1A.

344

345 **METHODS**

346 **Cell lines and growth conditions**

347 HEK293T and HEPG2 cells (ATCC) were maintained in DMEM High glucose (Gibco 11960)
348 supplemented with 10% FBS (Capricorn FBS-12A), 1% penicillin-streptomycin (Sigma P0781),
349 and 2 mM L-glutamine (Sigma G7513). H1581 cells (gift from Dr. R.Thomas) were maintained
350 in RPMI (Gibco 21875) with 10% FBS, 1% penicillin-streptomycin, 2 mM L-glutamine and 1
351 mM sodium pyruvate (Sigma S8636). Mycoplasma contamination was negative in all cell lines
352 used.

353 ***Xenopus laevis* and *Xenopus tropicalis*** *Xenopus laevis* frogs were obtained from Nasco.

354 *Xenopus tropicalis* frogs were obtained from Nasco, National *Xenopus* Resource (NXR) and
355 European *Xenopus* Resource Centre (EXRC).

356 All *X.laevis* and *X.tropicalis* experiments were approved by the state review board of Baden-
357 Württemberg, Germany (permit number G-141-18) and executed according to federal and
358 institutional guidelines and regulations. Developmental stages of the embryos were determined
359 according to Nieuwkoop and Faber (Xenbase). No statistical analysis was done to adjust sample
360 size before the experiments. No randomization of injection order was used during the
361 experiments.

362 **Constructs**

363 Alkaline phosphatase (AP) fusions with RSPOs (human RSPO1^{ΔC}-AP-pCDNA3, RSPO2^{ΔC}-AP-
364 pCDNA3, RSPO2^{ΔC}-AP-pCS2+, RSPO3^{ΔC}-AP-pCDNA3, murine RSPO4^{ΔC}-pCDNA3) were
365 generated by replacing the C-terminal domain (ΔC) by AP and used to produce conditioned
366 media. Human RSPO2 wild-type (RSPO2), the Furin1 and the Furin2 domain deletion mutants

367 (RSPO2^{ΔFU1/2}), and the TSP1 domain deletion mutant (RSPO2^{ΔTSP}) are ORFs lacking the C-
368 terminal domain, C-terminally tagged with a Flag-tag and subcloned into pCS2+²⁰. R1-TSP^{R2},
369 R1-TSP^{R2}-AP and R1-TSP^{R2}-Flag plasmids were cloned in pCS2+. Human RSPO2^{ΔFU1} (deletion
370 of amino acids encompassing the 6 cystines in the FU1 domain) and RSPO2^{ΔFU2} mutants
371 (deletion of amino acids encompassing the 8 cystines in the FU2 domain) were cloned in Flag-
372 tag or AP-tag pCS2+. Human RSPO1^{TSP1} and RSPO2^{TSP1}-HA were cloned in Streptag-HA-flag-
373 pCS2+. *Xenopus* Rspo2^{ΔFU1} (deletion of amino acids encompassing the 6 cystines in the FU1
374 domain) and Rspo2^{F107E} mutants were cloned in Myc-tag or AP-tag pCS2+. All constructs were
375 confirmed by sequencing. Conditioned media from all RSPO constructs were adjusted to equal
376 concentration by western blot and AP activity measurement, and further validated by WNT
377 reporter assay using HEK293T cells. The extracellular domain of BMPR1A (BMPR1A^{ECD}) was
378 subcloned in AP-pCS2+ for generating conditioned medium and used in *in vitro* binding assays.
379 Constitutively active forms of ACVR1, BMPR1A, and BMPR1B (QD) were generated by Gln-
380 Asp mutations as described⁶⁰. HA-tagged BMPR1A/ALK3 was a gift from Dr. D.Koinuma⁶¹.

381 For *Xenopus* mRNA microinjection, *Xenopus laevis* Bmp4-pCS2+, Rspo2^{ΔC}-myc-pCS2+,
382 Rspo2^{ΔFU1/2}-myc-pCS2+ and Rspo2^{ΔTSP}-myc-pCS2+ plasmids, Bmpr1a^{DN}-pCS2+, membrane-
383 RFP, Bmpr1a-EYFP-pCS2+, Rspo2^{ΔFU1}-myc-pCS2+, Rspo2^{F107E}-myc-pCS2+ were used for *in*
384 *vitro* transcription. Human Noggin-AP-pCS2+ and Chordin-AP-pCS2+ plasmids were used for
385 *Xenopus tropicalis* Crisphant rescue assay. Human ZNRF3 and ZNRF3^{ΔRING} constructs were
386 gifts from Dr. F.Cong (Novartis)²⁶, and ORFs were further subcloned in flag-pCS2+ for *in vitro*
387 transcription.

388 Cell transfection

389 For HEPG2 and H1581 cells, siRNAs and plasmids were transfected using DharmaFECT 1
390 transfection reagent (Dharmacon T-2001) and Lipofectamine 3000 (Invitrogen L3000)
391 respectively, according to the manufacturer protocols. For HEK293T cells, X-tremeGENE 9
392 DNA transfection reagent (Roche 6365787001) was used, according to the manufacturer
393 protocols.

394 **Generation of conditioned medium**

395 HEK293T cells were seeded in 15 cm culture dishes and transiently transfected with RSPOs-AP,
396 RSPOs-flag, BMPR1A^{ECD}-AP, DKK1 or WNT surrogate plasmids. After 24 hours, media were
397 changed with fresh DMEM, 10% FBS, 1% L-glutamine and 1% penicillin-streptomycin and
398 cultured 6 days at 32 °C. Conditioned media were harvested three times every two days,
399 centrifuged and validated by TOPFlash assay or western blot analyses. WNT3A conditioned
400 medium was produced in L-cells as previously described²⁰. For human RSPO2^{ΔFU1},
401 RSPO2^{ΔFU2}, *Xenopus* Rspo2^{ΔFU1} and Rspo2^{F107E} mutants conditioned media, HEK293T cells
402 were seeded in 12 well culture plates and transfected with 500 μg of each plasmid, and harvested
403 three times every two days. Production of the media was validated with western blot analyses
404 and AP activity analyses.

405 **Luciferase reporter assays**

406 BRE luciferase assays were executed using 300,000 ml⁻¹ of HEPG2 cells in 24-well plates.
407 PGL3-BRE-Luciferase (500 ng ml⁻¹) and pRL-TK-Renilla plasmids (50 ng ml⁻¹) were
408 transfected using Lipofectamine 3000. After 24 hours, cells were serum starved 2 hours and
409 stimulated 14-16 hours with 80 ng ml⁻¹ recombinant human BMP4 protein (R&D systems 314-
410 BP) along with RSPO1-4 conditioned medium. Luciferase activity was measured with the Dual

411 luciferase reporter assay system (Promega E1960). Firefly luminescence (BRE) was normalized
412 to Renilla. TOPFlash luciferase assays were carried out as previously described⁶². Data are
413 displayed as average of biological replicates with SD. Statistical analyses were made with the
414 PRISM7 software using unpaired t-test or one-way ANOVA test. Not significant (ns) $P > 0.05$,
415 * $P < 0.05$ ** $P < 0.01$, *** $P < 0.001$, and **** $P < 0.0001$.

416 **Western blot analysis**

417 Cultured cells were rinsed with cold PBS and lysed in Triton lysis buffer (20 mM Tris-Cl, pH
418 7.5, 1% Triton X-100, 150 mM NaCl, 1 mM EDTA, 1 mM EGTA, 1 mM β -glycerophosphate, 1
419 mM Na_3VO_4) or RIPA buffer with cOmplete Protease Inhibitor Cocktail (Roche 11697498001).
420 Lysates were mixed with Laemmli buffer containing β -mercaptoethanol and boiled at 95 °C for 5
421 min to prepare SDS-PAGE samples. Western blot images were acquired with SuperSignal West
422 pico ECL (ThermoFisher 34580) or Clarity Western ECL (Biorad 1705061) using LAS-3000
423 system (FujiFilm). Quantification of blots was done using ImageJ software.

424 **Cell surface biotinylation assay**

425 H1581 cells were seeded in 6 cm culture dishes and transfected with 50 nM of indicated siRNAs
426 for 3 days and 2 μg of BMPR1A-HA DNA for 2 days. Surface proteins were biotinylated with
427 0.25 mg ml^{-1} sulfo-NHS-LC-LC-Biotin (ThermoFisher 21338) at 4 °C for 30 min. The reaction
428 was quenched by 10 mM Monoethanolamine and cells were harvested and lysed with Triton X-
429 100 lysis buffer. 200-300 μg of lysate was incubated with 20 μl streptavidin agarose
430 (ThermoFisher 20359) to pull-down biotinylated surface proteins and subjected to Western blot.

431 **Surface receptor internalization assay**

432 H1581 cells were seeded in 15 cm culture dish and transfected with 10 µg of BMPR1A-HA
433 DNA for 2 days, and then split to 6 cm culture dishes. After 24 h, surface proteins were
434 biotinylated with 0.5 mg ml⁻¹ sulfo-NHS-SS-Biotin (ThermoFisher 21331) at 4 °C for 30 min.
435 After quenching excessive biotin with 10 mM monoethanolamine, pre-warmed control medium
436 or RSPO2 conditioned medium was added at 37 °C to induce internalization. After 20 min
437 stimulation, remaining surface-biotin was removed by 50 mM MesNa (2-
438 mercaptoethanesulfonate, membrane impermeable reducing agent, CAYMAN 21238) in MesNa
439 reaction buffer (100 mM Tris-HCl, pH 8.6, 100 mM NaCl and 2.5 mM CaCl₂) at 4 °C for 30 min
440 and MesNa protected-biotinylated proteins (internalized proteins) were analyzed. Cells were
441 harvested, and lysed with RIPA buffer (20 mM Tris-Cl, pH 7.4, 120 mM NaCl, 1% Triton X-
442 100, 0.25% Na-deoxycholate, 0.05% SDS, 50 mM sodium fluoride, 5 mM EDTA, 2 mM Na-
443 orthovanadate) supplemented with complete protease inhibitor. 500 µg lysate was incubated with
444 20 µl streptavidin agarose (ThermoFisher 20359) to pull-down biotinylated proteins and
445 subjected to Western blot.

446 ***Xenopus laevis* whole-mount *in situ* hybridization**

447 Whole-mount *in situ* hybridizations of *Xenopus* embryos were performed using digoxigenin
448 (DIG)-labeled probes as previously described⁶³. Antisense RNA probes against *rspo2* and *bmp4*
449 were generated by *in vitro* transcription as previously described²⁰. Probes against *bmpr1a* and
450 *znrf3* were prepared using full-size *Xenopus* *bmpr1a* ORF or *znrf3* ORF as a template. Mo and
451 mRNA injected embryos were collected at stage 11 (gastrula) or 32 (tadpole) for *in situ*
452 hybridization. Images were obtained using AxioCam MRc 5 microscope (Zeiss). Embryos in

453 each image were selected using Magnetic Lasso tool or Magic Wand tool of Adobe Photoshop
454 CS6 software, and pasted into the uniform background color for presentation.

455 ***Xenopus* microinjection and phenotype analysis**

456 *In vitro* fertilization, microinjection and culture of *Xenopus* embryos were performed as
457 previously described ⁶³. *X.laevis* embryos were microinjected with reporter DNAs, *in vitro*
458 transcribed mRNAs or antisense morpholino oligonucleotide (Mo) using Harvard Apparatus
459 microinjection system. Mos for *rspo2* ²⁰, *lrp6* ³⁹, *chordin*, *bmp4* ³⁸, *znrf3*⁴³ and standard control
460 were purchased from GeneTools. *rspo2*^{ΔTSP} Mo was designed based on *rspo2* sequence
461 ([Supplementary Table 3](#)). *X.laevis* 4-cell stage embryos were microinjected 5 nl per each
462 blastomere equatorially and cultured until indicated stages. Equal amount of total mRNA or Mo
463 were injected by adjustment with *ppl* or standard control Mo. Scoring of phenotypes was
464 executed blind from two individuals, and data are representative images from at least two
465 independent experiments. Embryos in each image were selected using Magnetic Lasso tool or
466 Magic Wand tool of Adobe Photoshop CS6 software, and pasted into the uniform background
467 color for presentation. Statistical analyses show Chi-square tests.

468 ***Xenopus tropicalis* CRISPR/Cas9-mediated mutagenesis**

469 The 5' region of genomic sequences from *X.tropicalis chordin* (NM_001142657.1) and *noggin*
470 (NM_001171898.1) were searched for guide RNA (gRNA) targeting sites using an online
471 prediction tool (<https://crispr.cos.uni-heidelberg.de>). Primers were designed for PCR-based
472 gRNA template assembly ([Supplementary Table 4](#)) ⁶⁴, or used as previously described ²⁷. A
473 primer lacking any target sequences was used as control gRNA. PCR reactions were performed

474 with Phusion Hot Start Flex DNA Polymerase (NEB M0535), followed by *in vitro* transcription
475 using MEGAscript T7 Transcription Kit (Invitrogen AM1334). Embryos were microinjected at
476 one to two-cell stages with a mixture of 50 pg of gRNA and 1 ng of recombinant Cas9 protein
477 (Toolgen) per embryo. Injected embryos were cultured until stage 30, fixed with MEMFA for
478 phenotypical analysis. Scoring of phenotypes was executed at stage 30 with blinding from two
479 individuals, and data are representative images from three independent experiments. Defects
480 were categorized by the severity of ventralization. ‘Severe’ showed small head, enlarged ventral
481 tissues and short body axis. ‘Mild’ showed one or two of the defects described above. ‘Normal’
482 showed no visible differences to the uninjected control. Statistical analyses show Chi-square test.

483 **Injected amount of reagents per *Xenopus* embryo**

484 Equal amounts of total RNA or Mo were injected by adjustment with *preprolactin* (PPL) mRNA,
485 control gRNA or standard control Mo. Per embryo; Figure 2b, 250 pg of *bmp4*, *rspo2* and *rspo2*
486 mutants mRNA; Figure 2c, 5 ng or 10 ng of *rspo2* Mo and 5 ng of *lrp6* Mo, 300 pg of reporter
487 DNA; Figure 2d, 15 ng of *bmp4* Mo, 2, 5, or 10 ng of *rspo2* Mo, 300 pg of reporter DNA; Figure
488 2f, 15 ng of *bmp4* Mo and 5 ng of *rspo2* Mo; Figure 2h, 50 pg of gRNA, 200 pg of *bmp4*
489 mRNA, 2 ng of *lrp6* Mo, 1ng of Cas9 protein; Figure 3c and 3e, 500 pg of *bmp4*; Figure 5b and
490 5c, 20 ng of *rspo2* Mo and *rspo2*^{ΔTSP} Mo, 300 pg of reporter DNA; Figure 5e and 5g, 8 ng of *chd*
491 Mo, 20 ng of *rspo2*^{ΔTSP} Mo and 200 pg of *bmpr1a*^{DN}; Figure 6c, 500 pg of *bmp4* and *bmpr1a*-
492 EYFP, 250 pg of membrane-RFP, *rspo2*, and *rspo2* deletion mutants mRNA; Figure 6e, 500 pg
493 of *bmpr1a*-EYFP, 250 pg of *rspo2*, and *rspo2* deletion mutants, and *gfp*; Figure 6g and 6i, 10 ng
494 of *rspo2* Mo and *rspo2*^{ΔTSP} Mo; Figure 7c, 40 ng of *znrf3* Mo, 100 pg and 200 pg of *rspo2*
495 mRNA, and 300 pg of reporter DNA; Figure 7d, 500 pg of *bmp4* and *bmpr1a*-EYFP, 250 pg of

496 membrane-RFP, 250 pg of *rspo2*, 100 pg of *znrf3*^{DN} mRNA; Figure 7h, 500 pg of *bmp4* and
497 *bmpr1a*-EYFP, 250 pg of membrane-RFP, *rspo2*, and *rspo2* mutants mRNA; Supplementary
498 Figure 2b, 15 ng of *bmp4* Mo and 5 ng of *rspo2* Mo; Supplementary Figure 3g and 3i, 50 pg of
499 gRNA, 10 pg and 25 pg of chordin and noggin DNA; Supplementary Figure 4b, 4d and 4f, 50 pg
500 of gRNA, 200 pg of *bmp4* mRNA, 2 ng of *lrp6* Mo, 1 ng of Cas9 protein; Supplementary Figure
501 4h, 250 pg of *bmp4* and *rspo3* mRNA; Supplementary Figure 6b, 20 ng of *rspo2*^{ΔTSP} Mo, 150 ng
502 and 250 ng of *rspo2* mRNA; Supplementary Figure 6d, 15 ng chd Mo, 10 ng *rspo2* Mo and 50
503 pg *bmpr1a*^{DN}; Supplementary Figure 7e, 80 ng of *znrf3* Mo, 200 pg of *ZNRF3* mRNA;
504 Supplementary Figure 10c, 500 pg of *bmpr1a*-EYFP.

505 ***Xenopus tropicalis* T7 Endonuclease I assay**

506 To validate CRISPR/Cas9-mediated genome editing, three embryos of each injection set were
507 lysed at stage 30 for genotyping PCR reactions as described⁶⁴ (Supplementary Table 4). All
508 target sequences were amplified with Roti-Pol Hot-TaqS Mix (Roth 9248). After denaturation
509 for 3 min at 94 °C and reannealing (ramp 0.1 °C per sec), the PCR products were incubated with
510 3 U of T7 Endonuclease I for 45 min at 37 °C. Cleavage results were visualized on a 2 % agarose
511 gel.

512 ***Xenopus laevis* western blot analysis**

513 Injected *Xenopus* embryos were harvested at stage 15 to 18, homogenized in NP-40 lysis buffer
514 (2% NP-40, 20 mM Tris-HCl pH 7.5, 150 mM NaCl, 10 mM NaF, 10 mM Na₃VO₄, 10 mM
515 sodium pyrophosphate, 5 mM EDTA, 1 mM EGTA, 1 mM PMSF, and cOmplete Protease
516 Inhibitor Cocktail) with a volume of 20 µl per embryo. Lysates were cleared with CFC-113

517 (Honeywell 34874), followed by centrifugation (14,000 rpm, 10 min at 4 °C), boiling at 95 °C
518 for 5 min with NuPAGE Sample Buffer. 0.5-1 embryos per lane were loaded for SDS-PAGE
519 analysis.

520 **Cycloheximide treatment on *Xenopus laevis* animal cap explants**

521 *Xenopus laevis* animal caps were dissected at stage 8 and treated with 30 µg ml⁻¹ of
522 cycloheximide (CHX) (Sigma C7698) until control embryos reached stage 10. CHX treatment
523 was validated since cell division was retarded compared to untreated control. *In situ*
524 hybridization and qRT-PCR were performed with same methods used in whole embryos.

525 ***In vitro* binding assay**

526 High binding 96-well plates (Greiner M5811) were coated with 2 µg ml⁻¹ of recombinant human
527 RSPO1 (Peprotech 120-38), RSPO2 (Peprotech 120-43), RSPO3 (Peprotech 120-44), RSPO4
528 (R&D systems 4575-RS) or FGF8b (Peprotech 100-25) recombinant protein reconstituted in
529 bicarbonate coating buffer (50 mM NaHCO₃, pH 9.6) overnight at 4 °C. Coated wells were
530 washed three times with TBST (TBS, 0.1% Tween-20) and blocked with 5% BSA in TBST for 1
531 hour at room temperature. 1.5 U ml⁻¹ of BMPR1A^{ECD}-AP or control conditioned medium was
532 incubated overnight at 4 °C. Wells were washed six times with TBST and bound AP activity was
533 measured by the chemiluminescent SEAP Reporter Gene Assay kit (Abcam ab133077) or
534 AquaSpark AP substrate (Serva 42593.01). For ZNRF3-BMPR1A binding assay, plates were
535 coated with recombinant human ZNRF3 Fc Chimera protein (R&D systems 7994-RF). RSPO2-
536 flag, RSPO2 deletion mutants-flag conditioned medium, or recombinant RSPO protein was
537 preincubated 4-6 hours with ZNRF3 prior to BMPR1A^{ECD}-AP treatment. Control conditioned

538 medium and vesicles were used as control. Data show average chemiluminescent activities with
539 SD from experimental triplicates. Statistical analyses show unpaired t-tests. K_d was obtained as
540 previously described ²⁴.

541 **Immunofluorescence**

542 150,000 H1581 cells were grown on coverslips in 12-well plates, followed by siRNA and DNA
543 transfection. After 48 hours cells were fixed in 4% PFA for 10 min. Cells were treated with
544 primary antibodies (1:250) overnight at 4 °C, and secondary antibodies (1:500) and Hoechst dye
545 (1:500) were applied for 2 hours at room temperature. Tyramide Signal Amplification for
546 detecting RSPO-HRP was carried out as previously described ^{24, 39}. Quantification was executed
547 using ImageJ. Dot plots show average and SD from every cells analyzed with unpaired t-test.

548 For *X.laevis* embryos, *bmpr1a*-EYFP and membrane-RFP mRNAs were coinjected with the
549 indicated mRNAs or Mos. Embryos were dissected for animal or ventrolateral explants at stage
550 9 or stage 11.5, respectively. Explants were immediately fixed with 4% PFA for 2 hours and
551 mounted with Fluoromount-G (ThermoFisher 00495802). Images were obtained using LSM 700
552 (Zeiss). Data are representative images from two independent experiments. For quantification,
553 Pearson's correlation coefficient for EYFP and RFP was analyzed using 16-30 random areas
554 harboring 10 cells chosen from 6-10 embryos per each set. Dot plots show an average and SD
555 from every plane analyzed with unpaired t-test.

556 **Cell surface binding assay**

557 Cell surface binding assays were carried out as previously described ³⁹ with few modifications.
558 In brief, human BMPR1A-HA and *Xenopus tropicalis* LGR4 DNA were transfected in

559 HEK293T cells, and incubated with 1.5 U ml⁻¹ conditioned media for 3 hours on ice. After
560 several washes and crosslinking, cells were treated with 2 mM Levamisole for 20 min to
561 inactivate endogenous AP activities and developed with BM-Purple (Sigma 11442074001). Cells
562 were mounted with Fluoromount G. Images were obtained using LEICA DMIL
563 microscope/Canon DS126311 camera.

564 **Quantitative real-time PCR**

565 Cultured cells were lysed in Macherey-Nagel RA1 buffer containing 1% β-mercaptoethanol and
566 total RNAs were isolated using NucleoSpin RNA isolation kit (Macherey-Nagel 740955).
567 Reverse transcription and PCR amplification were performed as described before⁶². For
568 *Xenopus laevis*, animal cap explants were harvested at stage 10 and qRT-PCR was executed as
569 previously described⁴³. Primers used in this study are listed in [Supplementary Table 1](#). Graphs
570 show relative gene expressions to GAPDH. Data are displayed as mean with SD from multiple
571 experimental replicates. Statistical analyses were performed using PRISM7 software with
572 unpaired t-test or one-way ANOVA test.

573

574 **DATA AVAILABILITY**

575 All data is available from the corresponding author upon reasonable request.

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584 **AUTHOR CONTRIBUTIONS**

585 H.L. designed and conducted *in vitro*-, human cell line and *Xenopus* experiments. C.S. conducted
586 human cell line and *Xenopus* experiments. R.S. designed and carried out *in vitro*- and human cell
587 line experiments. A.G. generated materials for the study. All authors analyzed and discussed the
588 data. C.N. conceived and coordinated the study and wrote the paper with contribution from H.L.

589 **COMPETING INTERESTS**

590 The Authors declare no competing interests.

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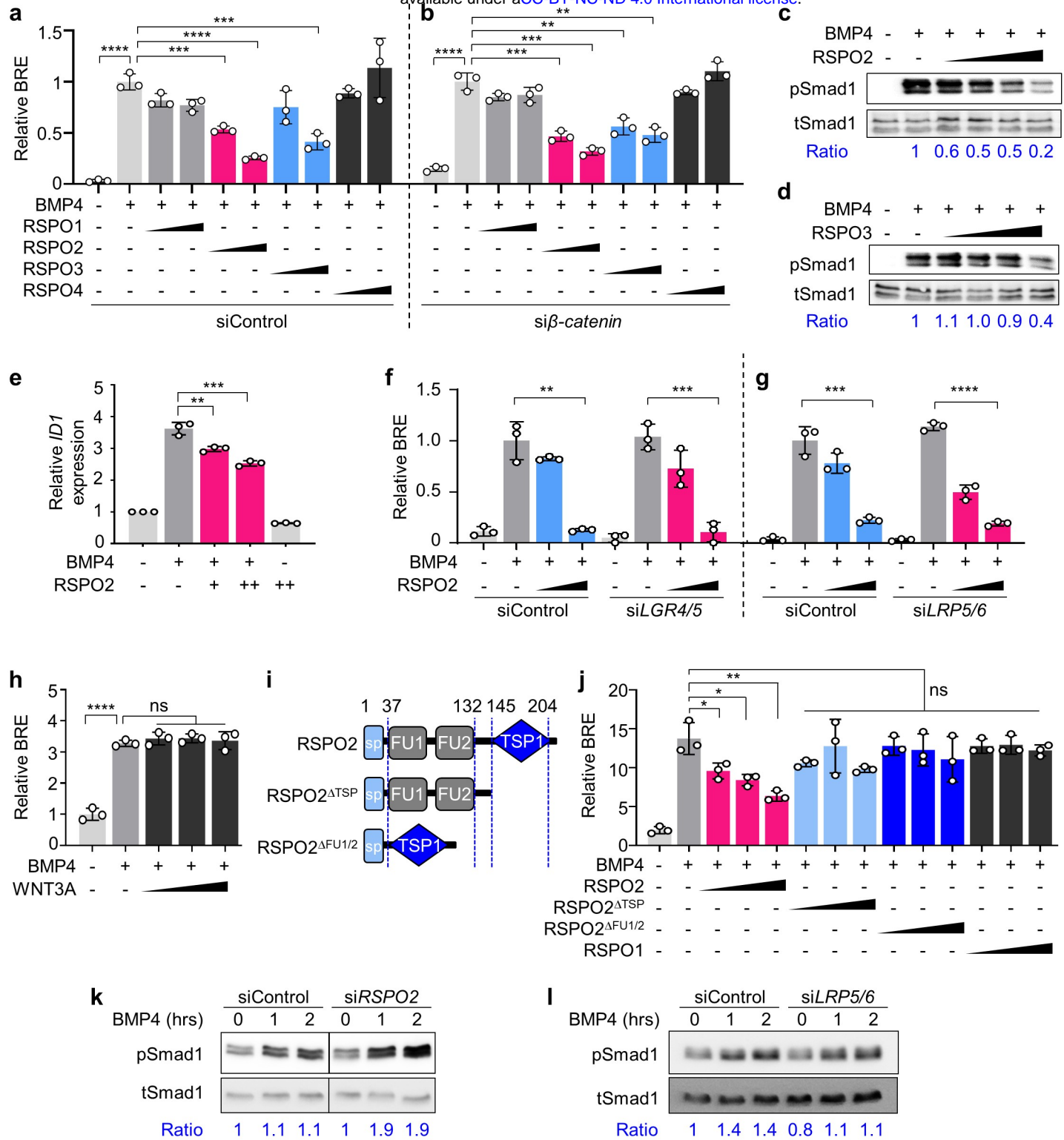


Fig. 1. RSPO2 and -3 antagonize BMP4 signaling WNT independently.

- (a, b) BRE reporter assay in HEPG2 cells upon siControl (a) or si β -catenin (b) transfection, with or without overnight BMP4 and RSPO1-4 treatment as indicated.
- (c, d) Western blot analyses of phosphorylated- (pSmad1) and total Smad1 (tSmad1) in HEPG2 cells stimulated by BMP4, treated with or without increasing amount of RSPO2 (c) or RSPO3 (d) overnight. Cells were starved 3-6 h before the stimulation. Ratio, relative levels of pSmad1 normalized to tSmad1. Representative data from two independent experiments are shown.
- (e) qRT-PCR analysis of BMP target *ID1* in HEPG2 cells upon BMP4, with or without overnight RSPO2 treatment.
- (f, g) BRE reporter assay in HEPG2 cells upon siLRP5/6 and siLGR4/5 knockdowns, with or without overnight BMP4 and RSPO2 treatment as indicated.
- (h) BRE reporter assay in HEPG2 cells stimulated overnight by BMP4 with or without increasing amount of WNT3A treatment. WNT3A activity was validated in **Supplementary Fig. 1b**.
- (i) Domain structures of RSPO2 and deletion mutants used in (j). sp, signal peptide; FU, furin domain; TSP1, thrombospondin domain 1.
- (j) BRE reporter assay in HEPG2 cells stimulated overnight with BMP4, and with or without RSPO2 WT or FU1/2- or TSP1 deletion mutants, respectively.
- (k, l) Western blot analyses of pSmad1 and tSmad1 in H1581 cells upon siRNA transfection as indicated, with 0 h, 1 h and 2 h of BMP4 stimulation. Ratio, relative levels of pSmad1 normalized to tSmad1. Representative data from two independent experiments are shown. Data for BRE reporter assays (a-b, f-h, j) are biological replicates and displayed as means \pm SD, and show a representative of multiple independent experiments. ns, not significant; *P < 0.05 **P < 0.01, ***P < 0.001, and ****P < 0.0001 from unpaired t-test or one-way ANOVA with Dunnett test. For the uncropped western blot images, see **Source file**.

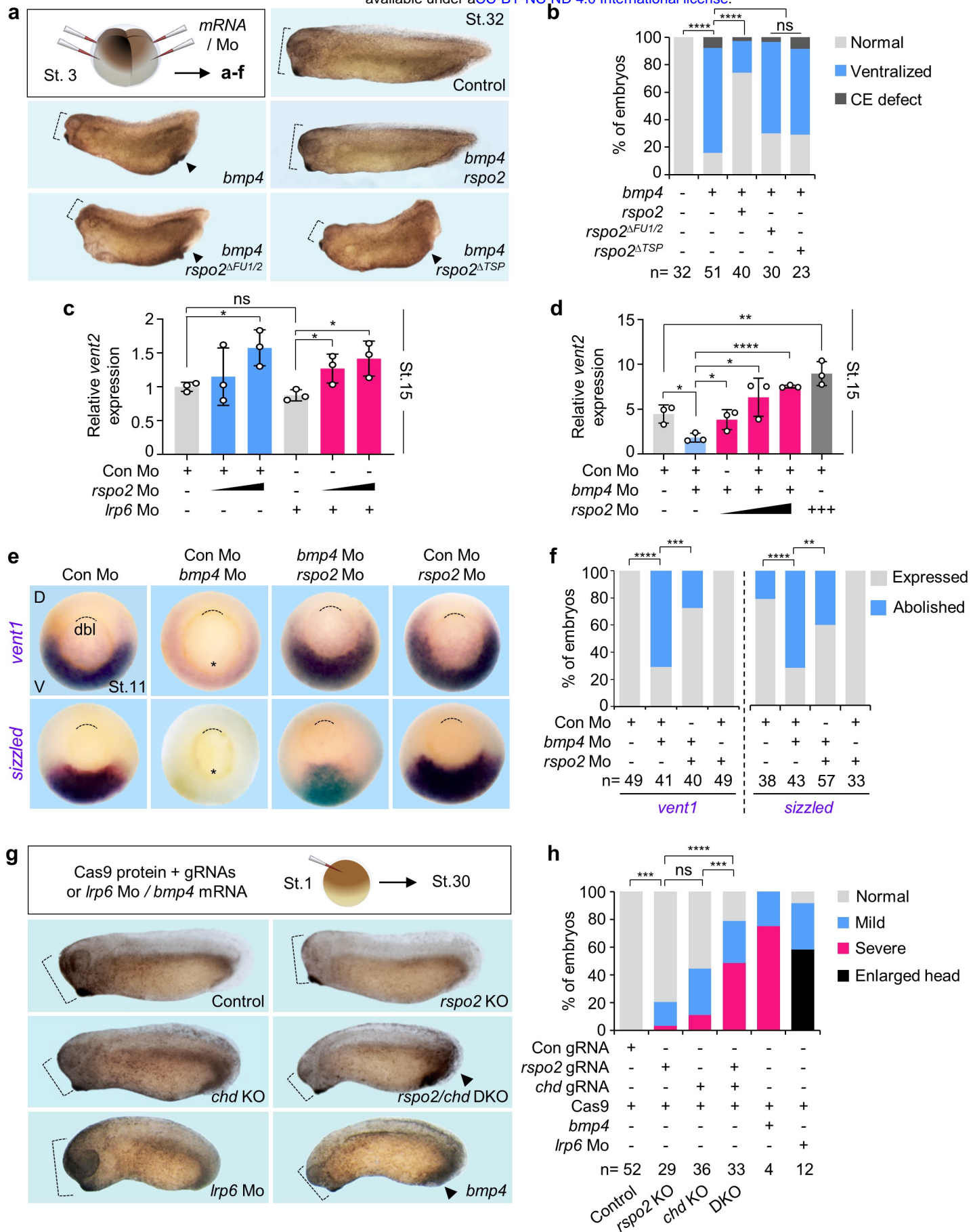


Fig. 2. *Rspo2* inhibits BMP4 signaling in *Xenopus* dorsoventral embryonic patterning.

(a) Microinjection strategy for (a-f), and representative phenotypes of *Xenopus laevis* tadpoles (St. 32) injected with the indicated mRNAs radially at 4-cell stage. Dashed lines, head size. Arrowheads, enlarged ventral structure.

(b) Quantification of embryonic phenotypes shown in (a). 'Ventralized' represents embryos with both small head and enlarged ventral structure, reminiscent of BMP hyperactivation. 'CE defect' refers to embryos with convergent extension (gastrulation) defects, unrelated to BMP signaling. Note that *rspo2* mRNA dosage used in (a) was below those that cause gastrulation defects. n, number of embryos.

(c, d) BMP-(*vent2*) reporter assays with *Xenopus laevis* neurulae (St.15) injected with reporter plasmids and the indicated Mo at 4-cell stage. Data are biological replicates and displayed as means \pm SD with unpaired t-test.

(e) *In situ* hybridization of *vent1* and *sizzled* in *Xenopus laevis* gastrulae (St.11, dorsal to the top, vegetal view) injected as indicated. D, dorsal, V, ventral. Asterisk, abolishment of the expression. Dashed line, dorsal blastopore lip (dbl).

(f) Quantification of embryonic phenotypes shown in (e). 'Expressed', normal, increased or reappearance of *vent1/sizzled* expression. 'Abolished', complete absence of *vent1/sizzled* expression. Data are pooled from two independent experiments. n, number of embryos.

(g) Microinjection strategy and representative phenotypes of *Xenopus tropicalis* tadpole (St.30) Crispants and tadpoles (St.30) injected with *bmp4* mRNA or *Irp6* Mo. At 1-cell stage, Cas9 protein with guide RNA (gRNA) targeting *rspo2* or *chd*, or both gRNAs were injected anically. Dashed lines, head size. Arrowheads, enlarged ventral structure.

(h) Quantification embryonic phenotypes shown in (g). 'Severe' showed small head, enlarged ventral tissues and short body axis. 'Mild' showed one or two of the defects described above. 'Normal' showed no visible differences to the uninjected control. The number of embryos is indicated on the bottom. Scoring of the embryos for quantification was executed with blinding from two individuals. ns, not significant. **P < 0.01, ***P < 0.001, ****P<0.0001 from χ^2 test comparing normal versus ventralized phenotypes (b) or normal versus severe and mild defects (h).

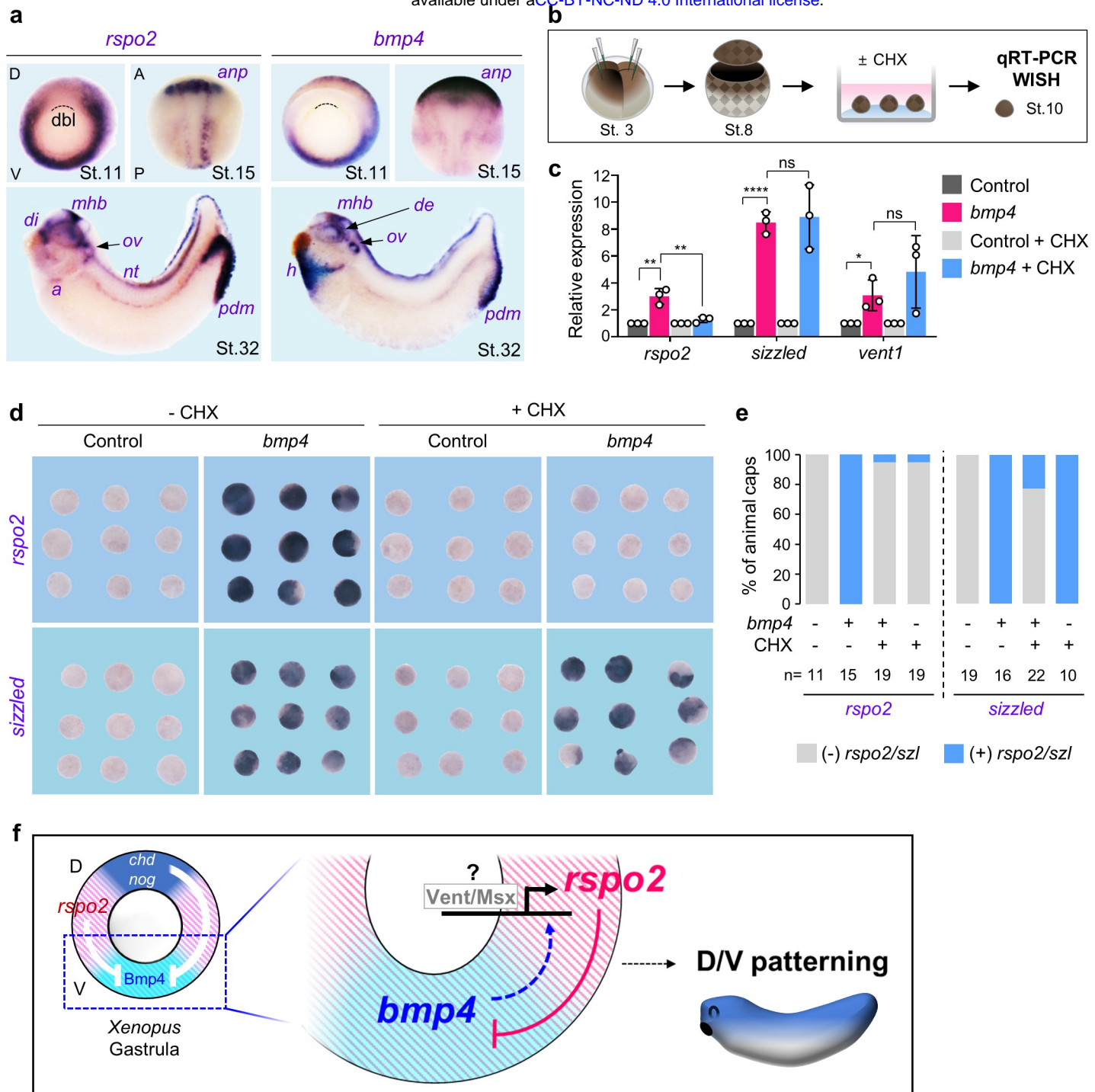


Fig. 3. Rspo2 is a negative feedback inhibitor in the BMP4 synexpression group.

(a) *In situ* hybridization of *rspo2* and *bmp4* in *Xenopus laevis* at gastrula (St. 11, dorsal to the top, vegetal view), neurula (St. 15, anterior to the top, dorsal view), and tadpole (St. 32, anterior to the left, lateral view). Dashed lines, dorsal blastopore lip (dbl); anp, anterior neural plate; di, diencephalon; mhb, mid-hindbrain boundary; ov, otic vesicle; a, atria; nt, neural tube; de, dorsal eye; h, heart; pdm, proctodeum.

(b) Microinjection and experimental scheme for (c-e). 2- or 4 cell stage *Xenopus laevis* embryos were anically injected with control (*ppl*) or *bmp4* mRNA. The animal cap (AC) explants were dissected from injected embryos at stage 8, and either treated or untreated with cycloheximide (CHX) until control embryos reached stage 10 for qRT-PCR (c) or *in situ* hybridization (d-e).

(c) qRT-PCR of *rspo2*, *sizzled*, and *vent1* expression in the AC explants injected and treated as indicated. Data are pooled from three independent experiments and displayed as means \pm SD with unpaired t-test. *P<0.05, **P < 0.01, ****P <0.0001.

(d) *In situ* hybridization of *rspo2* and *sizzled* in the AC explants injected and treated as indicated. (e) Quantification of (d). The number of the AC explants is indicated on the bottom.

(f) Model for Rspo2 function as a negative feedback inhibitor of BMP4 in *Xenopus* dorsoventral patterning.

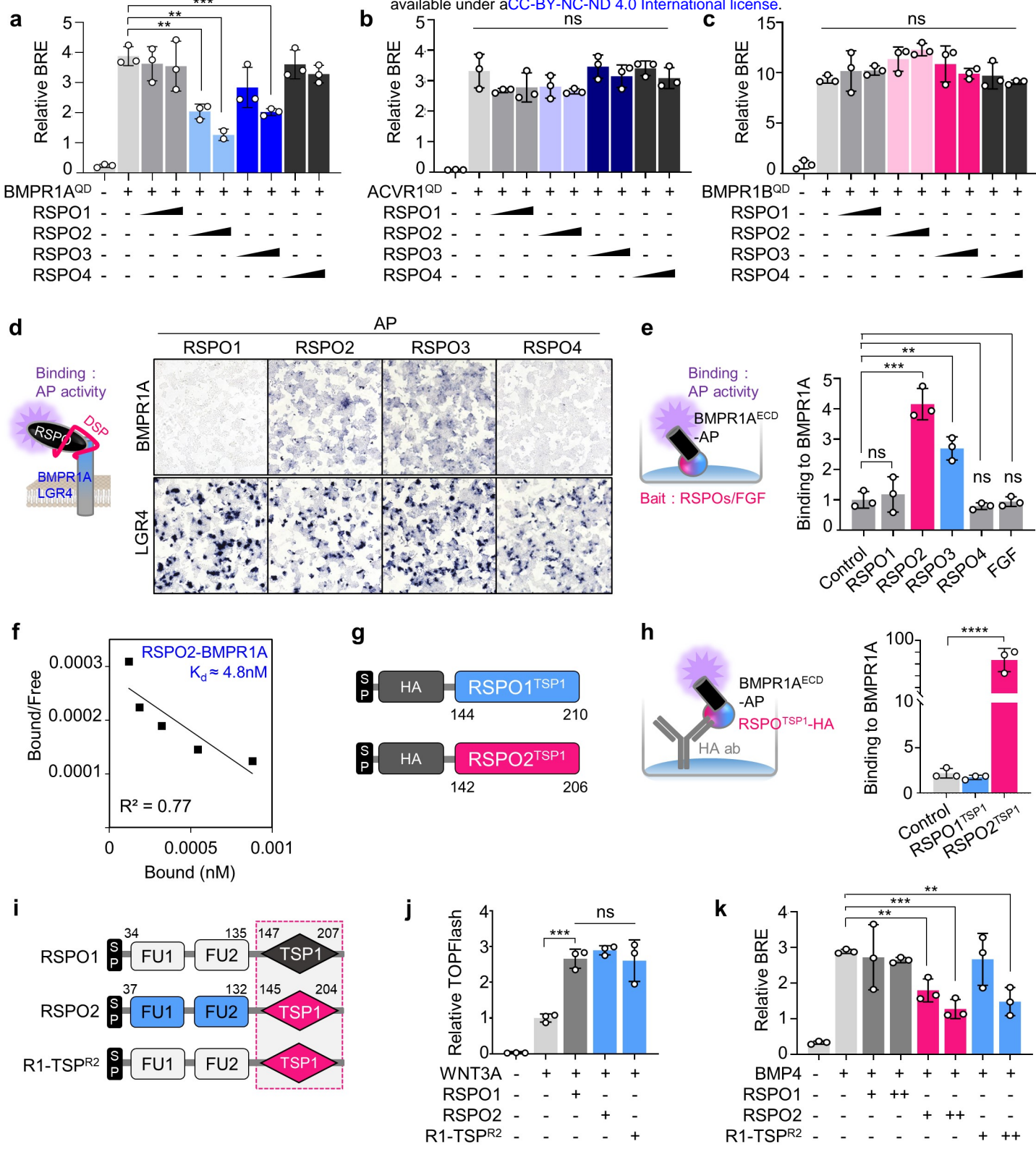


Fig. 4. RSPO2 and -3 interact with BMPR1A via the TSP1 domain.

(a-c) BRE reporter assays in HEPG2 cells transfected with constitutively active (QD) BMPR1A **(a)**, ACVR1 **(b)**, or BMPR1B **(c)** with or without BMP4 and RSPO1-4 treatment overnight.

(d) Cell surface binding assay in HEK293T cells. (Left) Scheme of the assay. Cells were transfected with BMPR1A or LGR4 DNA, and treated with same amount of RSPO1-4-AP upon DSP crosslinking as indicated. Binding was detected as purple stain on cell surface by chromogenic AP assay. (Right) Images of cells transfected and treated as indicated. Data shows a representative from four independent experiments. For quantification, see **Supplementary Fig. 5a**.

(e) *In vitro* binding assay between RSPO1-4, FGF and BMPR1A^{ECD}. (Left) Scheme of the assay. RSPOs and FGF recombinant proteins were coated on plate as baits, followed by BMPR1A^{ECD}-AP treatment overnight. (Right) Bound BMPR1A^{ECD} was detected by chromogenic AP assay. Normalized AP activity with control treatment was set to 1.

(f) Scatchard plot of RSPO2 and BMPR1A^{ECD} binding to validate K_d for RSPO2-BMPR1A.

(g) Domain structures of the RSPO1 and -2^{TSP1} with Strep-HA and flag tags used in **(h)**. SP, signal peptide; TSP1, thrombospondin domain 1.

(h) *In vitro* binding assay for RSPO^{TSP1} and BMPR1A^{ECD}. (Left) Scheme of the assay. HA-harboring RSPO1/2^{TSP1} were captured to HA antibody coated plate, and BMPR1A^{ECD}-AP was treated overnight. (Right) Bound BMPR1A to RSPO^{TSP1} was detected with absorbance.

(i) Domain structures of the RSPO1, -2 and R1-TSP^{R2}. SP, signal peptide; FU, furin domain; TSP1, thrombospondin domain 1. Dashed box indicates the TSP1 domain swapping.

(j) TOPflash reporter assay in HEPG2 cells upon WNT3A with or without **(i)** as indicated.

(k) BRE reporter assay in HEPG2 cells upon BMP4 with or without **(i)** as indicated. Data for reporter assays **(a-c, j, k)** are biological replicates; *In vitro* binding assays **(e, h)** are experimental replicates and displayed as mean \pm SD; ns, not significant, *P < 0.05, **P < 0.01, ***P < 0.001, ****P < 0.0001 from unpaired t-test.

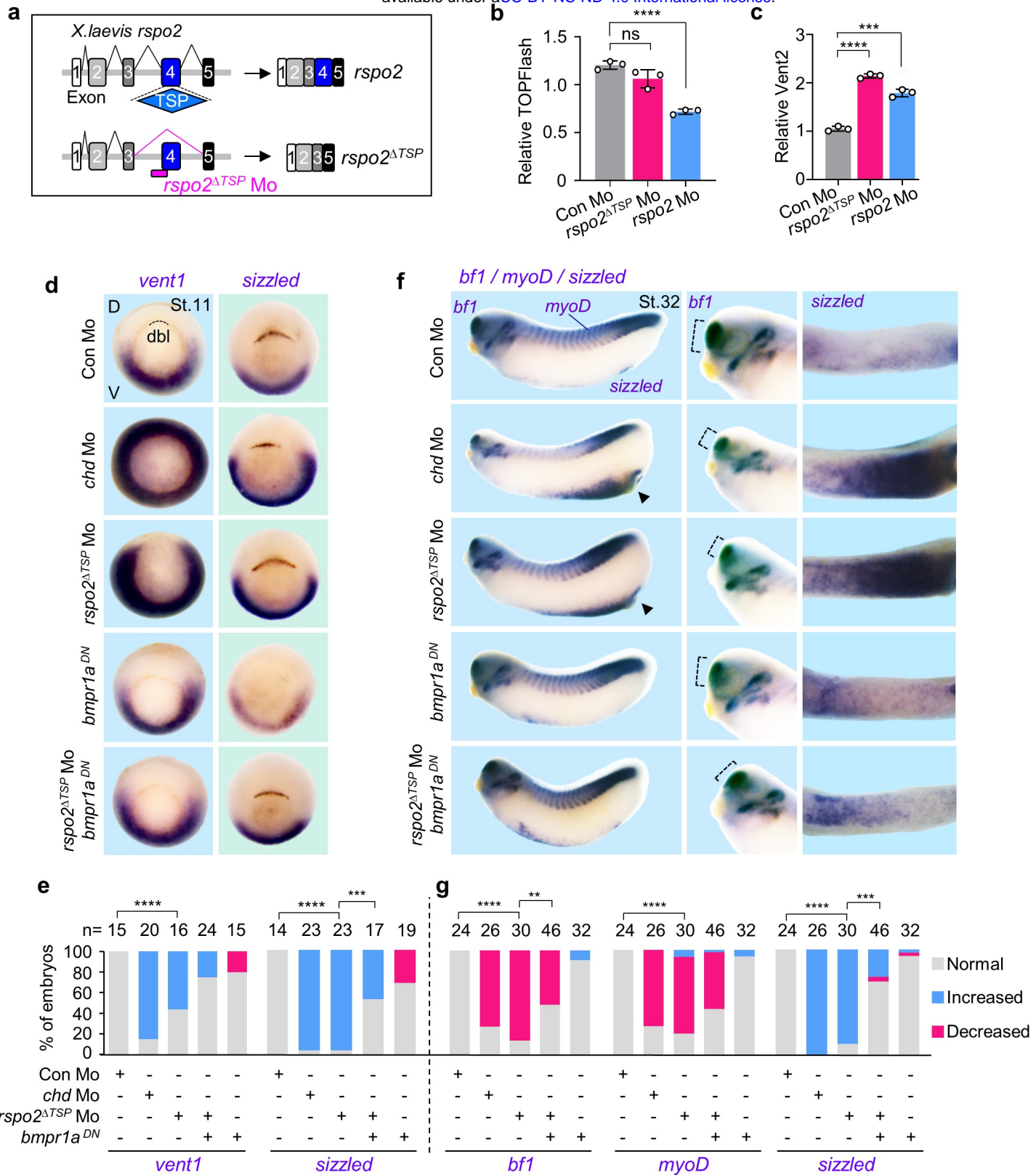


Fig. 5. Loss of Rspo2-TSP1 domain activates BMP signaling in *Xenopus* development.

(a) Scheme for *rspo2*^{ΔTSP} splicing Mo in *Xenopus laevis*.

(b) TOPFlash assay and (c) BMP-reporter (*vent2*) assay in *Xenopus laevis* neurulae (St.15) injected radially at 4-cell stage with reporter plasmids and Mo as indicated. Data are biological replicates, and displayed as mean \pm SD; ns, not significant, ***P < 0.001, ****P<0.0001 from unpaired t-test.

(d-g) *In situ* hybridization of BMP4 targets *vent1* and *sizzled* in *Xenopus laevis*. Embryos were injected radially and equatorially at 4-cell stage as indicated. Gastrulae (St.11) (d) and quantification (e); Tadpoles (St. 32) (f) and quantification (g). Dashed lines, dorsal blastopore lip (dbl) (d) or *bf1* expression (e); D, dorsal; V, ventral. For (f), left, lateral view; middle, magnified view of head; right, magnified view of ventral side. 'Increased/Decreased' represents embryos with significant expansion/reduction of *sizzled* or *vent1* signals toward the dorsal/ventral side of the embryo (e), or with significant increase/decrease of the signal strength (g). ns, not significant. n, number of embryos. Scoring of the embryos for quantification was executed with blinding from two individuals. ns, not significant; **P<0.01, ***P<0.001, ****P<0.0001 from χ^2 test comparing normal versus increased. Data are pooled from at least two independent experiments.

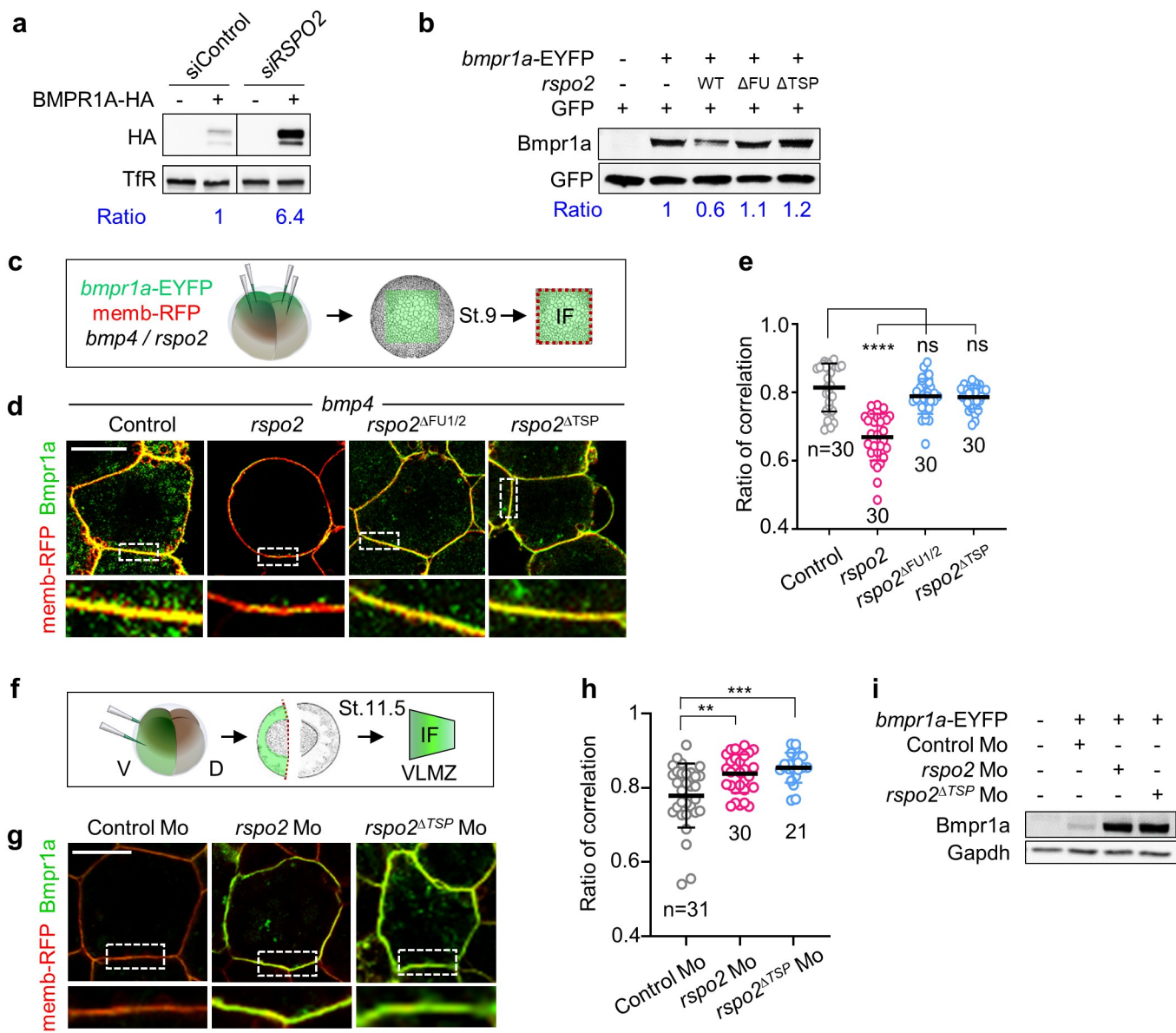


Fig. 6. RSPO2 removes cell surface BMPR1A.

(a) Western blot analysis in H1581 cells treated with siControl or siRSPO2 as indicated and transfected with or without BMPR1A-HA DNA. Transferrin receptor (TfR), a loading control. Ratio, relative levels of BMPR1A-HA normalized to TfR.

(b) Western blot analysis of Bmpr1a in *Xenopus laevis* (St. 15) neurulae injected anally at 2- to 4-cell stages as indicated. GFP mRNA was injected as an injection control. Ratio, relative levels of Bmpr1a normalized to GFP.

(c) Scheme for immunofluorescence microscopy (IF) in *Xenopus laevis* animal cap (AC) explants. Embryos were injected anally at 4-cell stage with *bmpr1a*-EYFP and memb-RFP mRNA along with *bmp4* and *rspo2* wild-type or mutant mRNA. AC explants were dissected at St.9 for IF. Membrane (memb)-RFP was used as a control comparing relative change of Bmpr1a-EYFP signal at cell surface.

(d) IF for Bmpr1a (green) and cell membrane (red) in AC explants injected as indicated, with a representative cell (top) and magnification (inset). Scale bar, 20 μ m. (e) Quantification of (d).

(f) Scheme for IF in *Xenopus laevis* ventrolateral marginal zone explants (VLMZ). Embryos were ventrally injected at 4-cell stage with *bmpr1a*-EYFP and memb-RFP mRNA with Mo. VLMZs were dissected at stage 11.5 for IF.

(g) IF for Bmpr1a (green) and cell membrane (memb-RFP, red) in VLMZ injected with mRNA and Mo as indicated. Scale bar, 20 μ m. (h) Quantification of (g).

(i) Western blot analysis of Bmpr1a in *Xenopus laevis* neurulae (St. 18) injected radially at 4-cell stage as indicated. Gapdh, a loading control.

Data are the number of areas analyzed (e, h) and displayed as mean \pm SD. ns, not significant; **P < 0.01, ***P < 0.001, ****P < 0.0001 from unpaired t-test. For the uncropped western blot images, see **Source file**.

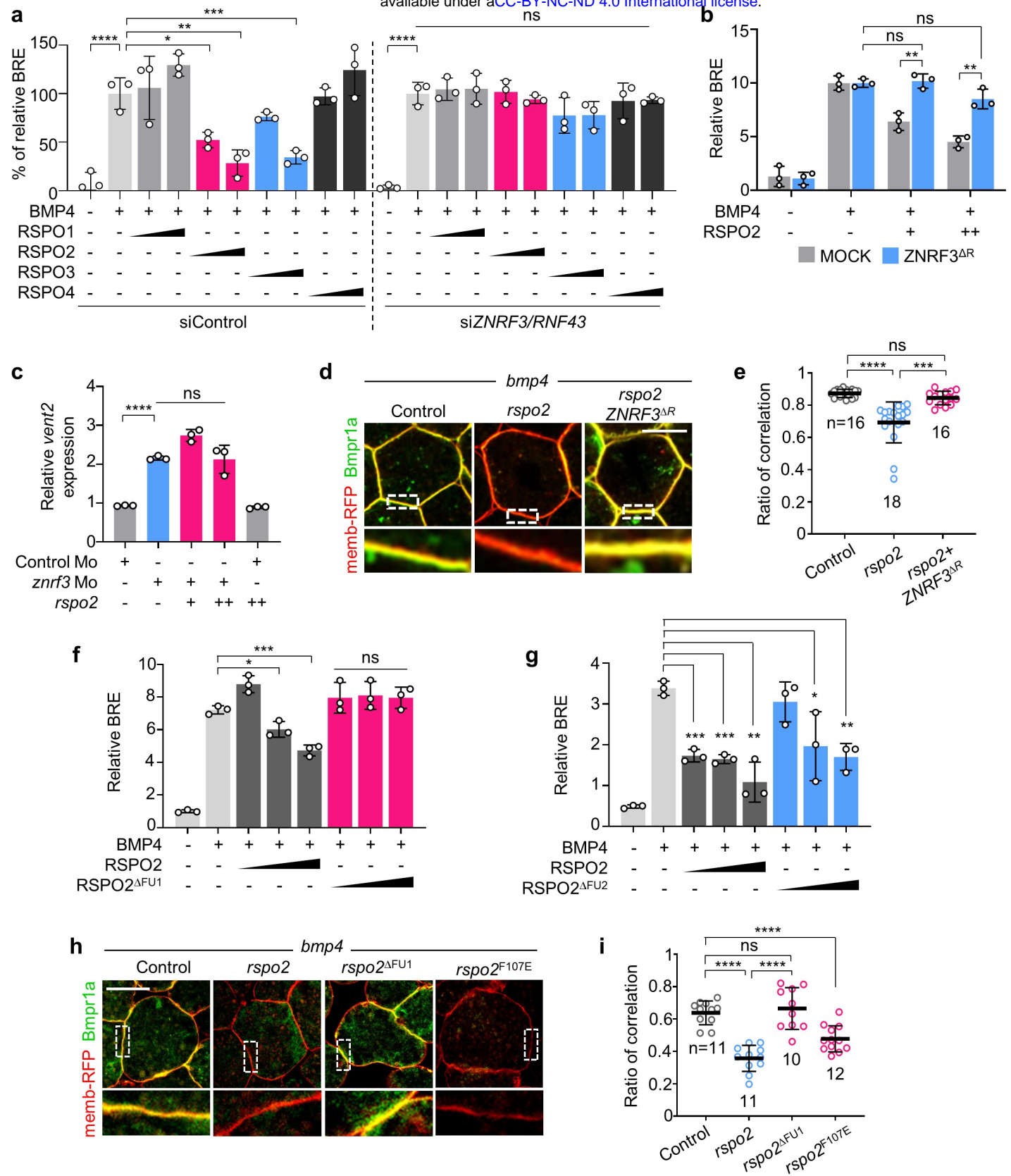


Fig. 7. RSPO2 requires ZNRF3 to antagonize BMP4-BMPR1A signaling.

- (a) BRE reporter assay in HEPG2 cells. Cells were transfected with siControl or siZNRF3/siRN43, and BMP4 with or without RSPO1-4 were added overnight as indicated. Normalized BRE activity upon BMP4 without RSPO2 stimulation was set to 100 %.
- (b) BRE reporter assay in HEPG2 cells upon ZNRF3^{ΔR} transfection, with or without overnight BMP4 and RSPO2 treatment as indicated.
- (c) BMP-reporter (*vent2*) assay in *Xenopus laevis* St.15 neurulae. Embryos were injected anically with reporter plasmids and the indicated Mo with or without *rspo2* mRNA at 4-cell stage. Normalized *vent2* activity of control Mo injected embryos with reporter plasmids was set to 1.
- (d) Immunofluorescence microscopy (IF) in *Xenopus laevis* animal cap explants for Bmpr1a (green) and the plasma membrane (red) from embryos injected with mRNA as indicated, with a representative cell (top) and magnification (inset). Scale bar, 20 μm. For scheme, see **Fig. 6c**.
- (e) Quantification of (d).
- (f-g) BRE reporter assay in HEPG2 cells treated with BMP4 and RSPO2/RSPO2^{ΔFU1}/RSPO2^{ΔFU2} overnight as indicated. For domain structure of RSPO2^{ΔFU1}/RSPO2^{ΔFU2}, see **Supplementary Fig. 8a**.
- (h) IF for Bmpr1a (green) and plasma membrane (red) in animal cap explants injected as indicated, with a representative cell (top) and magnification (inset) showing the plasma membrane. Scale bar, 20 μm. For domain structure of *Xenopus* Rspo2 mutants, see **Supplementary Fig. 8d**. (i) Quantification of (h).

Data are the number of areas analyzed (e, i) or biological replicates (a, b, c, f, g) and displayed as mean ± SD. ns, not significant; *P < 0.05, **P < 0.01, ***P < 0.001, ****P < 0.0001 from unpaired t-test. For the uncropped western blot images, see **Source file**.

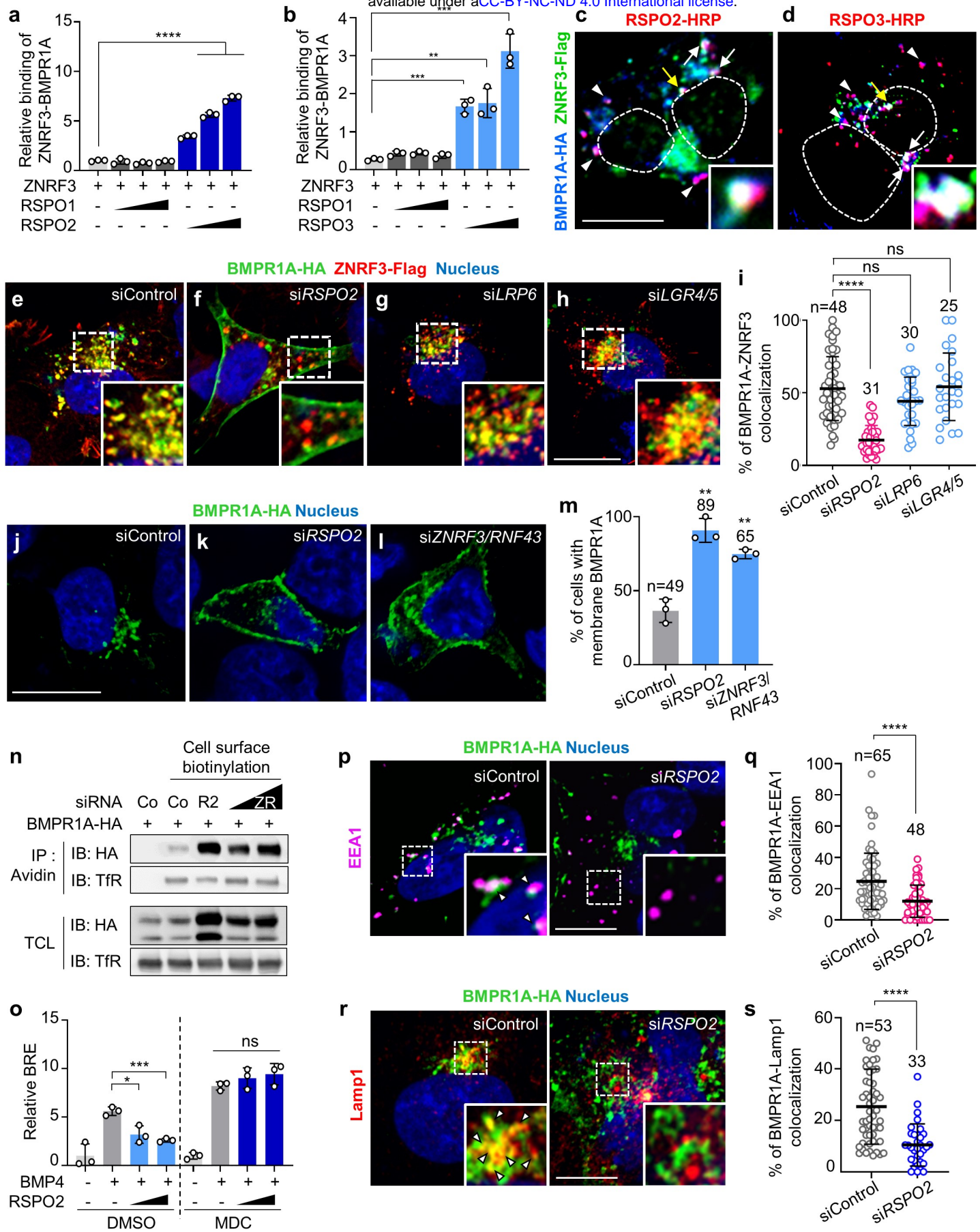


Fig. 8. RSPO2 bridges BMPR1A and ZNRF3 and triggers BMP receptor clearance from the cell surface.

- (a, b)** *In vitro* binding assay between ZNRF3 and BMPR1A^{ECD} mediated by RSPO1-3. ZNRF3-Fc protein was used as a bait, with sequential RSPO1-3 protein and BMPR1A^{ECD}-AP treatment. Bound BMPR1A^{ECD} to ZNRF3 was detected by chromogenic AP assay.
- (c, d)** IF in H1581 cells transfected with BMPR1A-HA and ZNRF3-flag DNA upon RSPO2 and -3-HRP treatment for 3 h. RSPOs (red) were visualized with tyramid signal amplification. BMPR1A (blue) and ZNRF3 (green) were stained against HA and flag antibody. White arrowheads, colocalized BMPR1A/RSPO2; white arrows, colocalized BMPR1A/RSPO2-3/ZNRF3; yellow arrow, colocalized BMPR1A/RSPO2-3/ZNRF3 in magnified inset; Dashed lines, nucleus. Scale bar, 20 μ m.
- (e-h)** IF of colocalized BMPR1A (green)/ZNRF3 (red) in H1581 cells treated with siRNA as indicated. Nuclei were stained with Hoechst. Scale bar, 20 μ m. **(i)** Quantification of BMPR1A colocalizing with ZNRF3 from **(e-h)**.
- (j-l)** IF of BMPR1A (green) in H1581 cells treated with siRNA as indicated. **(m)** Quantification of cells harboring membrane localized BMPR1A from **(j-l)**.
- (n)** Cell surface biotinylation assay in H1581 cells treated with BMPR1A-HA and siRNA as indicated. Co, control; R2, *RSPO2*; ZR, *ZNRF3/RNF43* siRNA. After labeling surface proteins with Biotin, lysates were pulled down with streptavidin beads and subjected to Western blot analysis. Transferrin receptor (TfR), a loading control. TCL, Total cell lysate. Data shows representative result from three independent experiments.
- (o)** BRE reporter assay in HEPG2 cells treated as indicated. MDC, monodansylcadaverin.
- (p)** IF of colocalized BMPR1A (green) / EEA1 (magenta) in H1581 cells treated with siRNA. White arrowheads, colocalized BMPR1A/EEA1 in magnified inset. **(q)** Quantification of **(p)**.
- (r)** IF of colocalized BMPR1A (green) / Lamp1 (red) in H1581 cells treated with siRNA as indicated. White arrowheads, colocalized BMPR1A/Lamp1 in magnified inset.
- (s)** Quantification of **(r)**.

Data for binding assays **(a,b)** are experimental replicates; IF **(i, m, q, s)** are the number of cells pooled from at least two independent experiments, and displayed as mean \pm SD. ns, not significant, *P < 0.05, **P < 0.01, ***P < 0.001, ****P < 0.0001 from unpaired t-test.