The phytohormone jasmonate and its metabolites regulate a wide spectrum of plant physiology, participating in normal development and growth processes, as well as defence responses to environmental and pathogenic stressors. Jasmonate is activated upon specific conjugation to the amino acid L-isoleucine (Ile), which produces the highly bioactive hormone (3R,7S)-jasmonoyl-L-isoleucine (JA-Ile) that is functionally and structurally mimicked by the Pseudomonas syringae phytopathogen coronatine.

The discovery of coronatine-insensitive mutants enabled the identification of COI1 as a key player in the jasmonate pathway, with further implications of regulated protein turnover of the AUX/IAA transcriptional repressors. Supported by its sequence homology and functional similarity to TIR1, COI1 has been assigned a critical role in the direct perception of the jasmonate signal.

Despite the importance of jasmonate signalling in plant physiology, the molecular mechanism of jasmonate perception remains elusive. Here we present crystal structures of COI1 bound to JA-Ile or coronatine, as well as peptides of a bipartite JAZ1 degron. Our structural and pharmacological studies reveal that the true jasmonate receptor is a co-receptor complex, consisting of the F-box protein COI1, the JAZ degron and a newly discovered third component, inositol pentakisphosphate.
dependent COI1–JAZ interaction. Consistent with these findings, truncation of the PY motif in JAZ1 has little effect on the ligand-binding activity (Supplementary Fig. 1c).

We have previously mapped the COI1-binding region of the JAZ proteins to the carboxy-terminal Jas motif, which is characterized by the SLX,FX,KRX,RX,PY consensus sequence preceded by two consecutive basic residues. A single Ala mutation of the central strictly conserved phenylalanine residue in the Jas motif is sufficient to abolish the formation of the high-affinity jasmonate co-receptor (Fig. 1a). Previous studies showed that the highly conserved PY sequence at the C terminus of the Jas motif has a role in JAZ co-receptor (Fig. 1a). Previous studies showed that the highly conserved central and C-terminal region and a more variable N-terminal region, to interact with COI1 and perceive the jasmonate signal. Consistent with our in vitro ligand-binding data, the minimal sequence in JAZ1 is sufficient for coronatine-induced COI1–JAZ1 interaction (Supplementary Fig. 1d). Therefore, we conclude that the interactions among COI1, coronatine, and the JAZ1 peptide are highly cooperative and that the short Glu 200–Val 220 sequence functions as the JAZ1 degron.

### Jasmonate-binding pocket on COI1

To elucidate the structural mechanism by which the COI1–JAZ1 co-receptor senses jasmonate, we crystallized and determined the structures of the COI1–ASK1–JAZ1 degron peptide complex together with either (3R,7S)-JA-Ile or coronatine (Supplementary Table 1). The crystal structure of COI1 reveals a TIR1-like overall architecture, with an N-terminal tri-helical F-box motif bound to ASK1 and a C-terminal horseshoe-shaped solenoid domain formed by 18 tandem leucine-rich repeats (LRRs; Fig. 2a, b). Similar to TIR1, the top surface of the COI1 LRR domain has three long intra-repeat loops (loop-2, loop-12 and loop-14) that are involved in hormone and polypeptide substrate binding. Unlike TIR1, however, a fourth long loop (loop-C) in the C-terminal capping sequence of the COI1 LRR domain folds over loop-2, partially covering it from above (Fig. 2b, c).

Despite their similar overall fold, COI1 has evolved a hormone-binding site that is distinct from TIR1. Configured in between loop-2 and the inner wall of the LRR solenoid, the ligand-binding pocket of COI1 is exclusively encircled by amino acid side chains (Fig. 2d–f). Many of the pocket-forming residues on COI1 are large in size and carry a polar head group (Supplementary Fig. 2). These properties allow them to mould a binding pocket into a specific shape while forming close interactions with each chemical moiety of the ligand. These close interactions are critical to proper hormone sensing of the complex—in yeast two-hybrid assays, mutation of any of these large side-chain amino acids on COI1 is sufficient to disrupt the interaction of COI1 with JAZ1 in the presence of coronatine (Supplementary Fig. 3).

In the binding pocket, both JA-Ile and coronatine sit in an ‘upright’ position with the keto group of their common cyclopentanone ring pointing up and forming a triangular hydrogen bond network with Arg 496 and Tyr 444 of COI1 at the pocket entrance (Fig. 2d–f). Without the JAZ degron peptide bound, the keto group of the ligand is accessible to solvent (Fig. 2g). The rest of the cyclopentanone ring of both JA-Ile and coronatine is sandwiched between the aromatic groups of Phe 89 and Tyr 444 of COI1, stabilized by hydrophobic packing. The cyclohexene ring of coronatine provides a rigid surface area for close packing with Phe 89, whereas the more flexible and extended pentenyl side chain of JA-Ile is more loosely accommodated by a hydrophobic pocket formed by Ala 86, Phe 89 and Leu 91 from loop-2 as well as Leu 469 and Trp 519 from the LRRs (Supplementary Fig. 4a). Differences at this interface probably explain the approximately tenfold higher affinity of coronatine over (3R,7S)-JA-Ile, as detected in our binding assays.
Deeper in the ligand-binding pocket, the common amide and carboxyl groups of JA-Ile and coronatine bind to the bottom of the binding site by forming a salt bridge and hydrogen bond network with three basic residues of COI1: Arg85, Arg348 and Arg409 (Fig. 2d, e). Together, these arginine residues constitute the charged floor of the ligand pocket. Tyr386 reinforces the interactions from above by making a hydrogen bond through its backbone amide group to the keto moiety of the ligand. In doing so, Tyr386 approaches the cyclopentanone ring of the ligand, narrowing the pocket entrance, and creating a hydrophobic cave below. The rest of the basin is carved out by Val411, Ala384 and the aliphatic side chain of Arg409 (Supplementary Fig. 3). In agreement with its important role in forming the JA-Ile co-receptor, this short N-terminal region of the JAZ1 degron completely covers the opening of the ligand-binding pocket, conferring high-affinity binding to the hormone. The close interaction between the hormone and the co-receptor complex provides a plausible structural explanation for the favourable binding of the (S,7S)-JA-Ile isomer, as the stereochemistry at the 7 position of (3R,7R)-JA-Ile may through a hydrogen bond formed between the backbone carbonyl of Pro202 in JAZ1 and the ligand-interacting COI1 residue Arg496, which is critical for the hormone-dependent COI1–JAZ interaction (Supplementary Fig. 3). In agreement with its important role in forming the JA-Ile co-receptor, this short N-terminal region of the JAZ1 degron completely covers the opening of the ligand-binding pocket, conferring high-affinity binding to the hormone.

Structural roles of the bipartite JAZ degron

The JAZ1 degron peptide adopts a bipartite structure with a loop region followed by an α-helix to assemble with the COI1–jasmonate complex. The hallmark of the JAZ1 degron is the N-terminal five amino acids identified in the radioligand binding assay. In a largely extended conformation, this short sequence lies on top of the hormone-binding pocket and simultaneously interacts with both COI1 and the ligand, effectively trapping the ligand in the pocket (Fig. 3a, b). At the N-terminal end, Leu201 of the JAZ1 peptide is embedded in a hydrophobic cavity presented by surface loops on top of COI1 (Fig. 3c). At the C-terminal end, Ala204 of JAZ1 uses its short side chain to pack against the keto group of the ligand and Phe89 of COI1 (Fig. 3c and Supplementary Fig. 4a). The same alanine residue of JAZ1 also donates a hydrogen bond through its backbone amide group to the keto moiety of the ligand emerging from the pocket (Fig. 3c). The middle region of the five-amino-acid sequence is secured to the COI1–jasmonate complex through hydrogen bonding. The JAZ1 degron peptide adopts a bipartite structure with a loop region followed by an α-helix to assemble with the COI1–jasmonate complex. The hallmark of the JAZ1 degron is the N-terminal five amino acids identified in the radioligand binding assay. In a largely extended conformation, this short sequence lies on top of the hormone-binding pocket and simultaneously interacts with both COI1 and the ligand, effectively trapping the ligand in the pocket (Fig. 3a, b). At the N-terminal end, Leu201 of the JAZ1 peptide is embedded in a hydrophobic cavity presented by surface loops on top of COI1 (Fig. 3c). At the C-terminal end, Ala204 of JAZ1 uses its short side chain to pack against the keto group of the ligand and Phe89 of COI1 (Fig. 3c and Supplementary Fig. 4a). The same alanine residue of JAZ1 also donates a hydrogen bond through its backbone amide group to the keto moiety of the ligand emerging from the pocket (Fig. 3c). The middle region of the five-amino-acid sequence is secured to the COI1–jasmonate complex through hydrogen bonding.
place the aliphatic chain unfavourably close to nearby JAZ1 and COI1 residues (Supplementary Fig. 4a).

Within the JAZ1 degron, two conserved basic residues, Arg 205 and Arg 206, were previously shown to have an important role in hormone-induced COI1 binding\(^1\). In the structure, Arg 205 contributes to COI1 binding by directly interacting with loop-12, whereas Arg 206 points in the opposite direction and inserts deeply into the central tunnel of the COI1 solenoid. Approaching the bottom of the ligand-binding pocket, the guanidinium group of the Arg 206 side chain joins the three basic COI1 residues that form the pocket floor and interacts directly with the carboxyl group of the ligand (Fig. 3d). Thus, the N-terminal seven amino acids (ELPIARR) of the JAZ1 degron peptide act as a clamp that wraps the ligand-binding pocket from top to bottom, closing it completely (Fig. 3b).

The highly conserved C-terminal half of the JAZ1 degron forms an amphipathic \(\alpha\)-helix that strengthens the JAZ1–COI1 interaction by binding to the top surface of the COI1 LRR domain, adjacent to the ligand-binding site (Fig. 3a). With its N-terminal end directly packing against loop-2 of COI1, the Jas motif helix blocks the central tunnel of the COI1 LRR solenoid like a plug. The N-terminal half of the Jas motif helix is characterized by three hydrophobic residues—Leu 209, Phe 212 and Leu 213—which are aligned on the same side of the helix and form a hydrophobic interface with COI1 (Fig. 3e). By soaking the COI1–ASK1 crystals with coronatine and a sufficiently high concentration of JAZ1 degron peptide lacking the N-terminal ELPIA sequence, we were able to trap a complex formed by COI1, coronatine and the isolated Jas motif helix in the crystal (Supplementary Table 1). This indicates that the \(\alpha\)-helix may provide a low-affinity anchor for docking the JAZ protein on COI1. In support of this idea, single-amino-acid mutations at the complementary surface on COI1 readily disrupt hormone-induced COI1–JAZ1 interaction (Fig. 3f).

**Inositol pentakisphosphate as a cofactor of COI1**

The crystal structure of TIR1 revealed an unexpected inositol hexakisphosphate (InsP\(_6\)) molecule bound in the centre of the protein underneath the auxin-binding pocket\(^2\). The sequence homology between COI1 and TIR1 suggests that COI1 might also contain a similar small molecule. Before crystallization, we analysed the recombinant COI1–ASK1 complex by structural mass spectrometry. Nano-electrospray mass spectra of the intact COI1–ASK1 complex revealed two populations differing by a mass of \(\sim 568\) Da, indicating that a small molecule was indeed co-purified with the proteins (Fig. 4a and Supplementary Fig. 5). The mass-spectrometry-derived molecular mass of the unknown compound is different from the mass of InsP\(_5\) (651 Da) but matches that of an inositol pentakisphosphate (Ins(1,2,4,5,6)P\(_5\); Fig. 4c)\(^2\). This conclusion was further supported by the TOCSY spectrum of synthetic Ins(1,2,4,5,6)P\(_5\) (Fig. 4d) and the subsequently acquired negative-ion electrospray ionization mass spectrometry spectrum of the compound (Supplementary Fig. 6).

Consistent with the binding of a small molecule cofactor, the crystal structure of COI1 showed strong unexplained electron densities clustered in the middle of the COI1 LRR domain. Like InsP\(_5\) in TIR1, these extra densities in COI1 are located directly adjacent to the bottom of the ligand-binding pocket of the jasmonate co-receptor, interacting with multiple positively charged COI1 residues (Fig. 4e). Unexpectedly, these islands of electron density cannot be explained by an Ins(1,2,4,5,6)P\(_5\) molecule.
Instead, their intensity, overall symmetry and poor connectivity indicate that they belong to multiple free phosphate molecules. Because a high concentration of ammonium phosphate was used as the major precipitant for crystallizing the jasmonate co-receptor, we postulate that the InsP₅ molecules that co-purified with COI1 was later displaced by phosphate molecules in the crystallization drops. In support of this scenario, the concave surface of the COI1 solenoid fold surrounding the phosphates is highly basic and decorated with residues conserved in plant COI1 orthologues, indicating a functionally important surface area (Fig. 4f and Supplementary Figs 2 and 7).

**InsP₅ potentiates jasmonate perception by COI1–JAZ1**

The highly selective co-purification of two different inositol phosphates, InsP₅ and InsP₆, with two homologous plant hormone receptors, COI1 and TIR1, implies that the proper function of the two F-box proteins might require the binding of specific inositol phosphates. To assess the functional role of Ins(1,2,4,5,6)P₅ in the COI1–JAZ1 co-receptor, we took advantage of our crystallographic observation and developed a protocol to strip the co-purified InsP₅ from COI1 without denaturing the protein. The resulting COI1–ASK1 complex was then tested in a ligand-binding-based reconstitution assay. As shown in Fig. 5a, untreated COI1 formed a high-affinity jasmonate co-receptor with JAZ1. Addition of exogenous Ins(1,2,4,5,6)P₅ did not significantly change its activity. In contrast, the dialysed COI1 sample completely lacked ligand binding by itself and showed only trace activity in the presence of JAZ1. Supplementation with either synthetic Ins(1,2,4,5,6)P₅ (Fig. 5b) or the purified and NMR-analysed InsP₅ sample (data not shown) rescued the interaction in a dose-dependent manner and with a half-maximum effective concentration (EC₅₀) of 27 nM (Fig. 5c). From this reconstitution result, we conclude that Ins(1,2,4,5,6)P₅ binding is crucial for the jasmonate co-receptor to perceive the hormone with high sensitivity.

![Figure 5](image)

**Discussion**

Our structural and pharmacological analyses reveal not only the essential components of the receptor system but also the detailed mechanism by which these components cooperatively assemble and recognize the hormonal signal through a network of interactions. Our data identify the true jasmonate receptor as a three-molecule co-receptor complex, consisting of COI1, JA degron and inositol pentakisphosphate, all of which are indispensable for high-affinity hormone binding. Our analyses also define the JAZ degron boundaries as a unique bi-partite sequence that binds COI1 and directly participates in hormone recognition. Unexpectedly, the N-terminal clamp region of the JAZ1 degron that is critical for hormone binding is diverse among JAZ proteins. This variable sequence might create a family of COI1–JAZ co-receptors that respond differentially to the hormone.

The crystal structure of the COI1–JAZ co-receptor in complex with JA-Ile revealed a markedly different binding mode of the hormone as predicted by computational modelling. Although COI1 shares high sequence homology with TIR1, subtle structural differences and the integration of two additional factors critical for ligand binding give rise to a hormone-binding pocket in COI1 that is challenging to model. For the same reason, the structural nature of the ligand-free form of the F-box protein cannot be modelled with accuracy. The direct interactions of the hormone with both COI1 and the JAZ protein as observed in the crystal nonetheless support a molecular glue mechanism previously proposed for the auxin system.

Discovery of the inositol pentakisphosphate cofactor of COI1 has important implications for the role of inositol phosphates in plant hormone signalling. COI1 co-purifies with a single isoform of...
InsP_3. Ins(1,2,4,5,6)P_5 indicating selectivity at the receptor level. However, both inositol-1,2,4,5,6-pentakisphosphate and inositol-1,4,5,6-tetrakisphosphate support high-affinity hormone binding in our reconstitution assays, leaving the identity of the physiologically relevant form of inositol phosphate an open question.

Finally, our study is the latest in a series of receptor structures for plant hormones, including auxin20, gibberellins21,22 and abscisic acid24–28. Despite different structural mechanisms, a common theme of hormone-mediated protein interactions emerges as a unique strategy favoured by plant systems throughout evolution.

**METHODS SUMMARY**

The Methods provides detailed information about all experimental procedures, including: (1) description of protein preparation, purification and mutagenesis; (2) description of protein crystallization, data collection and structure determination; (3) details for conducting in vitro radioligand binding assay; (4) details for conducting yeast-two-hybrid assay; (5) description of inositol phosphate purification scheme; (6) details for conducting in vitro inositol phosphate reconstitution assays; (7) description of structural mass spectrometry analysis of the intact protein complex; (8) description of NMR analysis of the inositol phosphate; and (9) description of mass spectrometry analysis of the inositol phosphate.

**Full Methods** and any associated references are available in the online version of the paper at www.nature.com/nature.

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**Author Contributions** L.B.S., G.A.H. and N.Z. conceived and L.B.S. conducted and interpreted the structural mass spectrometry experiments. L.B.S., X.T. and N.Z. analysed crystallographic data. J.W. and S.Y.H. purified the COI1–ASK1 complex and conducted crystallographic experiments. X.T. crystallized and determined the structures of the COI1–ASK1–JAZ1 hormone complexes. L.B.S. and N.Z. analysed structural mass spectrometry experiments. L.B.S., H.M. and M.S. conducted and interpreted the structural mass spectrometry experiments. L.B.S., H.M., T.R.H., F.-F.H. and J.R. conceived and conducted experiments for inositol phosphate purification and identification. Y.K. synthesized jasmonate stereoisomers. E.M. and L.B.S. undertook FRET assays. J.R., S.Y.H. and G.A.H. conceived and conducted experiments for materials should be addressed to N.Z. (nzheng@u.washington.edu).

**Author Information** Structural coordinates and structural factors have been deposited in the Protein Data Bank under accession numbers 300K, 300L and 300M. Reprints and permissions information is available at www.nature.com/reprints. The authors declare no competing financial interests. Readers are welcome to comment on the online version of this article at www.nature.com/nature. Correspondence and requests for materials should be addressed to N.Z. (nzheng@u.washington.edu).
METHODS
Protein preparation. The full-length Arabidopsis thaliana COI1 and ASK1 were co-expressed as a glutathione S-transferase (GST) fusion protein and an untagged protein, respectively, in Hi5 suspension insect cells. The COI1–ASK1 complex was isolated from the soluble cell lysate by glutathione affinity chromatography. After on-column tag cleavage by tobacco etch virus protease, the complex was further purified by anion exchange and gel filtration chromatography and concentrated by ultrafiltration to 12–18 mg ml⁻¹. Full-length JAZ substrate proteins were expressed as 6×His-fusion proteins in Escherichia coli and purified on Ni-NTA resin with subsequent dialysis into 20 mM Tris-HCl, pH 8.0, 200 mM NaCl and 10% glycerol. For truncation mutants, a stop codon was introduced in JAZ1 proteins using the Quick-Change II site-directed mutagenesis kit (Stratagen). Synthetic JAZ degron peptides were prepared by United Biochemical Research, Inc. JAZ degron fusion peptides were prepared with N-terminal 6×His tag and C-terminal GST fusion tag and expressed in E. coli. The protein was isolated by glutathione affinity resin for pull-down assay with untagged COI1–ASK1 complex.

Site-directed mutagenesis. Individual amino acid residues in the LRR domain of COI1 proteins were mutated to alanine using the Quick-Change II site-directed mutagenesis kit (Stratagen). Mutant proteins were co-expressed with JAZ1 (JAZ1:pB42AD) in yeast to detect protein–protein interactions.

Crystallography, data collection and structure determination. The crystals of the COI1–ASK1–JAZ1 peptide complexes bound to either caffeine or JA were grown at 4°C by the hanging-drop vapour diffusion method with 1.5 mM protein complex samples containing COI1–ASK1, JAZ1 peptide and hormone compound at a 1:1:1 molar ratio mixed with an equal volume of reservoir solution containing 100 mM BTP, 1.7–1.9 M ammonium phosphate, 100 mM NaCl, pH 7.0. Diffraction quality crystals were obtained with the micro-seeding method at 4°C. The crystals all contain eight copies of the complex in the asymmetric unit. The data sets were collected at the BL8.2.1 beamline at the Advanced Light Source in Lawrence Berkeley National Laboratory as well as the GM/CA-CAT 23-ID-B beamline at the Advanced Photon Source in Argonne National Laboratory using crystals flash-frozen in the crystallization buffers supplemented with 15–20% ethylene glycol at ~170°C. Reflection data were indexed, integrated and scaled with the HKL2000 package. All crystal structures were solved by molecular replacement using the program Phaser and the TIRI–ASK1 structure as search model. The structural models were manually built in the program CNS and refined using PHENIX. All final models have 96–98% of residues in the favoured region and 0% in disallowed region of the Ramachandran plot.

Hormone and inositol phosphate reagents. H-coronatine was synthesized by Amersham. Coronatine was purchased from Sigma; JA-Il conjugates were chemically synthesized as previously described. Synthetic inositol phosphates were purchased as sodium salts from Cayman Chemicals.

Radioligand binding assay. Radioligand binding was assayed on purified proteins, with 2 μg COI1–ASK1 complex and JAZ proteins at a 1:3 molar ratio, and/or 10 μM synthetic peptides. Reactions were prepared in 100 μl final volume and in a binding buffer containing 20 mM Tris-HCl, 200 mM NaCl and 10% glycerol. Saturation binding experiments were conducted with serial dilutions of H-coronatine in binding buffer. Nonspecific binding was determined in the presence of 300 μM coronaetine. Competition binding experiments were conducted with serial dilutions of JA-Ile in the presence of 100 nM H-coronatine with nonspecific binding determined in the presence of 300 μM coronaetine. Total binding was determined in the presence of vehicle only. Two-point binding experiments were performed in the presence of 100 nM or 300 nM H-coronatine with nonspecific binding determined in the presence of 300 μM coronaetine. Following incubation with mixing at 4°C, all samples were collected with a cell harvester (Brandel, Gaithersburg, MD) on polyethyleneimine (Sigma)-treated filters. Samples were incubated in liquid scintillation fluid for >1 h before counting with a Packard Tri-Carb 2200 CA liquid scintillation analyser (Packard Instrument Co.). Saturation binding experiments were analysed by nonlinear regression, competition binding experiments by nonlinear regression with Kc, calculation as per the method of ref. 34, and concentration-response data by sigmoidal dose–response curve fitting, all using GraphPad Prism version 4.0 for Mac OSX.

Yeast two-hybrid assay. The coding sequences (CDS) of the Arabidopsis thaliana gene COI1 (At2g39940) and coi1 site-directed mutants were cloned into the yeast two-hybrid bait vector pGILDA (Clontech) using Xmal and Xhol restriction enzyme recognition sequences previously added to the 5’ and 3’ end of the COI1 CDS, respectively, creating DNA-binding domain (LexA–COI1 and LexA–coi1) protein fusions. The CDS of Arabidopsis thaliana JAZ1 gene (At1g19180) was cloned into the yeast two-hybrid prey vector pB42AD (Clontech) creating a transcriptional activation domain (AD–JAZ1) fusion protein. Individual wild-type and mutant COI1 constructs were co-transformed with JAZ1 constructs into Saccharomyces cerevisiae strain EGY48 (p8pLacZ) using the frozen-EZ yeast transformation II kit (Zymo Research). Transformants were selected on SD-glucose medium (BD Biosciences) supplemented with -Ura/-Trp/-His drop-out solution (BD Biosciences). To detect the interaction between COI1 and JAZ1, transformants that had been selected in SD-Glu medium were re-suspended in sterile water. Ten microlitres of each suspension was spotted onto inducing media (SD-Galactose/Raffinose–UWHi BD Biosciences) supplemented with 80 μg ml⁻¹ 5-X-Gal and 50 μM coronaetine (Sigma). Yeast two-hybrid assay plates were incubated in the dark at 20°C and photographed 7 days later. Induced yeast cells were analysed for COI1 and JAZ1 expression levels by western blotting using epitope-specific antibodies (data not shown).

Inositol phosphate purification. Phenol was melted at 68°C and equilibrated with equal parts 0.5 M Tris-HCl, pH 8.0 until a pH of 7.8 was reached. The equilibrated phenol was then topped with 0.1 volume 100 mM Tris-HCl, pH 8.0 and stored at 4°C. For extraction, 30–40 mg of 1 M mg⁻¹ COI1–ASK1 protein was mixed in small batches with equal parts equilibrated phenol at room temperature. The samples were inverted and incubated for 30 min until phase separation occurred. With 30 s vortexing, the samples were incubated at room temperature for 30 min and spun at 15,000 r.p.m. for 5 min. The aqueous phase was removed as a primary extraction. Equal parts of a solution containing 25 mM Tris-HCl, pH 8.0 was added to the phenol and collected as above as a secondary extraction. The primary and secondary extractions were then combined and diluted 10X in 25 mM Tris-HCl, pH 8.0, then further purified by gravity flow on Q sepharose high-performance anion exchange resin (GE Healthcare). Following column wash with 10× column volumes of 0.1 N formic acid, stepwise elution was performed with 2× column volumes of 0.1 N formic acid (Thermo Scientific) with increasing concentrations of ammonium formate (Sigma), from 0 to 2 M.

Fractions were analysed for phosphate content by the wet-ashing method with perchloric acid in Pyrex culture tubes (13 X 100 mm). Typically, samples of 50–100 μl were ashed with 100–200 μl 70% perchloric acid (purified by redistillation, Sigma). Ashing was performed by heating the sample over a Bunsen-type burner with continuous shaking to prevent bumping. When the sample stopped emitting white smoke, the reaction was considered complete and then heated to dryness. 500 μl of distilled water was added to the room temperature tubes and vortexed. 100 μl samples containing up to 10 nmol inorganic phosphate were assayed for phosphate by a modification of a published procedure. A total of 125 μl of 50% molar bimolecular reagent was added and the samples were incubated and covered at room temperature for 12–14 h (overnight) for full colour development (total volume 225 μl). Plates were read at 650 nm and unknowns were determined from the linear regression of the standard curve (0–10 nmol NaH2PO4 per well). All assays were done in triplicate. Final fractions containing phosphate were combined and lyophilized repeatedly to remove residual ammonium formate.

Inositol phosphate reconstitution assays. COI1–ASK1 complex was separated from pre-bound inositol phosphate by dialysis. Briefly, proteins were mixed with 10% glycerol and incubated in 2 M ammonium phosphate, 100 mM Bis-Tris propane pH 7.0, 200 mM NaCl, 10% glycerol, at 4°C for >24 h with a minimum of 3X buffer changes at 100X sample volume. Samples were then transferred to 20 mM Tris-HCl, pH 8.0, 200 mM NaCl, 10% glycerol, at 4°C for >24 h with a minimum of three buffer changes at 100X sample volume. Inositol phosphate rescue experiments were conducted according to the radioligand binding assays described above in the presence of 300 nM H-coronatine with nonspecific binding determined in the presence of 300 μM coronaetine.

Structural mass spectrometry analysis of the intact protein complex. Nano-electrospray ionization mass spectrometry (MS) and tandem MS (MS/MS) experiments were performed on a Synapt HDMS instrument. Before MS analysis, 50 μl of a 16 mg ml⁻¹ solution of COI1–ASK1 in 20 mM Tris-HCl pH 8.2, 0.2 M NaCl and 5 mM DTT, was buffer-exchanged twice into 0.5 M ammonium acetate solution by using Bio-Rad Bioskop columns. To improve desolvation during ionization, samples were diluted 1:4 in 0.5 M ammonium acetate and isopropanol was added to a final concentration of 5%. Typically an aliquot of 2 μl solution was loaded for sampling via nano-ESI capillaries which were prepared in-house from borosilicate glass tubes as described previously. The conditions within the mass spectrometer were adjusted to preserve non-covalent interactions. The following experimental parameters were used: capillary voltage up to 1.26 kV, sampling cone voltage 1.26 kV, sampling cone voltage 150 V and extraction cone voltage 6 V, MCP 1590. For tandem MS experiments peaks centred at m/z 4,564 and 4,588 were selected in the quadrupole and collision energy up to 65 V was used. Argon was used as a collision gas at maximum pressure. All spectra were calibrated externally by using a solution of sodium iodide (100 μg ml⁻¹). Spectra are shown with minimal smoothing and without background subtraction.

Nuclear magnetic resonance (NMR) analysis. NMR spectra were acquired on a Varian INOVA A600 spectrometer equipped with a cold probe using 200 μM samples of synthetic peptide or synthetic inositol-1,2,4,5,6-pentakisphosphate (Cayman...
Chemical) dissolved in D$_2$O. TOCSY spectra were acquired with mixing times of 35 or 50 ms, processed with NMRPipe$^{37}$ and visualized with NMRView$^{38}$.

**Mass spectrometry analysis of inositol phosphate purified from COI1–ASK1.**

MS experiments were conducted on a Finnigan LTQ linear ion-trap mass spectrometer (ITMS) with Xcalibur operating system. Methanol was continuously infused (10 µl min$^{-1}$) to the ESI source, where the skimmer was set at ground potential, the electrospray needle was set at 4.5 kV, and the temperature of the heated capillary was 275 °C. The sample was diluted with equal volume of 2% ammonia in methanol and 10 µl was flow injected. The automatic gain control of the ion trap was set at $2 \times 10^4$, with a maximum injection time of 50 ms. Helium was used as the buffer and collision gas at a pressure of $1 \times 10^{-3}$ mbar (0.75 mTorr). The MS$^n$ ($n = 2, 3, 4, 5$) experiments were carried out with an optimized relative collision energy ranging from 12% to 16% with an activation q value at 0.25. The activation time was set at 30–60 ms. The mass spectra were acquired in the profile mode and were accumulated for 3–5 min for MS$^n$ spectra. The mass resolution of the instrument was tuned to 0.6 Da at half peak height.

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