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An extracellular steric seeding mechanism for Eph-ephrin signalling platform assembly

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Abstract

Erythropoetin-producing hepatoma (Eph) receptors are cell surface protein tyrosine kinases mediating cell-cell communication. Upon activation they form signalling clusters. We report crystal structures of the full ectodomain of human EphA2 (eEphA2), alone and in complex with the receptor-binding domain of the ligand ephrinA5 (ephrinA5^{RBD}). Unliganded eEphA2 forms linear arrays of staggered parallel receptors involving two patches of residues conserved across A-class Ephs. eEphA2-ephrinA5^{RBD} forms a more elaborate assembly, whose interfaces include the same conserved regions on eEphA2, but re-arranged to accommodate ephrinA5^{RBD}. Cell surface expression of mutant EphA2s demonstrated that these interfaces are critical for localization at cell-cell contacts and activation-dependent degradation. Our results suggest a 'nucleation' mechanism whereby a limited number of ligand-receptor interactions seed an arrangement of receptors which can propagate into extended signalling arrays.

Ephs constitute a quarter of all known human receptor tyrosine kinases. They direct key processes during development and repair of the nervous system, blood vessel formation, insulin secretion, immune system function, intestinal homeostasis and bone tissue integrity¹. They are grouped into two classes, A and B², but domain composition is conserved across the family (Fig. 1a). The extracellular region comprises an N-terminal ligand-binding domain (LBD), a cysteine-rich region and two fibronectin type III domains (FN1 and FN2). FN2 connects to the transmembrane helix (TM), followed by an intracellular region comprising a juxtamembrane region (JM), a tyrosine kinase domain and a sterile-alpha motif (SAM) domain often linked to a C-terminal PDZ binding motif. Structures have been determined for examples of all Eph domains in isolation, with the exception of the cysteine-rich region^{3,4}.

Activation of Eph receptors depends on the presence of their ligands (ephrins) and involves the packing of Ephs into signalling clusters^{1,4}. Ephrins consist of an N-terminal extracellular receptor-binding domain (ephrin^{RBD}) and a C-terminal extension linked to the plasmamembrane by a lipid anchor (class A ephrins) or transmembrane helix (class B ephrins). Binding affinities vary for different Eph-ephrin pairs, but in general binding within classes is favoured⁵. Crystal structures for 1:1 complexes of Eph LBDs bound to ephrin^{RBD}s show a hydrophobic ephrin-binding groove on the receptor and provide insight into ligand-receptor specificity, but do not define a mechanism for ligand-dependent signalling⁶. Members of the

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E.S. conducted crystallographic and cellular studies. K.H. performed crystal mounting and oversaw X-ray data collection. G.S. led the MALS analysis. A.R.A. and E.Y.J. participated in study design and oversaw all aspects of the work.

Atomic coordinates and structure factors will be deposited within the Protein Data Bank.

Eph family of receptor tyrosine kinases appear to require a more complex form of oligomerisation to trigger signalling than the ligand-induced dimerisation model developed from studies of systems such as the human growth hormone receptor⁷. Insights into Eph receptor clustering upon ephrin binding are as yet limited⁴. The crystal structure of the EphB2 LBD - ephrinB2^{RBD} complex revealed a second, putative tetramerization, interface⁶. However, five subsequent Eph LBD - ephrin^{RBD} complex crystal structures have not shown evidence for a consistent mode of oligomerisaton⁸⁻¹¹. Biophysical and functional analyses have also implicated additional regions of Eph ectodomains in ephrin mediated signalling¹². We therefore sought to provide structural insight into a full Eph receptor ectodomain architecture and its interactions. We focus on EphA2 which is expressed during embryonic development and in adult epithelia¹³ where it regulates cell adhesion^{14,15}. EphA2 also acts as a powerful oncogene in many tumours by promoting vascularisation and metastasis¹⁶⁻¹⁹.

RESULTS

Structure of EphA2 ectodomain (eEphA2)

To improve crystallization, we expressed eEphA2 (residues K27-N534) in glycosylationimpaired mammalian cells^{20,21}, trimmed the resultant glycans down to single Nacetylglucosamine moieties and di-methylated the amino groups²². The structure (one molecule per asymmetric unit) was refined to a final $R_{work} = 24\%$ ($R_{free} = 30\%$) using data to 3.0 Å resolution. Detailed data and refinement statistics are given in Table 1. A view of the final electron density map is shown in Supplementary Fig. 1. All five domains are wellordered and the overall linear arrangement results in an elongated molecule (length ~172 Å, width ~22-48 Å, Fig. 1b). The previously uncharacterized cysteine-rich region shows the topology of a sushi (CCP) domain followed by an EGF-like domain. Sequence alignment of class A Ephs mapped onto the eEphA2 structure reveals patches of surface conservation on the LBD and sushi domains (Fig. 1c). Conserved sushi domain residues contribute to a crystal contact with a parallel neighbouring molecule staggered by one domain (Fig. 1c). In total, interfaces between these molecules bury 2661 Å² of surface area and stabilise linear arrays of staggered, parallel Ephs. These crystal contacts for full length EphA2 ectodomain do not re-iterate the "head-to-head" contacts recently observed for crystals of EphA2 LBD¹⁰. Two putative N-linked glycosylation sites, located in the FN1 domain and in the linker between domains FN1 and FN2, are not involved in crystal contacts.

Structure of eEphA2 in complex with ephrinA5^{RBD}

To compare unliganded eEphA2 with a ligand-bound form, we attempted to crystallize it with a panel of ephrinA ligands and succeeded in producing diffracting crystals of eEphA2 in complex with human ephrinA5 RBD (ephrinA5^{RBD}). Di-methylation of the amine groups and extensive optimization yielded a crystal (containing one eEphA2-ephrinA5^{RBD} per asymmetric unit) from which diffraction data were collected to 4.8 Å resolution. The resultant complex structure was refined (by rigid body and TLS) to R_{work} = 31%, R_{free} = 31% (Fig. 1d). Crystallographic details are summarized in Table 1 and an example of the final electron density map is shown in Supplementary Fig. 1. The structure of ephrinA5^{RBD} in the complex shows no evidence of major structure changes when compared to the previously reported structure²³ of unliganded ephrinA5^{RBD}. Superposition of ligand-bound and unbound eEphA2s reveals domain re-orientations (Supplementary Fig. 2). The largest re-orientation, a relative rotation of 71° by FN2, demonstrates the hinge-like character of the FN1-FN2 linker. Other points of flexion are located between the LBD and sushi domain and within the EGF-like domain.

The crystal consists of parallel and anti-parallel arrays of eEphA2-ephrinA5^{RBD} complexes (Supplementary Fig. 3) formed by interfaces A-F (Supplementary Fig. 4 and Supplementary

Table 1). Interfaces A-E mediate parallel arrays (Fig. 1e and Fig. 2a), interface F stabilizes anti-parallel packing (Fig. 2b). The previously reported Eph-LBD - ephrin^{RBD} interaction³ forms interface A and broadly matches that previously reported for the complex of EphA2 LBD with ephrinA1^{RBD10} (Fig. 2a, c). Interface C is a small contact between eEphA2 and a second ephrinA5^{RBD} (Fig. 2a, c). Interfaces B, D and E connect neighbouring eEphA2 molecules via their LBDs, sushi domains and FN1 domains, respectively (Fig. 2a, c-e). Interfaces B and D use the conserved surface patches which form the staggered LBD-sushi contact in the unliganded eEphA2 crystals, but in the presence of ephrinA5^{RBD} these patches mediate in-register LBD-LBD (interface B) and sushi-sushi (interface D) interactions (Fig. 1e). Finally, the FN1-FN2 domains from eEphA2 form interface F with ephrinA5^{RBD} bound (via interface A) to an anti-parallel eEphA2 neighbour (Fig. 2b, f).

EphA2 cell-cell contact localization is not dependent on the intracellular domains

Recent work has shown that EphA2 localizes at the cell-cell contacts in the mouse eye lens where it binds to ephrinA5 and maintains tight cellular packing²⁴. EphA2 is also known to localize at cell-cell contacts when transfected into a range of epithelial cell lines²⁵⁻²⁷ and we therefore chose to exploit this system to dissect the interaction characteristics of EphA2 in vivo using a human embryonic kidney epithelial cell line (HEK293T) and fluorescent protein (mVenus)-tagged constructs (Fig. 3). A series of deletion constructs all showed cell surface expression (Supplementary Fig. 5). As anticipated, full length EphA2 accumulated at the contact region between transfected cells. We did not observe EphA2 accumulation in regions of the cell surface not involved in contacts, or in contact regions between EphA2expressing and non-transfected native cells (Fig. 3a-c). These observations imply that trans presentation of EphA2 (i.e. from opposing cells) is required for clustering. We found that deleting individual cytoplasmic domains, or the entire intracellular region, did not prevent cell surface expression and did not affect the pattern of localization at cell-cell contacts (Supplementary Fig. 6). In contrast, although cell surface localization is still maintained, deletion of the N-terminal four extracellular domains (LBD-FN1) abolishes the specific localization at the cell-cell interface, suggesting that the ectodomain is essential (Supplementary Fig. 6).

Mutations in interfaces A, B, D, F affect cell-cell contact localization

To investigate which of the interfaces in our crystal structures have biological relevance, we produced a panel of EphA2 point mutants designed to disrupt interfaces A-F (Supplementary Table 1). Interface B and D mutations result in homogeneous EphA2 cell surface distribution, completely abolishing the wild type localization pattern (Fig. 3d-f and Supplementary Fig. 6). These are the major interfaces for the in-register arrays of liganded eEphA2, the same surfaces form the staggered arrays for unliganded eEphA2, and *in vivo* could mediate *in cis* interaction, i.e. of Eph receptors on the same cell. Previous mutagenesis studies have indicated interface B residues can contribute to EphA3 – ephrinA5 binding affinity¹², although it is not the main ephrin binding site. Mutations in the minor interfaces C and E had no effect on EphA2 localization (Supplementary Fig. 6).

Introduction of an N-linked glycosylation site (EphA2 A190N + L192S) to disrupt interface A, the well-established ephrin binding site for binding *in trans*, abolished cell-cell contact localization (Supplementary Fig. 6). As HEK293 cells express class A ephrins endogenously²⁸, these results suggest that binding to endogenous ephrins *in trans* is necessary to drive localization of expressed EphA2 to cell-cell contacts, a conclusion further supported by studies using insect (SF9) cells (Supplementary Fig. 6c). Interface F (the second largest interface in the complex crystals) would *in vivo* be consistent with a *cis*-interaction between ephrinA5 and EphA2 FN1-FN2; mutations at this interface caused an intermediate phenotype where protein is partially de-localized over the entire cell surface

(Supplementary Fig. 6). This is consistent with previous functional studies suggesting the FN2 domain of EphA3 binds ephrinA5 *in cis*²⁹.

Specific proteolytic cleavage of EphA2 depends on clustering

Previous studies have demonstrated that activation of Eph receptors leads to their internalization and degradation³⁰ and specific activation-dependent proteolytic cleavage has been reported for EphB2^{31,32}. We performed immunoblot assays which show that EphA2 is proteolytically cleaved in HEK293 cells. Upon wild type EphA2 expression at least three C-terminal tyrosine-phosphorylated EphA2 proteolytic fragments are observed, corresponding to ~70, ~75 and ~95 kDa (including the ~28 kDa mVenus tag, Supplementary Fig. 7). For mutants in interfaces A, B and D the fragments of ~75 kDa and ~95 kDa are absent. For mutants in interface F only the ~95 kDa fragment is absent. Notably, the cleavage patterns observed for EphA2 wild type and mutants correlate with their cell surface localization patterns; interface F mutants having less severe impairment of cell-cell contact localization than interface A, B, and D mutants (Supplementary Fig. 6 and 7). The correlation of specific cleavage with localization to cell-cell contacts suggests that this process depends on the ability of the receptor to cluster via interfaces A, B, D and F.

DISCUSSION

Previously reported data for Eph function point to a requirement for receptor clustering into extended cell surface assemblies^{4,33-35}. However, full length Eph ectodomains (eEphA2, eEphB2 and eEphB3) show no propensity to form oligomers in solution (as assessed by multiple angle light scattering, MALS, and analytical ultracentrifugation) either alone or in complex with monomeric, soluble ephrins (ephrinA5^{RBD}, ephrinA2^{RBD} and ephrinB2^{RBD}) (Supplementary Fig. 8 and data not shown). These data accord with previous reports that full length Eph receptors only cluster if the ephrin ligand is presented in a membrane-bound or oligomeric form³⁶. Nevertheless, at very high concentration in crystals both unliganded and complexed eEphA2 form array-like networks of interfaces which functional experiments demonstrate are relevant to the cell surface signalling. The eEphA2 arrays involve staggered interactions between the LBD and sushi domains. Ephrin binding snaps the Eph arrays into register with LBD-LBD and sushi-sushi interactions involving the same eEphA2 surface patches. Whilst we were completing our analysis an independently determined crystal structure of unliganded human EphA2 ectodomain was deposited in the Protein Data Base³⁷. The 3FL7 lattice includes crystal contacts similar to the in-register form we observe indicating that this mode of Eph-Eph interaction can also occur without ephrin-binding. This implies that in the absence of ligand transient cis associations between cell surface Ephs may occur as a dynamic equilibrium between LBD-LBD/sushi-sushi and LDB-sushi interactions. For such a system ephrin-binding at "nucleation" points could trigger more widespread recruitment of EphA2 into in-register arrays containing unliganded receptor and facilitating, in a cooperative manner, additional ephrin-binding (Fig. 4).

Indeed, it has been reported that once an Eph receptor signalling cluster is nucleated by ephrin-binding, further Eph are recruited to the cluster in an ephrin-independent manner³³. The above steric 'seeding' mechanism provides a model for how relatively low levels of ephrins can trigger the formation of extended assemblies of EphA2. These assemblies may be further stabilized by transmembrane³⁸ and cytoplasmic³⁹ region interactions. Our results illustrate a nucleation mechanism for ectodomain-driven array-based signalling which could be broadly applicable to other cell surface receptor systems.

METHODS

Vectors and Cloning

Secreted polyhistidine-tagged ectodomains of the human EphA2 (eEphA2, residues 27-534, Uniprot: P29317) and ephrin A5 (ephrinA5^{RBD}, residues 27-166, Uniprot: P52803) were cloned into the AgeI-KpnI sites of pHLsec²¹ using standard PCR methods (EphA2 residues 1-23 and ephrinA5 residues 1-20 belong to the secretion signals in EphA2 and ephrin A5 as predicted by SignalP⁴⁰). For cell surface expression transmembrane constructs of human EphA2, fused to a C-terminal monomeric fluorescent protein mVenus (green)⁴¹, were cloned into pHLsec (coding additionally for a C-terminal poly-histidine tag). The following constructs were produced as mVenus fusions: EphA2 (residues 27-976), EphA2^{Δ SAM} (residues 27-889), EphA2^{Δ KSAM} (residues 27-564), EphA2^{Δ e} (residues 436-976). EphrinA5 (residues 27-166) was fused at its C-terminus to the monomeric transmembrane helix of receptor protein tyrosine phosphatase RPTPµ⁴² (residues 742-769, Uniprot: P28827) followed by the monomeric fluorescent protein mCherry, and cloned into pHLsec.

Protein Crystallization

Details of the protein expression and purification are given in the Supplementary Material Online. eEphA2 was treated with endoglycosidase F1^{43,44} and lysine-methylated ²². Crystals diffracting to 3 Å resolution grew at 20 °C from drops⁴⁵ where protein solution was mixed 1:1:1 with reservoir solution (20 % polyethylene glycol 6000, 1 M LiCl, 0.1 M Tris pH 8) and 0.4 M non-detergent sulfobetaine 256 (Hampton). Selenomethionine labeled protein²¹ was purified and crystallized as the native protein. Crystals of eEphA2 and ephrinA5^{RBD} complex grew in conditions similar to the that specified below, but diffracted only to 6-8 Å resolution. Only one crystal diffracted well to 4.8 Å resolution. This crystal grew at 20 °C in a drop where two parts of protein solution were mixed with one part water, one part reservoir solution (8 % polyethylene glycol 6000, 0.8 M LiCl, 0.08 M citrate pH 5) and one part 1 % polyvinylpyrrolidone K15 (Hampton).

Data collection and processing

Data statistics are summarized in Table 1. Selenomethionine derivative crystals of eEphA2 were flash-frozen after briefly dipping the crystal containing loop into perfluoropolyether oil PFO-X125/03 (Lancaster). A data set was collected at 100 K at the selenium edge (wavelength = 0.98 Å) at the ESRF beamline BM14 using the inverse beam option (rotation every 20 images, corresponding to 10 degrees). This crystal diffracted to 3 Å resolution. The data were processed using XDS⁴⁶ and MOSFLM⁴⁷. Molecular replacement using models for the ligand binding domain of EphA2 (PBD entry 3C8X), the fibronectin type III domain 2 of EphB4 (PDB entry 2E7H), and fibronectin type III domain 2 of EphA8 (PDB entry 1X5L) yielded initial phases that were used to locate selenium sites in PHASER⁴⁸. Electron density modification was done with DM⁴⁹ and manual improvement of the protein model in COOT⁵⁰. Cycles of phase calculation plus model building and refinement iteratively lead to a satisfactory model. A second crystal yielded better data at high resolution at the ESRF beamline ID23-EH2 (100 K, wavelength = 0.87 Å) and was used for the final rounds of refinement. eEphA2 carries two predicted N-glycosylation sites, the first is on the FN1 domain where N-acetyl-glucosamine (NAG) was placed into the electron density. This NAG molecule is in close proximity to, but not in contact with the FN2 domain of a neighbouring EphA2 chain. The second putative N-glycosylation site located in the poorly defined linker region between FN1 and FN2 and lacked electron density indicating the presence of sugar, possibly due to the relative disorder of this region. This region is also not involved in crystal contacts. COOT and MOLPROBITY⁵¹ were used for model validation. 3 residues (0.6 %) are located in outlier regions of the Ramachandran plot.

Crystals of eEphA2 in complex with ephrinA5^{RBD} were frozen in a cryoprotectant solution containing one part reservoir solution and one part reservoir solution supersaturated with LiCl. After brief equilibration in cryoprotectant, the crystals were mounted on a loop which was then dipped into perfluoropolyether oil PFO-X125/03 and flash-frozen. The best dataset was collected at 100 K on ESRF beamline ID14-EH2 (wavelength = 0.93 Å) and processed using XDS. A model of human EphA2 ligand binding domain in complex with ephrin A2 receptor binding domain (PDB code 3CZU) was used for molecular replacement in PHASER (RFZ=11.1, TFZ=9.6, LLG=162). We placed the models of individual eEphA2 domains sequentially into the improving density and refined each using rigid body refinement in PHASER. Molecular replacement was initially done using the 3.0 Å resolution eEphA2 model presented here. After deposition of the 2.5 Å resolution eEphA2 model in the Protein Data Bank (PBD code 3FL7) by the Structural Genomics Consortium, Toronto, we used this higher resolution model. The model was refined using PHENIX⁵² (Supplementary Material Online). 6 residues are in outlier regions of the Ramachandran plot as found in the input molecular replacement model downloaded from the Protein Data Bank. The two putative N-glycosylation sites lie outside crystal contact areas.

To calculate the angle between the orientations eEphA2 FN2 adopts in the unliganded and ephrinA5^{RBD}-bound structures, we first generated two models of liganded eEphA2 superposed onto the FN1 and FN2 domains of unliganded eEphA2. We then used LSQKAB within the CCP4 suite⁵³ to calculate the rotation applied to superpose the two resulting structures via their FN2 domain.

Multiangle Light Scattering (MALS)

Proteins were purified by size-exclusion chromatography and concentrated to approximately 1.6 ml/ml (eEphA2) and 1.8 mg/ml (eEphA2/ephrinA5^{RBD} complex). Separation for the MALS was achieved using an analytical Superdex S75 10/30 column (GE Heathcare) and the eluate was passed through online static light scattering (DAWN HELEOS II, Wyatt Technology, Santa Barbara, CA), differential refractive index (Optilab rEX, Wyatt Technology) and Agilent 1200 UV (Agilent Techologies) detectors. Data were analysed using the ASTRA software package (Wyatt Technology).

Live cell assays

HEK293T cells were grown in Dulbecco's Modified Eagle's Medium (DMEM high glucose, Sigma) supplemented with L-glutamine, non-essential amino-acids (Gibco) and 10% foetal calf serum (FCS, Sigma). Plasmid DNA was transfected with lipofectamine 2000 following the recommended protocol (Invitrogen). Final concentrations were 2 µg plasmid DNA per ml, 5 µl lipofectamine per ml and 1.5% FCS. After 4-6 hours, cells were returned to growth medium, which contains 10% FCS, and plated on poly-L-lysine-coated glass bottom culture plates (MatTek). To observe *trans* interactions of different constructs, cells were transfected separately and plated out together at this stage. 15-20 hours later cells were imaged using a Zeiss LSM510 confocal microscope (with variable emission detectors [Meta-Head]). For this time-scale, we do not observe cell-rounding and death upon EphA2 expression for either HEK293 or COS7 cells (data not shown). All images were taken with the same detector and laser settings and the images showing fluorescent signal were not subsequently modified, thus allowing direct comparison.

Immunoblot analysis

To assess cell surface expression of truncated EphA2 constructs, cells were incubated for 10 minutes at 4 °C to slow down endocytosis and then incubated 30 min with 1 mg/ml No-Weigh Sulfo-NHS-LC-Biotin (Pierce) at 4 °C. The biotinylation reaction was stopped by adding 100 mM Tris pH 7.5. Cells were lysed in phosphate buffer saline supplemented with

100 mM Tris pH 7.5 and 1% Triton X-100 (Sigma). Cell lysates were incubated for 30 minutes with streptavidin-coated agarose beads (Pierce) at 4 °C, the beads washed three times with lysis buffer and then re-suspended in standard protein denaturing gel loading buffer for immunoblot analysis using anti-pentahistidine (penta-His, Qiagen, dilution 1:1000). Cells intended for anti-phosphotyrosine or anti-green fluorescent protein immunoblot analyses were re-suspended in chilled phosphate buffer saline 15-20 hours post transfection, immediately mixed with standard protein gel loading buffer and analyzed using anti-phosphotyrosine (4G10® Platinum, Millipore, dilution 1:2000), anti-GFP (Cat. No. A-11122, Invitrogen, dilution 1:1000), anti-actin (ACTN05 (C4), AbCam, dilution 1:400), anti-mouse IgG peroxidase-conjugated secondary antibody (Sigma, dilution 1:1000) and anti-rabbit IgG peroxidase-conjugated secondary antibody (AbCam, dilution 1:5000). To rule out that FCS or trace amounts of trypsin used for cell propagation are responsible for EphA2 cleavage, we performed control experiments in cells cultured with 0% or 1.5% FCS, in the presence of 0%, 0.5% or 5% of the trypsin-EDTA solution used for tissue culture maintenance (Invitrogen, Cat. No. 25300-054) (data not shown).

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

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Figure 1. Crystal structures of the complete EphA2 extracellular region (eEphA2) alone and in complex with ephrinA5 $^{\rm RBD}$

a, Domain composition of ephrinAs and Eph receptors. **b**, Crystal structure of eEphA2 comprising EphA2 LBD, Cys, FN1 and FN2 domains. Colours are as in a. The previously uncharacterized Cys domain is composed of a sushi domain (CCP) and an EGF domain. **c**, Arrangement of two eEphA2 molecules within the crystal. Their contacts bury 2661 Å² of molecular surface. The front molecule is displayed and oriented as in a. The back molecule is shown in surface view and coloured according to human Eph class A receptor sequence conservation (red = strongest conservation, white = weakest conservation). **d**, eEphA2 (colours as in a) in complex with ephrinA5^{RBD} (cyan). The orientation of the sushi domain is the same as in b and c. **e**, Arrangement of neighbouring complexes within the crystal. The eEphA2 molecule on the right is oriented and coloured as in d. Its neighbour on the left is in surface view and coloured according to EphA2 sequence conservation. Their contacts bury 1240 Å².



Figure 2. The eEphA2-ephrinA5^{RBD} complex: structural changes within eEphA2 and formation of array-like clusters

a, **b**, Crystal packing of eEphA2-ephrin $A5^{RBD}$. The ribbon diagrams are coloured as in Fig. 1. The cartoon above each panel illustrates the crystal contacts designated interfaces A-F (dotted lines) for eEphA2 (purple) and ephrin $A5^{RBD}$ (cyan). Panel a shows six symmetry-related eEphA2-ephrin $A5^{RBD}$ complexes forming a parallel array; view is approximately along unit cell axis c. The protein arrangement reflects EphA2 and ephrinA5 interaction *in trans*, i.e. where the proteins are located on opposing cells. Panel b shows six symmetry-related eEphA2-ephrin $A5^{RBD}$ complexes forming an anti-parallel array, view as in f, but tilted by ~45° around unit cell axis a. The arrangement shows EphA2 and ephrinA5 interacting both *in trans* and *in cis*, consistent with both proteins presented on both cells. **c-f**, Interfaces A-F are shown as black lines. Colours are as in a. Interfaces A-E form the crystal contact interactions shown in a, interface F is a major interface but contributes only to the contacts shown in b.



Figure 3. EphA2 localization at cell-cell contacts depends on ectodomain clustering

Representative results are presented of functional studies using HEK293T cells. A full set of results is provided in Supplementary Fig. 6. Filled white arrows indicate cell-cell contacts between transfected cells. Open arrows point to cell-cell contacts between a transfected cell and a non-transfected cell. **a-c**, mVenus-tagged EphA2 accumulates at cell-cell contacts of transfected cells. **d-e**, Mutants in interfaces A, B, D and, to some extent, interface F do not cluster at cell-cell contacts of transfected cells but are distributed evenly on the cell surface (the interface B mutant G131Y is shown as an example). **a**, **d** Fluorescence images are shown. **b**, **e**, Phase contrast images of the corresponding cells. **c**, **f**, Overlay images of a, b, d, e.



Fig.4. A mechanism for the nucleation and propagation of EphA2 signalling assemblies

a, **b**, Eph and ephrin are shown schematically with ectodomains colour coded as in Fig. 1a and the Eph intracellular region in grey. Lightening flashes indicate the strength of activation. In panel a transient arrays of staggered Ephs snap into in-register signalling arrays on nucleation by ephrins. The intermediate phenotype of interface F mutations in functional assays may reflect weaker interactions between Eph receptors and their ligands, resulting in a weakened signalling activity. In panel b additional *cis* (interface F) interactions inter-connect anti-parallel arrays further stabilising the signalling assemblies.

	eEphA2 (refinement)	eEphA2 (phasing)	eEphA2+ephrinA5 ^{RBD}
Data collection			
RCSB PDB code	XXX	XXX	XXX
Space group	P1	P1	P2221
Cell dimensions			
a, b, c (Å)	41.92, 44.65, 93.42	41.47, 44.35, 92.08	173.63, 59.63,112.16
$\alpha,\beta,\gamma(^{\circ})$	93.02, 98.22, 112.12	93.41, 97.22, 112.44	90, 90, 90
Resolution (Å) $*$	46-3.0 (3.1-3.0)	45-3.0 (3.1-3.0)	173.6-4.8 (4.9-4.8)
R _{meas} *	0.143 (0.443)	0.097 (0.645)	0.112 (0.743)
<i>Ι</i> /σ <i>Ι</i> [*]	9.87 (2.60)	8.97 (1.74)	12.95 (2.83)
Completeness (%)*	97.6 (94.5)	97.8 (97.7)	98.2 (75.5)
Redundancy *	3.3 (2.3)	2.2 (2.0)	6.2 (6.3)
Refinement			
Resolution (Å) *	45.9-3.0 (3.3-3.0)	-	40.5-4.8 (6.1-4.8)
No. reflections	12037	-	5643
$R_{ m work}/R_{ m free}$	0.236/0.299	-	0.312/0.314
No. atoms	3743	-	4889
Protein	3728	-	4889
Ligand/ion	15	-	-
Water	-	-	-
average B-factors			
Protein	100.4	-	247.2
Ligand/ion	107.6	-	-
Water	-	-	-
R.m.s. deviations			
Bond lengths (Å)	0.004	-	0.003
Bond angles (°)	0.697	-	0.675

 Table 1

 Data collection and refinement statistics

*Values in parentheses are for highest-resolution shell.