Enhanced internalization of ErbB2 in SK-BR-3 cells with multivalent forms of an artificial ligand

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ABSTRACT

Targeting and down-regulation of ErbB2, a member of EGF receptor family, is regarded as one of the key aspects for cancer treatment because it is often overexpressed in breast and ovarian cancer cells. Although natural ligands for ErbB2 have not been found, unlike other ErbB receptors, EC-1, a 20-amino acid circular peptide, has been shown to bind to ErbB2 as an artificial ligand. Previously we showed EC-1 peptide did not induce the internalization of ErbB2 in SK-BR-3 cells. In this report, we designed divalent and multivalent forms of EC-1 peptide with the Fc portion of the human IgG and bionanocapsule modified with ZZ-tag on its surface to improve the interaction with ErbB2. These forms showed higher affinity to ErbB2 than that of EC-1 monomer. Furthermore, prominent endosomal accumulation of ErbB2 occurred in SK-BR-3 cells when stimulated with EC-Fc ligand multivalently displayed on the surface of the bionanocapsule, whereas SK-BR-3 cells as themselves displayed stringent mechanism against ErbB2 internalization without stimulation. The multivalent form of EC-1 peptide appeared to internalize ErbB2 more efficiently than divalent form did. This internalization was unaffected by the inhibition of clathrin association, but inhibited when the cholesterol was depleted which explained either caveolar or GPI-AP-early endocytic compartment (GEEC) pathway. Because of the lack of caveolin-1 expression, caveolar machinery may be lost in SK-BR-3 cell line. Therefore, it is suggested that the multivalent form of EC-1 induces the internalization of ErbB2 through the GEEC pathway.

KEYWORDS: ErbB2 • EC-1 peptide • internalization • multivalent display • bionanocapsule

INTRODUCTION

The epidermal growth factor receptor (EGFR) family consisting of ErbB1 (EGFR), ErbB2, ErbB3 and ErbB4 is known to play a crucial part in the development and progression of cancers. The signal for cell survival, differentiation and proliferation is achieved through both homo- and heterodimerization of the receptors [1, 2]. Anomalous increase in the expression of the ErbB2 is an identified factor for the development of several cancer types [1, 3] and high ErbB2 levels parallel the worse prognosis for breast cancer patients [4-6]. ErbB2 also associates with ErbB3 in the heterodimerization and is considered the most active dimer that initiates cancers [7, 8]. ErbB2 is overexpressed in breast (30%) and ovarian (15–30%) cancer [9-11]. The myriad roles played by ErbB2 have made it one of the most preferred targets for the treatment of cancer. ErbB2 stringently internalizes efficiently recruiting back to the cell surface [12]. The mechanism of ErbB2 recycling and degradation is complex and appears to vary depending on the cell types. Various groups have shown that geldanamycin treatment efficiently internalizes the ErbB2 ligand [13-15]. Although the
internalization of ErbB2 is enhanced when multivalently cross-linked with antibody such as anti-ErbB2 antibody sc-08 [13, 16–19], there is no known natural ligand that promotes the ErbB2 internalization till date. By phage display method, small peptides, which possess binding affinity to ErbB2, have been isolated [20–22]. EC-1 peptide, a cyclic 20-amino-acid-peptide, is also one of the artificial ErbB2 ligands isolated from random peptide library of phage display [23]. EC-1 peptide is considered to abolish the tyrosine phosphorylation of the intracellular domain of ErbB2 and to inhibit the proliferation of SK-BR-3 cells overexpressing ErbB2.

Previously, EC-1 peptide fused to GFP was shown to stimulate internalization of ErbB2 in SK-BR-3 cells whereas ErbB2 in SK-BR-3 cells retained on the surface [24]. Because anti-ErbB2 antibody sc-08 stimulated the internalization of ErbB2 in both cell lines, the mechanism of internalization stimulated by EC-1 peptide was considered quite different from that by anti-ErbB2 antibody sc-08. In this study, we designed multivalent form using EC-1 peptide fused to human IgG-Fc domain (EC-Fc) and bionanocapsule displaying ZZ-tag on the surface [25]. We then asked whether the multivalent forms stimulated the internalization of ErbB2 in SK-BR-3 cells.

**Materials and methods**

**Cell cultures**

Human breast carcinoma derived cell lines, MCF-7, MDA-MB-453 and SK-BR-3, and human ovarian carcinoma cell line SK-OV-3 were from the ATCC (VA). SK-BR-3 and SK-OV-3 cells were grown at 37°C in RPMI-1640 medium supplemented with 10% FBS and MCF-7 in DMEM medium supplemented with 10% FBS under an atmosphere of 5% CO2 and MDA-MB-453 cells in Leibovitz’s L-15 medium supplemented with 10% FBS without conditioning CO2. All culture media were supplemented with 2 mM glutamine just before use.

**Construction of expression plasmids**

The expression plasmid for EC-Fc was constructed as follows. The DNA fragment coding human IgG-Fc domain was excised from the plasmid pBOS93 [26] with restriction enzymes AgeI and NotI and then ligated at the 3’-end of a synthetic oligonucleotide coding for the EC-1 peptide. The resultant sequence coding EC-Fc was ligated at the 3’-end of the sequence coding the signal peptide derived from human RNase1 [27]. Finally, the sequence coding EC-Fc with a secretion signal peptide was cloned downstream of CMV promoter to construct the plasmid pBOS54 with hygromycin resistant gene. Simultaneously, the expression vector for Fc domain without EC-1 moiety was constructed in the same procedure to construct the plasmid pBOS53.

**Preparation of EC-Fc protein**

Chinese hamster ovary (CHO) cells were transfected with the expression plasmid pBOS54 for EC-Fc protein. The transformed cells were cultured in the presence of 100 μg/ml Hygromycin B (Wako Pure-Chemicals, Osaka, Japan) for several weeks and cells stably expressing EC-Fc were pooled. Ten million of the transformed cells were seeded in 500 ml of CHO-S-SFMII ( Gibco, Carlsbad, CA, USA) containing 100 μg/ml hygromycin at 37°C for 5 days. At the end of culture, the medium was collected and centrifuged to remove precipitates. The supernatants were passed over a 1 ml column of Protein G agarose (Invitrogen, Carlsbad, CA, USA) equilibrated with PBS. The column was then washed with PBS and the bound protein was eluted with 0.1 M sodium phosphate buffer, pH 2.6. Fractions containing EC-Fc were readily neutralized with sodium phosphate buffer, pH 8.0 by adding 1/10 volume of each fraction. The buffer was then replaced with PBS using a PD-10 column (GE Healthcare, Piscataway, NJ). The preparation of Fc protein without EC-1 moiety was based on the same procedure.

**Competition EIA to estimate Kd value**

The value of dissociation constant (Kd) between the ligand and ErbB2 was estimated as previously described with slight modification [24, 26]. Briefly, each well of 96-well plate was coated with 200 ng of sErbB2 in 0.1 M sodium bicarbonate buffer, pH 8.6, overnight by following with PBS containing 1% BSA. After the rinse with PBS containing 0.1% Tween 20 (PBST), EC-Fc and varying concentration of sErbB2, which were premixed 30 min before, was applied to the wells. After the incubation for 30 min at 25 °C, the wells were washed with PBST and 50 μl of protein A-HRP conjugate (Sigma-Aldrich) diluted to 1:10,000 was added and incubated for 1 hr. Wells were then washed eight times with PBST and 100 μl of OPD solution was added for the HRP reaction. After 15 min, the reaction was quenched by 50 μl of 0.5 M sulphuric acid and the absorbance at 492 nm of each well was measured by a microplate reader MTP-120 (Corona, Japan). Each experiment was performed in triplicate and the mean values and standard deviations were calculated.

**Preparation of multivalently displayed EC-1 peptide on bionanocapsule**

To prepare the multivalent form of EC-1 peptide, EC-Fc proteins were displayed on the surface of bionanocapsule (BNC), which was composed of the recombinant surface antigen of hepatitis B virus [29]. To confer the high affinity for IgG Fc domain to BNC, the surface antigen was designed to fuse with the ZZ motif in the protein A derived from *Staphylococcus aureus* [25]. The resultant ZZ-tagged BNC (ZZ-BNC) was labelled with FITC by incubating with 1 mg/ml of FITC at 25°C for 15 min and the reaction mixture was further incubated at 4°C overnight in 0.1 M sodium bicarbonate buffer, pH 8.6. The molar ratio of FITC to ZZ-BNC was 110 to 1. After incubation, free FITC was removed by PD-10 column equilibrated with PBS. To prepare multivalently displayed form of EC-1 peptide, the molar ratio of ligand and BNC was optimized using Fc protein and human IgG (Sigma-Aldrich). To optimize the ratio FITC-labelled ZZ-BNC was mixed and incubated with Fc protein or human IgG at different ratio of 1:10, 1:20, 1:40, 1:60, 1:80 and 1:100 for 1 hr at 4°C. The precipitates formed were removed by centrifugation at 13,000 rpm for 5 min and the supernatant was transferred to a 96-black-well plate and fluorescence intensity at 530 nm of each well was measured by a microplate reader MTP-800 (Corona, Japan). Each experiment was performed in triplicate and the mean values and standard deviations were calculated. To characterize binding capacity of EC-Fc protein to ZZ-BNC, EC-Fc protein and ZZ-BNC were mixed in different ratios and incubated for 1 hr at 4°C. After centrifugation, the EC-Fc/BNC complex in the supernatant was immunoprecipitated with...
anti-HBsAg microbeads (Abott) by incubation overnight at 4°C. The beads were then washed three times in PBS followed by centrifugation for 30 sec at 4°C. Then the beads were suspended in Laemnli buffer containing β-mercaptoethanol, boiled for 5 min at 95°C, and subjected to SDS-PAGE and Western blot with anti-human IgG Fc antibody. As the quantitative control of Western blot, 125, 250 and 500 ng of EC-Fc was applied at the same time. Thus, the efficient ratio of EC-Fc and ZZ-BNC was determined and the multivalent form of EC-1 peptide was prepared as EC-Fc/BNC. BNC bound to Fc protein (Fc/BNC) was prepared simultaneously as a control.

Confocal microscopic observation of ErbB2 internalization

For confocal microscopic observation, cells were grown on 35 mm glass base dish (Iwaki Science Products, Japan). With the conditioned media prepared from CHO cells expressing EC-Fc or Fc proteins in HEPES buffered RPMI-1640 supplemented with 1% BSA, the target cells were incubated for 90 min at 37°C. The cells were fixed with 4% paraformaldehyde in PBS, permeabilized with 0.2% Triton X-100 and were blocked with 1% BSA or 10% FBS in PBS. Then the cells were washed with PBS and incubated with FITC-labelled anti-human IgG Fc antibody (Sigma-Aldrich) diluted 1:500 for 30 min at 37°C. Prior to the preparation of EC-Fc/BNC or Fc/BNC, BNC was labelled with RITC (Sigma-Aldrich). Cells were incubated with RITC-labelled EC-Fc/BNC or Fc/BNC for 1, 4 and 8 hrs. The cells were washed with PBS and were fixed with 4% paraformaldehyde in PBS and then permeabilized with 0.1% TritonX-100 for 5 min. The fixed and permeabilized cells were blocked with 1% BSA in PBS and incubated with anti-ErbB2 antibody sc-06 (Santa Cruz Biotechnology, CA) for 1 hr followed by Alexa 488–labelled anti-rabbit IgG (Molecular Probe Inc., OR) for 30 min. After washing the cells with PBS, the visualization was carried under a confocal microscope, LSM 510 Meta (Carl Zeiss, Germany) equipped with Argon laser and LSM software (Carl Zeiss, Germany) with 2 nM (10 μg/ml) of EC-Fc/BNC for 5 hrs at 4°C or 37°C. After incubation, the cells were washed twice with ice-cold PBS to remove unbound BNC and then collected by the treatment with 0.025% trypsin. After centrifugation at 6,000 rpm for 1 min the supernatant was discarded and the cell pellet was washed twice with ice-cold PBS. The cells were then lysed in the lysis buffer, further incubated for 20 min at 4°C and sonicated twice. The cell extracts were clarified by centrifugation for 5 min at 4°C. Twenty microlitres of anti-HBsAg microbeads suspension was added to the extracts and incubated overnight at 4°C. After centrifugations for 30 sec at 4°C, the beads were washed three times in PBS, suspended in Laemnli buffer supplemented with β-mercaptoethanol, heated 5 min at 95°C, and subjected to SDS-PAGE followed by Western blotting. Transferin receptor internalization was taken as a control for the immunoprecipitation experiments because transferrin is constitutively replenished from the surface of the cell to the endosome and from recycling endosome to the cell surface.

Analysis of cellular uptake of EC-Fc/BNC

The receptor-mediated cellular uptake of EC-Fc/BNC was evaluated. The cells in 60-mm dish were washed twice with ice-cold PBS and incubated with 2 nM (10 μg/ml) of EC-Fc/BNC for 5 hrs at 4°C or 37°C. After incubation, the cells were washed twice with ice-cold PBS to remove unbound BNC and then collected by the treatment with 0.025% trypsin. After centrifugation at 6,000 rpm for 1 min the supernatant was discarded and the cell pellet was washed twice with ice-cold PBS. The cells were then lysed in the lysis buffer, further incubated for 20 min at 4°C and sonicated twice. The cell extracts were clarified by centrifugation for 5 min at 4°C. Twenty microlitres of anti-HBsAg microbeads suspension was added to the extracts and incubated overnight at 4°C. After centrifugations for 30 sec at 4°C, the beads were washed three times in PBS, suspended in Laemnli buffer supplemented with β-mercaptoethanol, heated 5 min at 95°C, and subjected to SDS-PAGE followed by Western blotting.

Flow cytometric analysis of ErbB2 internalization

For flow cytometric analysis, the cells seeded in a 100-mm dish were washed with PBS and were collected with enzyme free cell dissociation buffer (GIBCO) after incubating at 37°C for 5 min. The dissociation reaction was stopped by adding serum to the sample. Five hundred thousand cells were suspended in HBS (PBSB) supplemented with 1% BSA, containing 0.1% sodium azide in PBS (PBSB). After suspended in PBSB supplemented with EC-Fc, EC-Fc/BNC, Fc and Fc/BNC the cells were subsequently incubated on ice for 30 min. For internalization experiment, the cells were fixed and then permeabilized with 0.2% Triton X-100 in PBS. Then washed with PBSB, the cells were incubated with anti-human Fc antibody labelled with FITC at a dilution of 1:500 in PBSB followed by incubation on ice for 30 min. The cells were then processed for flow cytometric analysis by a FACS Calibur (Becton, Dickinson and Company).

Western blotting and image analysis

Proteins resolved on acrylamide gel were transferred to a polyvinylidene difluoride (PVDF) membrane (Bio-Rad, VA). The membrane was blocked using the JACoP plugin of ImageJ.

Biotinylation assay for the internalization of ErbB2

Biotinylation was performed as described previously [24]. Briefly, 5 × 10^5 cells were plated and cultured in a 60-mm tissue culture dish with complete medium. At the 90% confluency, the cells were washed twice in Hanks' balanced salt solution (HBSS) for 10 min at 4°C. Sulfo-NHS-SS-Biotin (Pierce Chemical) dissolved in HBSS at a concentration of 0.5 mg/ml was added to the cells at 4°C with mild shaking for 20 min. This reaction was repeated twice. The cells were washed with HEPES buffered RPMI supplemented with 1% BSA and 2 mM glutamine (RPMI-BSA) for 10 min at 4°C. Cells were incubated with either 30 μg/ml EC-Fc protein, or anti-ErbB2 monoclonal antibody sc-08 diluted to 1:100 in RPMI-BSA for 1 hr at 37°C. For the multivalent display studies the SK-BR-3 cells were incubated with 20, 100 and 200 nM of EC-Fc and 1, 5 and 10 nM of EC-Fc/BNC, where EC-Fc/BNC = 20.1. The cells incubated with anti-ErbB2 antibody sc-08 were washed and incubated for 1 hr at 37°C with HRP conjugated anti-mouse IgG (Cell Signaling Technology, MA) diluted to 1:400 with RPMI-BSA. The incubation was stopped by transferring the dishes back on ice, and the cells rinsed twice with HBSS. The biotin on the cell surface was cleaved under reducing condition of 20 mM DTT in 50 mM Tris-HCl, pH 8.7 containing 100 mM NaCl and 2.5 mM CaCl₂ for 20 min at 4°C, which was repeated twice. The cells were then washed three times with HBSS, and lysed in the lysis buffer consisting of 50 mM Tris-HCl, pH 7.4, 150 mM NaCl, 2 mM EDTA, 1% Triton X-100, 10 mM Naf, 1 mM vanadate, and protease inhibitor cocktail (Sigma-Aldrich) by the incubation for 20 min at 4°C. The lysates were collected and sonicated twice, and cell extracts were clarified by centrifugation for 5 min at 4°C. Protein concentrations in the extracts were determined by BCA assay (Pierce Chemical). Twenty microlitres of avidin agarose (Sigma-Aldrich) was added to the extracts and incubated overnight at 4°C. After centrifugations for 30 sec at 4°C, the agarose was washed three times in the lysis buffer, suspended in Laemnli buffer supplemented with β-mercaptoethanol, heated 5 min at 95°C, and subjected to SDS-PAGE followed by Western blotting. Transferin receptor internalization was taken as a control for the immunoprecipitation experiments because transferrin is constitutively replenished from the surface of the cell to the endosome and from recycling endosome to the cell surface.
with 10% skim milk in 10 mM Tris-HCl, pH 7.4, 150 mM NaCl containing 0.1% Tween-20 (TBST). The blots were then probed with a biotinylated polyclonal anti-human IgG antibody diluted to 1:1,000 (Dako, Denmark) or anti-preS-1 monoclonal antibody (Beacle, Japan) conjugated to HRP diluted to 1:15,000 in blocking condition, followed by a HRP-Avidin (Zymed, CA) in TBST containing 10% skim milk. The HRP signal was developed with Western Lightning™ and Western Lightning Plus Chemiluminescence Reagent (Perkin Elmer, MA) and the intensity of the bands was visualized by LAS image 4000 (Fujifilm, Japan). Quantitative assessments of the relative intensity of the blots were performed with NIH ImageJ.

Results

Preparation of EC-Fc protein

EC-Fc fusion protein was designed as EC-1 peptide fused to the amino terminus of IgG Fc domain (Fig. 1A). EC-Fc protein secreted from recombinant CHO cells were immunoprecipitated with protein A-agarose. The bound proteins were analysed by Western blotting with anti-human IgG to detect the Fc domain. As shown in Figure 1B, EC-Fc was observed as a monomer at approximately 30 kDa under reducing condition, whereas dimer at approximately 60 kDa was observed together with the tetramer, hexamer and higher order of oligomeric forms under non-reducing condition in either Coomassie Brilliant Blue (CBB) staining or Western blotting. The Fc domain without EC moiety prepared as a control was also observed as monomer at approximately 30 kDa under reducing condition, whereas only dimer formed as a result of the S-S-bond in the hinge region was observed at approximately 60 kDa under non-reducing condition. No further oligomeric forms were observed for Fc. These results suggested that the hinge region in the Fc domain should be effective to prepare dimeric form of Fc and EC-Fc proteins through disulfide bonds. The oligomeric forms observed as tetramer, hexamer and higher order can be because of the intermolecular disulfide bonds between cysteine residues in each EC-1 peptide. It is worthwhile noticing that no trimers were observed probably because extra-disulfide linkages were formed only between the EC-1 moieties of each EC-Fc proteins. The reduced protein in the CBB staining and Western blotting indicated an increment in the theoretical size of the artificial ligands, Fc and EC-Fc, which could be attributed to the N-linked glycosylation of the Fc portion of the human IgG. In this context, the oligomers should have two EC-1 moieties as correctly cyclized peptide through disulfide bonds in a molecule (Fig. 1C). The majority of EC-Fc was considered divalent form of EC-1 peptide even if the oligomeric forms such as hexamer and higher oligomer were formed because high oligomerization should precipitate EC-Fc and prevent it from working as ligand of ErbB2. For the susceptibility to the reducing conditions, EC-Fc was treated with DTT at various concentrations ranging from 0.5 to 5 mM at 4°C and 25°C (Fig. 1D). As the result, no order of disulfide bonds appeared to be present in the sensitivity to the reduction reagent in oligomeric forms because the reduction did not stop at a dimer even in low concentration of DTT. The specific binding affinity of EC-Fc to sErbB2 was evaluated by competition EIA (Fig. 1E). Scatchard plot revealed a high affinity binding site with a Kd of 26 nM, whereas a low affinity site was estimated to be 360 μM.

Internalization of ErbB2 was induced by EC-Fc in SK-BR-3 cells

Because the specific binding of EC-Fc to ErbB2 was confirmed, we then checked uptake of ErbB2 upon incubation with EC-Fc. For this assay, surface biotinylated cells were prepared and used. After incubation with EC-Fc, the biotin-labelled molecules retained on the cell surface were stripped off by the reducing reagent, whereas the molecules internalized were kept labelled with biotin so that they should be immunoprecipitated for the further analysis. The anti-ErbB2 antibody sc-08, which is known to induce internalization of ErbB2 in either SK-OV-3 cells or SK-BR-3 cells [13, 34], was used as a positive control. When evaluated on SK-BR-3 cells, EC-Fc was found to induce the internalization of ErbB2 (Fig. 2A). This internalization was estimated approximately 50% of that induced by anti-ErbB2 antibody sc-08. Internalized ErbB2 was observed when SK-OV-3 cells were treated with EC-Fc (Fig. 2B). In SK-OV-3 cells, the internalized ErbB2 induced by EC-Fc was almost equivalent with that induced by antibody sc-08. EC-Fc at 1 μM showed internalization of ErbB2 equivalent to that induced by anti-ErbB2 antibody sc-08 at 6.67 nM. This internalization was dependent on the doses of EC-Fc in a similar manner of monovalent EC-1 peptide fused to eGFP (EC-eGFP) [24]. When the biotinylated cells were treated with 5 μM EC-Fc, the internalized fraction of ErbB2 was increased up to three folds compared to that of the untreated cells at 37°C (Fig. 2C). We previously described that the internalization of ErbB2 was only observed in SK-OV-3 cells, when SK-BR-3 cells and SK-OV-3 cells were treated with EC-eGFP [24]. Taking the internalization demonstrated previously into consideration, this level of internalization was conceivable in SK-OV-3 cells, so that 5 μM EC-Fc should correspond to 10 μM EC-eGFP. The dimeric form of EC-Fc induced ErbB2 internalization whereas EC-eGFP treatment ErbB2 was retained on the surface of the SK-BR-3 cells without internalization.

Internalization of ErbB2 was enhanced by multivalent form of EC-Fc in SK-BR-3 cells

The multivalent form of EC-1 peptide was prepared exploiting the affinity of ZZ-BNC for IgG Fc region (Fig. 3A). When ZZ-BNC was mixed with EC-Fc, the ligand EC-1 was multivalently displayed on the surface of ZZ-BNC (EC-Fc/BNC). On the other hand, ZZ-BNC displaying null ligand was just prepared by mixing Fc protein with ZZ- BNC (Fc/BNC). First of all, binding capacity and binding efficiency of the ZZ-BNC with ligand was optimized and characterized. To optimize the ratio of Fc fusion molecule to BNC, FITC-labelled
Fig. 1 Construction and preparation of EC-Fc. (A) Schematic diagrams of EC-Fc. EC-1 peptide was fused to the human IgG Fc domain containing hinge region, so that the fusion protein EC-Fc should form dimer. The secretional expression of EC-Fc was designed with the signal peptide derived from human RNase I under the control of CMV promoter. The unique restriction enzyme sites Xba I, Nde I, Age I and Not I were used to construct the expression plasmid. (B) EC-Fc (theoretical MW = 30.0 kDa) or IgG Fc (theoretical MW = 27.6 kDa) was produced, purified with protein A-column and subjected to SDS-PAGE and Western blotting. Anti-human IgG Fc antibody recognized the purified EC-Fc without significant degradation. β-ME: betamercaptoethanol; CBB: Coomassie Brilliant Blue staining; WB: Western blotting. (C) Reduction of EC-Fc oligomeric form with DTT treatment. WB: Western blotting; M: monomer; D: dimer; T: tetramer; O: oligomer (B, C). Molecular weight markers range from 6.5 to 200 kDa (B, C). (D) Schematic representation of the probable dimer, tetramer and hexamer formation of EC-Fc and dimer formation of Fc through disulphide linkages. (E) Evaluation of the affinity of EC-Fc for ErbB2 by competitive EIA. Kd values between EC-Fc and sErbB2F were estimated from the Scatchard plots.
ZZ-BNC was mixed with Fc protein at variable molar ratios of 1:10, 1:20, 1:40, 1:60, 1:80 and 1:100, respectively, and the residual fluorescent intensity in the supernatant was measured. As shown in Figure 3B, the soluble Fc/BNC was reduced to 30% at a molar ratio of ZZ-BNC to Fc protein at 1:100 when judged from the fluorescence in the supernatant. Similarly, when human IgG was used, the soluble IgG/BNC was found reduced to 10% at a molar ratio of ZZ-BNC to IgG protein at 1:100. The amount of Fc protein bound to ZZ-BNC in the supernatant was further estimated from intensity of the band detected by Western blotting (Fig. 3C). As the result, the amount of Fc protein bound to ZZ-BNC was determined maximum when the molar ratio of ZZ-BNC to Fc protein was at 1:20. Thus, EC-Fc/ZZ-BNC was prepared by mixing ZZ-BNC and EC-Fc at the molar ratio of 1:20 for further experiments.

ErbB2 internalization mediated by EC-Fc/BNC was evaluated by the cellular uptake of EC-Fc/BNC (Fig. 3D–F). The SK-BR-3 cells were incubated with either EC-Fc/BNC or Fc/BNC for 5 hrs at 37°C or 4°C (Fig. 3D). As a result, EC-Fc/BNC showed significant cellular uptake at 37°C in SK-BR-3 cells. The EC-Fc/BNC cellular uptake into SK-BR-3 cells was found not only dose dependent in the range from 1 to 10 nM (Fig. 3E) and time course dependent up to 8 hrs at 2 nM (Fig. 3F). These results suggest that multivalent EC-1 peptide displayed on BNC might make its internalization into the cells efficient. The time course changes in the internalization of BNC in SK-BR-3 cells were evaluated by incubating cells with EC-Fc/BNC for various time periods under a confocal microscope (Fig. 3G). To demonstrate that observed localization or internalization of the receptor is not an artefact of fixation and cell permeabilization, we assessed the internalization of the EC-Fc/BNC with live cell imaging (Fig. S1). The colocalization of EC-Fc/BNC was confirmed by merging the location of BNC and EC-Fc, whereas Fc/BNC did not show any internalization. The internalization of ErbB2 induced by EC-Fc/BNC was further confirmed under confocal microscope (Fig. 3H). SK-BR-3 cells were incubated with RITC-labelled BNC displaying EC-Fc at 2 nM for 4 hrs. Then the cells were further probed for the ErbB2 by anti-ErbB2 monoclonal antibody sc-08 followed by Alexa 488–labelled secondary antibody. The internalized ErbB2 was observed to be colocalized with BNC.
together with BNC in SK-BR-3 whereas the Fc/BNC did not show any detectable internalization of ErbB2 as well as for BNC. As the result, the internalization of ErbB2 is concluded to be dependent on the uptake of BNC displaying EC-1 peptide multivalently. Internalization of EC-Fc/BNC was found to be dependent on both time course and ErbB2 in SK-BR-3 cells. Simultaneously, the internalization of ErbB2 was confirmed to be induced when EC-Fc/BNC was added. This observation implies that the cellular uptake of EC-Fc/BNC should be associated with the receptor-mediated endocytosis.

**Comparative study of the effect by EC-Fc and EC-Fc/BNC**

The specific binding of EC-1 ligand to cells overexpressing ErbB2 was assessed by immunostaining (Fig. 4A). Three cell lines MCF-7, SK-BR-3 and SK-OV-3 cells were selected and the binding of EC-Fc protein and EC-Fc/BNC was evaluated. The expression level ErbB2 was approximately $10^4$ sites/cell in MCF-7 [30], $10^5$ sites/cell in SK-BR-3 and SK-OV-3, respectively [31, 32]. The cells were incubated with EC-Fc or Fc or EC-Fc/BNC or Fc/BNC for 1 hr at 37°C followed by staining with anti-human IgG Fc-labelled with FITC. The fluorescence was observed only at the ligase of SK-BR-3 and SK-OV-3 cells but not with MCF-7 whereas Fc and Fc/BNC did not show any positive staining in all cell lines. No detectable fluorescence at the confocal microscopy to MCF-7 attributes to the low ErbB2 expression in the cells. The surface binding of the EC-Fc/BNC as well as EC-Fc was then qualitatively analysed by flow cytometry. As shown in Figure 4A in SK-BR-3 and SK-OV-3 cells, EC-Fc (green line) as well as EC-Fc/BNC (blue line) exhibited to bind to cell surface specifically. The binding efficiency of EC-Fc was found to be better than EC-Fc/BNC for SK-BR-3 cells. As the results, the specific binding of EC-Fc and EC-Fc/BNC to ErbB2 and overexpressing cells was successfully demonstrated.

ErbB2 internalization depending on the doses of EC-Fc and EC-Fc/BNC was assessed in the range of 20–200 nM in SK-BR-3 cells (Fig. 4B). ErbB2 internalization was observed with both EC-Fc and EC-Fc/BNC. When the intensity of the band analysed densitometrically showed the internalization of ErbB2 was more efficient by the stimulation for 2 hrs up to 100 nM of the ligand multivalently displayed on BNC than by EC-Fc ligand alone (Fig. 4C).

**EC-Fc/BNC internalized through GEEC pathway in SK-BR-3 cells**

Cells adopt divergent pathways for the endocytosis of the cargos and receptors. The key pathways were categorized into clathrin-dependent and clathrin-independent mechanisms [33]. The clathrin-independent pathway is further classified into caveolar and GPI-anchored early endocytic compartments (GEEC) pathways [33, 34]. ErbB2 is thought to be internalizing in SK-OV-3 through clathrin-mediated internalization [24]. In this study we found that the mechanism of internalization of ErbB2 present in SK-OV-3 cells is deficient in SK-BR-3 cells. We tried to identify the difference between the two cell lines using DNA microarray (Fig. S2). As a result the expression of claudin 16, caveolin-1 and caveolin-2 were found to be extensively downregulated in SK-BR-3 cells when compared to SK-OV-3 cells. This absence of caveolin-1 in SK-BR-3 cells was further confirmed by immunostaining using anti-Cav1 antibody (Fig. 5A). SK-BR-3 did not show the presence of caveolin-1 suggesting that the impaired caveolar mechanism prevailing in the cell, which was consistent with the finding by the previous reports [13, 35]. Because caveolin-1 was detected in SK-OV-3 cells the internalization pathway deficient in SK-BR-3 cells might be attributed to caveolae.

The mechanistic approach for the uptake of the EC-Fc/BNC was assessed in SK-BR-3 cells using inhibitors for other pathways. SK-BR-3 cells were treated with EC-Fc/BNC in the presence or absence of inhibitors for clathrin, and GEEC pathway and the internalization was assessed. When the cells were treated with 100 nM CPZ, an amphiphilic drug, which inhibits the clathrin-mediated pathway, the internalization of the EC-Fc/BNC was unaffected (Fig. 5B).
Most of the cargos irrespective of their route merges with Rab5 and early endosome antigen-1 (EEA-1) enriched in early endosomes, which is further sorted into various intracellular destinations [36, 37]. The colocalization of EC-Fc/BNC with EEA-1 was assessed with anti–EEA-1 antibody under confocal microscope at various time periods starting from 5, 30 and 90 min and the colocalization was found to be maximum at 30 min (Fig. S3).

When the cells were treated with 5 mM mCD, which dislodges the cholesterol from the surface and inhibit the cdc42 activity, for 30 min the internalization was blocked and the cells untreated showed considerable uptake of EC-Fc/BNC. The bound fractions of EC-Fc/BNC were treated with and without acid stripping to ensure the colocalized fraction inside the cells (Fig. 5C). The colocalization was calculated to more than 60%, which was reduced in the internalization of EC-Fc/BNC with the mCD treatment. To assess the surface binding of the EC-Fc/BNC after the inhibitor treatment the cells were stained for the ErbB2 and EC-Fc/BNC at various concentration of the mCD (Fig. S4). The inhibitor treatment did not deteriorate the binding affinity of the EC-Fc/BNC to ErbB2. These results were further confirmed with flow cytometric studies (Fig. S5).

**Discussion**

The EC-Fc was designed as dimer due to the hinge region in IgG Fc domain (Fig. 1A). However, we found higher oligomeric forms such as tetramer and hexamer. These higher order forms were supposed to be divalent as described in ‘Results’ and Figure 1C. Further multivalent form of EC-1 peptide was prepared by displaying EC-Fc on BNC which was optimized to a ratio of EC-Fc to BNC = 1:20. These divalent and multivalent forms of EC-Fc induced internalization of ErbB2 into SK-BR-3 cells whereas...
The internalization of ErbB2 was found dependent on the concentration in the range of 20–200 nM for EC-Fc and appeared to be enhanced when 20–100 nM EC-Fc was displayed on BNC (Fig. 4B). EC-Fc/BNC appeared more efficient in inducing the internalization of ErbB2 when compared to EC-Fc ligand alone. The time course dependence of the BNC internalization was found to be also in the same range of EC-Fc/BNC concentration. The binding affinity of the EC-Fc was estimated to be about 26 nM whereas monovalent EC-1 peptide as EC-eGFP fusion showed 1 M (Table 1). The affinity of the multivalent display for ErbB2 was 33 nM, which did not ameliorate the affinity of EC-Fc for the ErbB2. Therefore, the affinity should elucidate the enhanced internalization of ErbB2 induced by EC-Fc/BNC in SK-BR-3 cells. It is believed that the heterodimerization between ErbB2 and ErbB3 is one of the reasons for retention of ErbB2 on the cell surface, so that monovalent form of the EC-1 peptide could not induce internalization of ErbB2 in SK-BR-3 cells. Shuttling between homodimers and heterodimers should increase depending on the distance of the dimers. EC-Fc supposedly keeps the receptors close enough to facilitate homodimer formation, which will allow the internalization of ErbB2. Therefore, the cells treated with EC-Fc/BNC showed considerable increment in the internalization of

Table 1 Affinity for ErbB2 of EC-1 peptide in various forms and internalization of ErbB2 in SK-OV-3 and SK-BR-3 cells

| Forms of EC-1 peptide | EC-eGFP | EC-Fc | EC-Fc/BNC |
|-----------------------|---------|-------|-----------|
| Kd (nM)               | 1.0 x 10** | 2.6 x 10 | 3.3 x 10 |

Internalization of ErbB2 observed in

| SK-OV-3 cells | SK-BR-3 cells |
|---------------|---------------|
| +1            | +5            |
| ++            | +             |
| +             | ++            |

1Internalization was observed in the range of μM orders of ligands.

2Internalization was observed in the range of nM orders of ligands.

3Internalization was not observed.

monovalent form of EC-1 peptide did not induce internalization [24]. The internalization of ErbB2 was found dependent on the concentration in the range of 20–200 nM for EC-Fc and appeared to be enhanced when 20–100 nM EC-Fc was displayed on BNC (Fig. 4B). EC-Fc/BNC appeared more efficient in inducing the internalization of ErbB2 when compared to EC-Fc ligand alone. The time course dependence of the BNC internalization was found to be also in the same range of EC-Fc/BNC concentration. The binding affinity of the EC-Fc was estimated to be about 26 nM whereas monovalent EC-1 peptide as EC-eGFP fusion showed 1 M (Table 1). The affinity of the multivalent display for ErbB2 was 33 nM, which did not ameliorate the affinity of EC-Fc for the ErbB2. Therefore, the affinity should elucidate the enhanced internalization of ErbB2 induced by EC-Fc/BNC in SK-BR-3 cells. It is believed that the heterodimerization between ErbB2 and ErbB3 is one of the reasons for retention of ErbB2 on the cell surface, so that monovalent form of the EC-1 peptide could not induce internalization of ErbB2 in SK-BR-3 cells. Shuttling between homodimers and heterodimers should increase depending on the distance of the dimers. EC-Fc supposedly keeps the receptors close enough to facilitate homodimer formation, which will allow the internalization of ErbB2. Therefore, the cells treated with EC-Fc/BNC showed considerable increment in the internalization of

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the BNC, although the effect of EC-Fc moiety appeared conceivable because Fc/BNC did not show significant internalization of ErbB2 as the background (Fig. 3D–F).

It is thought that the processing and degradation of the ErbB2 is different in various cell lines [24, 34, 35]. The CPZ at 100 nM, which inhibits the formation of clathrin-coated pits, inhibited the internalization of ErbB2 by EC-1 peptide fused to eGFP in SK-OV-3 [24]. On the contrary, the same concentration of CPZ did not block the internalization of ErbB2 in SK-BR-3 cells. In this study, we confirmed the absence of caveolin-1 in SK-BR-3 cells using microarray and immunostaining where it showed around 300-fold increment in the expression of Cav-1 in SK-OV-3 cells (Fig. S1). Recent studies support our observation that the SK-BR-3 cells lack caveolae [13, 35]. ErbB2 internalization after geldanamycin treatment showed GEEC or clathrin-independent compartments (CLIC) pathway, which ultimately fuses with the classical pathway for the degradation in lysosomes, is independent of caveolar and clathrin pathway [35].

In this study we employed mpCD, a cholesterol dislodging oligosaccharide, for deciphering the mechanism behind the internalization of the EC-Fc/BNC in SK-BR-3 cells, as an inhibitor for the blockade of GEEC pathway. Cholesterol depletion with mpCD inhibits both caveolar [38] and GEEC pathway [39, 40]. Earlier it was shown that cholesterol level determines the internalization of GPI-anchored proteins and with clostridium difficile toxin B [34, 39]. Because SK-BR-3 cells lack caveolar machinery, the blockade with mpCD will only correspond its effect in blocking the internalization of GEEC-mediated mechanism. The internalization of ErbB2 is sensitive to the depletion of cholesterol, which is essential for most of the clathrin-independent pathways. GEECs are formed as compartments free of clathrin, caveolin and dynamin, which are short lived (2–5 min) and ultimately fuses with EEA-1 enriched early endosome [33, 38]. It was shown that cdc42 is responsible for the dynamin-dependent and clathrin-independent internalization of GPI-anchored proteins. It was also reported earlier that cdc42 activation recruits actin polymerization machinery rendering this pathway sensitive to inhibitors of cholesterol depletion and actin polymerization [39]. This pathway specifies the visible difference in the uptake of the EC-Fc/BNC between SK-OV-3 and SK-BR-3 cells.

Another important molecule, which plays a key role in the GEEC pathway, is EEA-1 that contains a FYVE finger, which interacts with PI3K [41, 42]. PI3K phosphorylates Rab5 that helps the EEA-1 to localize to early endocytic compartments [36]. The colocalization of EC-Fc/BNC with EEA-1, which corresponds the preliminary step in the endosomal pathways before transferring it to the sorting endosomes, was observed (Fig. S2). Because both caveolar and clathrin pathways are impaired in SK-BR-3 cells as described earlier, the colocalization should correspond to GEEC pathway.

In this study, we revealed the presence of GEEC pathway as ErbB2 internalization, which critically differs between SK-BR-3 cells and SK-OV-3 cells using EC-1 peptide in various forms. As for the nanoparticle internalization, the pathways of clathrin and caveolar-independent endocytosis have not been described until now [43]. The internalization of the carrier particle was described repeatedly on folate receptor (FR) tagged nanoparticles [44]. In these reports, mechanism of the carrier internalization emphasizes the FR internalization through GEEC pathway. Similarly, EC-Fc should utilize the mechanism of internalization of ErbB2 in this study even when they are displayed on BNC (Fig. 4B).

It is considered that ZZ-BNC is an efficient nano-machine for molecular targeting in drug delivery system displaying antibodies on the surface [25, 29, 45]. Multivalency of EC-Fc was successfully demonstrated to target ErbB2 enhancing the internalization of ErbB2, in this report. In this context, delivery of therapeutic substances into specific cells of various diseases including cancer can be feasible if EC-Fc/BNC is used as a drug delivery system. Multivalent display strategy proposed here should be useful for molecular targeting to introduce therapeutic drug into the cells followed by the internalization of receptors.

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Conflict of interest

The authors report no conflicts of interest. The authors alone are responsible for the content and writing of the paper.

Supporting information

Additional Supporting Information may be found in the online version of this article:

Fig. S1 Live cell imaging. For live cell imaging, the SK-BR-3 cells were cultured in 18 mm coverslips and washed twice with phenol red free RPMI (Sigma) containing 1 % glucose, 25 mM HEPES and 1 % BSA supplemented with 100 U/ml penicillin and 100 µg/ml streptomycin. The EC-Fc BNC tagged with RITC was incubated on ice for 30 min prior to the visualization in order to enable the binding of the multivalent forms of BNC to the cell surface. The cells were then washed thrice with warm media and were visualized in 100X 1.3 N.A. oil immersion objective Olympus IX81 inverted microscope (Olympus) with temperature control followed by data acquisition with Metamorph software (Molecular Devices). The imaging was carried out with DP71 cooled CCD camera attached to the microscope and the fluorescence of RITC was acquired along with the DIC field image. The time-lapse imaging
Fig. S2 Microarray analysis. Expression of genes in SK-BR-3 cells was analyzed by microarray procedure. Total RNA preparation and analysis was performed as described previously [46, 47]. DNA microarray was carrying 1,795-oligonucleotide probes for human cell surface proteins. cDNAs were synthesized with Superscript II reverse transcriptase (Invitrogen) with oligo dT primers. Aminomethyl-dUTP was incorporated into cDNAs followed by coupling with Cy-3 dye (Aambion, TX, USA) and were processed for hybridization at 55°C for 15 hours. The fluorescent images for the hybridization were captured using FLA8000 scanner (Fuji Film, Japan) and analyzed with GenePix Pro5.1 software (Axon Instruments, CA). We found caveolins, Cav-1 and 2, and claudin 16 (CLDN-16) were downregulated in SK-BR-3 cells when compared to SK-OV-3 cells while Erb2B and GAPDH expression were equally expressed in both the cell lines. RT-PCR (Fig. S1A) and qRT-PCR (Fig. S1B) further confirmed these results. The conditions for the RT-PCR are as follows: 94°C for 5min; followed by 30 cycles of 94°C for 30 sec, 55°C for 30 sec, 72°C for 30 sec and 72°C for 7 min. Quantitative real time PCR was performed with SYBR Green Realtime Master Mix (Toyobo) in triplicates containing 5ng of cDNA along with 400 nM primers using LightCycler™ (Roche). The thermal cycling condition was as follows: 95°C for 1 min followed by 40 cycles of 95°C -10 sec, 55°C for 10 sec, 72°C for 25 sec and 60°C for 1 min. The following sets of the primers were used for the PCR reaction. Claudin-16, (Forward) 5'-GCTTGGCACATGAGGGATCT-3' (Reverse) 5'-TCGATCTGACATCTCTAC-3'; Cav-1 (Forward) 5'-GACTCGGAGGACATCTCCTAC-3'; (Reverse) 5'-GTTGATGCGGACATTGCTGA-3'; Cav-2, (Forward) 5'-ACGTACAGCTCTTCATGGAC-3' (Reverse) 5'-CAAGTTGGAGGCTGACAGAAG-3'.

Fig. S3 Time dependent colocalization of EC-Fc BNC in early endosomes. Colocalization of EC-Fc/BNC-RITC with EEA-1 was found in the early endosomes. SK-BR-3 cells were treated with EC-Fc/BNC-RITC for 5, 30, 90 min, and the colocalization of the EC-Fc/BNC-RITC with the early endosomal marker, EEA-1 was assessed with anti EEA-1 antibody (BD Biosciences) and the secondary antibody labeled with AlexaFlour-488 (Molecular Probes). Error bars indicate standard error. JACoP plugin in Image J was employed for percentage colocalization assessment. Percentage colocalization was found to be maximum up to approximately 70% by 30 min. The data represented here is the representative of two independent experiments.

Fig. S4 Binding of EC-Fc BNC to the cell surface after treatment with varying concentration of mβCD. The binding of EC-Fc/BNC's to ErbB2, in the presence of 2 to 10 mM of cholesterol dislodging oligosaccharide-mβCD, was not affected. SK-BR-3 cells were treated with 5 and 10 mM of mβCD for 30 minutes followed by treatment with EC-Fc/BNC labeled with RITC are shown in Figure S2. The cells were stained with anti ErbB2 antibody (green) and red spots indicate EC-Fc/BNC labeled with RITC. Bars depict 10 µm.

Fig. S5 FACS analysis for the surface binding and internalization of EC-Fc BNC with inhibitor treatment. Surface binding and internalization of EC-Fc/BNC labeled with FITC in SK-BR-3 cells was quantified using FACS analysis. First, the binding of EC-Fc/BNC on SK-BR-3 cells was treated with or without 5 mM of mβCD on ice for 30 min. The cells were then fixed and analyzed with FACS to determine the surface bound fraction (Fig. S5A). The histograms showed overlapping of the peak tops indicating that the surface binding is majorly unaffected by the mβCD treatment. The peak (blue), which corresponds to approximately 20% of the total cell counts, orientating towards the untreated cells (grey peak) indicates the presence of a different population of cells that are incapable of binding to EC-Fc/BNC after mβCD treatment. Simultaneously, another set cells were treated with EC-Fc/BNC labeled with FITC in the presence or absence of 5 mM of mβCD at 37°C for 30 minutes. Cells were then trypsinized to remove the surface fraction, fixed and permeabilized, and the internalized fraction was analyzed adopting the same procedure with FACS (Fig. S5B, C). mβCD treated SK-BR-3 cells showed a peak point in the histogram which overlaid the peak area of untreated cells indicating the mβCD inhibits the internalization, while the cells without mβCD treatment drifted from the untreated cells towards the right. Total intensity of the internalized fraction was assessed from the FACS data were plotted on to a graph (Fig. S5). mβCD suppressed the internalization to approximately 50% when compared to the absence of mβCD treatment. M: mβCD; T: trypsin. Error bars indicate standard error.

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