Overexpression of the receptor tyrosine kinase HER2 plays a critical role in the development of various tumors. Biparatopic designed ankyrin repeat proteins (bipDARPins) potently induce apoptosis in HER2-addicted breast cancer cell lines. Here, we have investigated how the spatiotemporal receptor organization at the cell surface is modulated by these agents and is distinguished from other molecules, which do not elicit apoptosis. Binding of conventional antibodies is accompanied by moderate reduction of receptor mobility, in agreement with HER2 being dimerized by the bivalent IgG. In contrast, the most potent apoptosis-inducing bipDARPins lead to a dramatic arrest of HER2. Dual-color single-molecule tracking revealed that the HER2 “lockdown” by these bipDARPins is caused by the formation of HER2-DARPin oligomer chains, which are trapped in nanoscopic membrane domains. Our findings establish that efficient neutralization of receptor tyrosine kinase signaling can be achieved through intermolecular bipDARPin crosslinking alone, resulting in inactivated, locked-down bipDARPin-HER2 complexes.
Human epidermal growth factor receptor 2 (HER2/ErbB2/neu) has gained tremendous importance as a biomarker in diagnostics and a target in the therapy of breast cancers, but also in gastric and gastroesophageal cancers, over the past decade. This receptor tyrosine kinase has a distinct role within the ErbB family of receptors, in that it has no ligand. Derived from the rigidity seen in all crystal structures, in molecular dynamics, and binding experiments, HER2 has therefore been assumed to always be in a dimerization-competent state. Under physiological conditions, HER2 likely does not homodimerize, and thus its role is restricted to that of a signal-amplifying co-receptor for other, ligand-activated members of its family. Upon malignant overexpression, however, spontaneous formation of homodimers and heterodimers, even in the absence of ligands of the other ErbB family member, enables sustained proliferation and survival signaling. This stimulus, in turn, frequently transforms the cancer cell signaling architecture towards singular dependence on HER2 signaling. Such HER2-addicted cancers are more aggressive but, at the same time, also susceptible to targeted therapies. Therefore, HER2 overexpression, previously merely a correlate of bad prognosis, nowadays generally also implies the availability of several targeted treatment options.

Among these, monoclonal antibodies (mAbs) have been successfully developed that bind to the HER2 extracellular domain and thus block activity. Prominent examples that have been approved for therapy are the humanized mouse mAbs trastuzumab (TZB) and pertuzumab (PZB). Interestingly, the mechanisms of HER2 inhibition by TZB and PZB are complementary (targeting complexes with the non-liganded and liganded states of HER3, respectively), and therefore most efficacious therapies employ the combination of both mAbs. Of note, subtle changes in the relative orientation of the Fab domains can transform F(ab')2-like molecules derived from TZB into active pro-proliferative agents, underlining the importance of binding geometry for the modulation of HER2 activity by affinity reagents.

Recently, we have reported a novel strategy based on designed ankyrin repeat proteins (DARPins) to inhibit HER2 activity in breast cancer cells. To this end, we engineered biparatropic antitumor DARPin (bipDARPins) consisting of two binding moieties, which recognize the extracellular subdomains I and IV of HER2, respectively (Fig. 1a). These binding moieties are connected by a short linker (see Supplementary Fig. 1 for a detailed overview), with the aim to trap and stabilize an inactive conformation. Compared to TZB and PZB, bipDARPins much more efficaciously promote apoptosis in HER2-dependent tumor cell lines by dephosphorylating HER2 (and not only HER3) and subsequently preventing re-activation of phosphoinositide-3 kinase through RAS. While the signaling pathways underlying this effect have been characterized in detail, the exact mechanism responsible for the potent inhibitory function of bipDARPins, in contrast to less active molecules, has so far remained largely unclear. Less active molecules not only include the antibodies TZB and PZB but also the same DARPin units with longer linker in between or in different orientation.

X-ray structures of single DARPin moieties in complex with the isolated, soluble extracellular subdomains of HER2 led to the conclusion that the most active bipDARPins cannot bind both epitopes in an intramolecular fashion, implying that rather two neighboring HER2 molecules are engaged (Fig. 1c, d). In contrast, intramolecular trapping would be possible for the less active molecules with longer linkers. Possible mechanisms of action considered had initially included locking an inactive HER2 conformation similar to the tethered conformation seen in the other ErbB receptors, but we have shown subsequently that the extracellular domain is very rigid, making such a conformation highly unlikely. This suggests that inhibition is instead achieved via restrained dimerization (Fig. 1c, d), in which interactions between the cytosolic tyrosine kinase domains required for allosteric activation are prevented. However, this binding mode also may allow elongation into daisy chain-like arrangements (Fig. 1d), which we have previously proposed to cause efficient inhibition of HER2 signaling.

To delineate the mechanism responsible for HER2 inhibition by bipDARPins, we explored here their effects on the spatiotemporal organization and dynamics of HER2 in the plasma membrane. Chemical crosslinking in combination with immunoprecipitation revealed oligomerization induced by active bipDARPins, but not by TZB and PZB. Making use of cell surface-specific post-translational labeling techniques, we quantified the diffusion properties of HER2 in the plasma membrane of living cells by fluorescence recovery after photobleaching (FRAP) and by single-molecule localization microscopy (SMLM). Efficient arrest of

Fig. 1 Concept of bipDARPins and possible modes of binding to HER2. a Monovalent, monoparatopic DARPin (9.26 binding to extracellular subdomain I, vermilion; G3 binding to extracellular subdomain IV, yellow) and design of bipDARPin 6L1G (right), connecting 9.26 and G3 through a short linker. b Ligand-activated (e.g., by heregulin, mint) HER2:HER3 (sky blue, blue) heterodimers are able to stimulate proliferation and survival. c, d Bivalent intermolecular binding of bipDARPins may induce the formation of inactive dimers or oligomerization into daisy chains. In both cases, the kinase domains are separated and thus inactive. HRG heregulin, TM transmembrane helix, KD kinase domain.
HER2 diffusion was observed upon treatment with bipDARPins, while control experiments with TZB and PZB yielded only a slight reduction in mobility. Dual-color single-molecule tracking (SMT) and co-tracking demonstrate that bipDARPins induce clustering and immobilization of HER2 at the cell surface. Detailed analyses of local diffusion properties uncovered transient nanoscale immobilization, potentially caused by entrapment in plasma membrane subcompartments. These findings presented here strongly suggest that the signaling-incompetent states of HER2, induced by bipDARPin binding, are signaling-inactive oligomers formed directly through crosslinking. In addicted cell lines, this state leads to pan-HER inhibition.

**Results**

**Chemical crosslinking suggests that bipDARPins induce HER2 oligomer formation.** To characterize the stoichiometries of HER2 complexes formed with bipDARPins, we performed chemical crosslinking analyses with bis[sulfosuccinimidyl]suberate (BS3) using BT474 cells, a model for HER2-addicted breast cancer10 with a highly elevated expression level of approximately 9×10^5 copies of HER2 per cell13. In the absence of treatment, as well as after addition of the HER3-binding ligand HRG, we primarily detected non-crosslinked (monomeric) HER2, and only minor nonspecific crosslinking adducts (Fig. 2a). As expected, the homo-bivalent monoparatopic control DARPin GL4G (Supplementary Fig. 1) led to a band at the expected size of two cross-linked HER2 molecules. Remarkably, the high-potency bipDARPIn 6L1G not only yielded a similar band but also an additional band staining for both HER2 and DARPin at a size corresponding to complexes of 3-4 HER2 molecules. Control treatments, either with the monovalent moieties contained within 6L1G, DARPins 9.26 and G3, the combination of these monovalent DARPins, or with the non-HER2-binding control DARPin OL1O (that binds bacterial maltose binding protein) did not alter the band pattern observed for HER2 alone.

We also investigated HER2 crosslinking with BS3 after incubation with TZB or PZB (Fig. 2b). As previously described14, TZB seemed to be preferably crosslinked, forming a ternary complex with two HER2 molecules, running at a higher apparent size than the presumed HER2 dimer without TZB. In contrast, PZB seemed not to lead to an increase in crosslinked species that would contain two or more HER2 molecules. Instead, HER2 and PZB appeared to be preferably interconnected in a 1:1 ratio. Together, these results suggested that bipDARPins not only dimerized HER2 in an inactive state but also induced daisy chain formation as depicted in Fig. 1f.

**FRAP measurements reveal immobilization of HER2 by bipDARPins.** To further explore bipDARPin-induced oligomerization of HER2 suggested by these crosslinking experiments, we probed receptor diffusion dynamics at the surface of live cells by single-molecule and ensemble fluorescence imaging techniques. To ensure selective labeling of proteins at the cell surface with photostable fluorophores, we fused HER2 to an N-terminal HaloTag15 (HT-HER2), which was reacted with membrane-impermeable, negatively charged dyes conjugated to the HaloTag ligand (HTL). Thus, the variation of labeling density and optimal dual-color labeling was readily achieved by adjusting the labeling conditions, which allowed us to apply and compare different imaging techniques using the same HER2 expression construct and total cell-surface expression levels. As we included a long, flexible linker between the receptor’s N-terminus and the HaloTag, interference with any receptor function is minimized16.

To obtain an overall picture of receptor mobility in the plasma membrane, we used the classic FRAP technique17, with densely labeled HT-HER2. For this purpose, we stably transfected HEK293-derived T-Rex cells, which are based on the Flip-In system for inducible overexpression, with HT-HER2. To exclude a bias from receptor endocytosis and recycling, FRAP experiments were typically performed under conditions of ATP depletion18. Photobleaching was applied to a circular spot of 2 μm in diameter. We measured in the equatorial plane rather than at the basal membrane19, because a better signal-to-noise ratio could be achieved due to spatial integration over more molecules in the z-dimension (Fig. 3a, b). Photobleaching was performed in a single scan, yielding a decrease by 40–60% compared to the pre-bleach fluorescence intensity, while photobleaching during subsequent acquisition of fluorescence recovery was negligible. As the resulting distorted geometry of the photobleached volume within the membrane precludes application of established analytical models for fitting the recovery curves20,21, we directly compared the FRAP curves obtained under different conditions, as suggested elsewhere19.

In the absence of treatment, we consistently observed only 40–50% maximum recovery of photobleached HER2 within the observation time of 30–50 s that was sufficient to reach a plateau of the FRAP curve (Figs. 3–5). Strikingly, dramatic reduction of HER2 mobility was observed upon treatment with active bipDARPins, with maximum recovery levels of <10% reached within the observation time (Fig. 3a, c). This “lockdown” of HER2 mobility by bipDARPins was also observed for FRAP experiments at the basal cell membrane (Fig. 3b, c). The homo-bivalent antibodies TZB and PZB, which presumably dimerize HER2, each reduced the rate and maximum level of HER2 recovery by ~50%.

![Fig. 2 Oligomerization of HER2 (~180 kDa) by bipDARPins and antibodies on BT474 cells analyzed by chemical crosslinking and two-color western blot.](image-url)

- **a** Comparison of the size of HER2 complexes after treatment with the bipDARPin 6L1G (32 kDa) with those obtained with growth factor heregulin (HRG, 27 kDa), monovalent DARPin (9.26 and G3, ~12 kDa each) or a bivalent, monoparatopic DARPin (GL4G, ~30 kDa). Primary antibodies against HER2 and DARPins were used. **b** Crosslinked HER2 complexes upon treatment with antibodies trastuzumab (TZB, ~145 kDa) and pertuzumab (PZB, ~145 kDa), detected by primary antibodies against HER2 and human IgG.

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Remarkably, the combination of TZB and PZB also induced complete HER2 immobilization (Fig. 3d), in line with oligomerization that can be caused by alternating crosslinking with two mAbs of non-overlapping epitopes19.

Efficient lockdown of endogenous HER2 by bipDARPins in breast cancer cells. To confirm the biological relevance of bipDARPin-induced lockdown of HER2, we employed the small (6 kDa) affibody ZHER2 (ref. 22) for labeling endogenous, untagged HER2 in breast cancer cells. ZHER2 binds epitopes that are non-overlapping with any of the used DARPins or antibodies10,11,23, thus ensuring minimum interference. We applied ZHER2 for probing the receptor dynamics at the surface of BT-474 cells. FRAP experiments confirmed similar diffusion dynamics of ZHER2-labeled HER2 in the plasma membrane of BT-474 cells as observed for HT-HER2 in TREx-HEK293 cells (Fig. 4a). Strikingly, strong lockdown of HER2 upon treatment with the biologically most active DARPin 6L1G was confirmed for the endogenous receptor (Fig. 4a), in a clearly dose-dependent manner (Fig. 4b).

In comparison, the monoparatopic, homo-bivalent controls 6L46 and GL4G both led to a similar, moderate reduction in mobility when applied individually. By contrast, a mixture of these two binders abolished HER2 mobility completely, in agreement with receptor oligomerization by crosslinking (Fig. 4b). Similarly, as seen for HT-HER2 on TREx-HEK cells, TZB or PZB slightly reduced the mobility of HER2, while their combination lead to immobilization. Thus, the HER2 complexes formed upon binding of the single bipDARPin 6L1G resemble those resulting from mixtures of two homo-bivalent, monoparatopic agents (DARPins or antibodies) with non-overlapping epitopes, rather than the single agents.

BipDARPin-HER2 complex formation suffices for immobilization. HER2 immobilization occurred rapidly upon bipDARPin binding, as illustrated by several consecutive FRAP measurements before and after 6L1G addition (Fig. 5a). No significant increase in HER2 mobility was observed even 6 h after treatment (Fig. 5b), demonstrating that immobilization is not a transient effect based on an immediate cellular response to DARPin binding, but that it persists as long as 6L1G is present. This suggests that bipDARPins and HER2 rapidly (within the dead time of the experiment, ~160 s) form complexes, which remain at the cell surface for extended periods, in line with the long internalization half-life of HER2 (ref. 24) and stable surface and internal HER2 levels upon 6L1G treatment25.

To test whether the cortical membrane cytoskeleton contributed to HER2 lockdown, we performed FRAP experiments in the presence of Latrunculin B (LatB), which depolymerizes actin. However, LatB treatment neither affected HER2 mobility in untreated cells nor when added before or after treatment with 6L1G (Fig. 5c). Furthermore, treatment with a HER2-specific kinase inhibitor neither impacted the mobility of HER2 alone nor the ability of 6L1G to cause HER2 lockdown (Fig. 5d). This confirms that phosphorylation is not involved in HER2 immobilization. To test whether the high baseline fraction of intrinsically immobile HER2 was restricted to cell lines with extremely strong HER2 expression, we compared the results from BT-474 breast cancer cells (which strongly overexpress HER2) to those obtained from our TREx-HEK293 cell line without induction of HER2.
overexpression, which results in intermediate expression levels (Supplementary Table 1 and Supplementary Fig. 2). The recovery curves appear virtually identical, again showing ~40% of HER2 being immobile on a ~50 s timescale (Fig. 5e).

**BipDARPins induce immobilization of HER2 at the single-molecule level.** The strong loss of HER2 mobility identified by the FRAP experiments clearly supported our model that bipDARPins induce HER2 clustering into daisy chains. To further investigate the local diffusion properties of HER2, we used single-molecule techniques. First, we probed fluorescence fluctuations within a diffraction-limited confocal volume positioned on the apical plasma membrane, using low-density dual-color labeling of HT-HER2 with Cy3 and AlexaFluor 647 (AF647), respectively. In the absence of bipDARPins, distinct fluorescence bursts originating from fluorescently labeled HER2 molecules diffusing through the focus were observed after an initial bleaching phase to reach a steady state (Supplementary Fig. 3a, b). Strikingly, the number of observed bursts dramatically decreased after adding the biologically active bipDARPin 6L1G but not after adding the monovalent control DARPin OL1G (Supplementary Fig. 3b). Quantitative burst detection confirmed the loss of HER2 mobility upon treatment with bipDARPins (Supplementary Fig. 3c).

The strongly reduced number of bursts suggested that HER2 lockdown by bipDARPins reduces mobility at length scales on the order of the size of the confocal volume. To explore the spatiotemporal re-organization of HER2 by bipDARPins with sub-diffraction resolution and to directly visualize immobile HER2-DARPin complexes, we turned to SMLM. For this purpose, we transiently expressed HER2 fused to an N-terminal SNAP2 tag in HeLa cells. After simultaneous labeling with DY-549P1 and DY-649P1, we performed time-lapse dual-color total internal reflection fluorescence microscopy (TIRFM) for tracking and co-tracking individual HER2 in the basal plasma membrane of living cells. To ensure reliable single-molecule localization and tracking, cells with an observable particle density <1 µm⁻² in each channel were selected (see below). Given an estimated labeling degree of 40% for the SNAP2 tag and the expected level of endogenous HER2 in HeLa cells (Supplementary Table 1 and Supplementary Fig. 2), the fraction of labeled HER2 was <10% in these experiments.

To assess the spatiotemporal organization and dynamics of HER2, super-resolution images were rendered from single-molecule localizations obtained within 500 consecutive frames. These images revealed a strikingly speckled appearance for cells treated with the active bipDARPin 6L1G, compared to a monovalent control (Supplementary Movie 1, Supplementary Movie 2, and Fig. 6a). Since negligible photobleaching or photoblinking can be assumed during the observation time, the observed characteristics for HER2 in the presence of the active bipDARPin can be attributed to a dramatic loss in mobility, leading to repeated detection of the same molecule at the same location over time. By contrast, a much more spatially homogeneous distribution is observed in the negative control experiment in cells treated with the monovalent DARPin OL4G. Only few speckles were observed under these conditions, indicating largely unhindered diffusion.

We therefore analyzed in more detail the diffusion properties of HER2 molecules after treatment with different DARPins and mAbs, by SMT performed at comparable particle density to minimize bias (Fig. 6b, Supplementary Fig. 6b, and Supplementary Table 2). Immobile molecules were identified by a spatiotemporal clustering algorithm to quantify the immobile fraction (Fig. 6c and Supplementary Table 2). For the mobile fraction, an average diffusion constant was determined by mean squared displacement (MSD) analysis (Fig. 6c) and Supplementary Fig. 6d). The concentration dependence of the mobility reduction as illustrated by the relative fluorescence recovery after 48 s. 

![Fig. 4 Mobility changes of endogenous HER2 on the surface of BT-474 cells (labeled via the affibody ZHER2) upon treatment with DARPins and mAbs as quantified by FRAP.](image-url)

**Fig. 4 Mobility changes of endogenous HER2 on the surface of BT-474 cells (labeled via the affibody ZHER2) upon treatment with DARPins and mAbs as quantified by FRAP.** a) Dramatic loss of mobility upon treatment with bipDARPin 6L1G compared to untreated cells (Untr.). b) Concentration dependence of the mobility reduction as illustrated by the relative fluorescence recovery after 48 s. c) Monovalent, monoparatopic DARPins 6L46 and GL4G reduce HER2 mobility only slightly, while a mixture of both (GL4G + 6L46) fully immobilizes HER2 through crosslinking. d) Similarly, moderate mobility changes are observed upon treatment with therapeutic mAbs PZB and TZB but almost complete immobilization by their combination (PZB + TZB).
Immunization of HER2 correlates with oligomerization. These observations suggested that diffusion behavior directly reflects the differential oligomerization properties. To verify this correlation, we exploited simultaneous dual-color detection for analyzing dimerization and oligomerization of HER2 by DARPins and mAbs with sub-diffraction resolution. To this end, co-tracking of molecules localized in both spectral channels was performed to unambiguously identify HER2 complexes comprising two or more molecules. The fraction of such co-trajectories observed after treatment with different binders are compared in Fig. 6e. Only negligible dimerization levels (<0.05%) were detected in control experiments without any agent and in cells treated with the monovalent DARPin OL4G, thus confirming a limited intrinsic tendency of HER2 to homodimerize. By contrast, all other DARPins and mAbs significantly increased the fraction of co-locating HER2 molecules (Fig. 6e and Supplementary Table 2). Owing to the relatively low degree of labeling used in these experiments (<10%), however, these numbers cannot be directly converted into fractions of dimers and oligomers. Based on the relative numbers observed for different HER2 binders, however, their propensity to dimerize and oligomerize HER2 can be compared. Similar co-locomotion levels around 0.5% were observed for the homo-bivalent, monoparatopic DARPins GL4G and 6L46 as well as for TZB and PZB, all of which presumably dimerize HER2. A significantly higher value of 1.04% was observed for the bipDARPin 6L1G, indicating higher oligomerization levels, with an average of ~4 molecules/oligomer estimated from the ~2-fold increased dual-labeled fraction. Substantially stronger differences were observed for the immobile fraction. Here, the dual-labeled fractions for dimerizing agents were similar as for the mobile fractions, while it increased by a factor of ~6 for 6L1G (Fig. 6f and Supplementary Table 2). In line with this observation, we found a strong correlation of immobile and dual-labeled fraction size (Fig. 6g). Taken together, these results suggest that the bipDARPin 6L1G crosslinks HER2 into oligomers of different sizes (~4–10 copies), which is accompanied by a severe loss in mobility.

To confirm this hypothesis, we performed single-molecule localization, tracking, and co-tracking experiments with HER2 in the presence of the crosslinking mixture of 6L46 and GL4G. Under these conditions, largely identical spatiotemporal dynamics and diffusion of HER2 as compared to treatment with monoparatopic controls GL4G and 6L46 (Supplementary Fig. 1) and (likewise homo-bivalent, monoparatopic) antibodies TZB and PZB, all of which are expected to dimerize HER2; (iii) a markedly higher immobile fraction (>50%) was found for the biparatopic DARPin 6L1G, accompanied by a further decrease in the average diffusion constant of the mobile fraction.

HER2 immobilization occurs at a wide range of expression levels, as indicated by FRAP measurements on HER2-overexpressing BT-474 cells and non-induced HEK cells stably transfected with HaloTagged HER2. Note that the sensitivity of the FRAP method limits the cell lines that can be investigated to those with at least intermediate expression levels (Supplementary Table 1 and Supplementary Fig. 2). Data for untreated cells (Untr.) and 6L1G in c are identical to those shown in Fig. 4c.
6L1G were observed (Supplementary Figs. 6a, b and 7a). As expected, the crosslinking mixture of 6L46 and GL4G also induced a comparable increase of the immobile fraction, as well as a decrease of the average diffusion coefficient of the remaining mobile particles (Supplementary Figs. 6c, d and 7b, c). Likewise, very similar levels of dual-labeled mobile and immobile fractions were found in the presence of the 6L46–GL4G combination and 6L1G (Supplementary Figs. 6e, f and 7d, e), corroborating that similar levels of oligomerization are induced by these treatments.

HER2 lockdown is caused by increased trapping in membrane nanodomains. Given the high mobility of HER2 in the absence of DARPin, the dramatic effect of dimerization and oligomerization on the diffusion remains surprising. We therefore more carefully analyzed the diffusion properties of HER2 by unbiased SMT, which was applied to cells with very low cell surface expression levels to ensure high tracking fidelity. Confinement analysis revealed characteristic short-term arrests of mobile HER2, which was occasionally observed even for untreated cells or after incubation with the monovalent OL4G (Fig. 7a–c, Supplementary Movies 3 and 4, and Supplementary Table 2). Upon treatment with mAbs and homo-bivalent DARPins that presumably dimerize HER2, an enhanced occurrence of such transient arrest events was observed, while the dwell times of the arrested state remained rather constant (Fig. 7d–f, Supplementary Fig. 9a, b, and Supplementary Table 2). Treatment with the apoptosis-inducing bipDARPin 6L1G strongly further enhanced the occurrence of HER2 transient arrest events but only moderately increased the dwell time of the arrested state. Thus, mobility within the 32-s time frame of these experiments was still observed.

Fig. 6 Lockdown and oligomerization of HER2 diffusion as resolved by single-molecule localization microscopy. a Representative SMLM super-resolution images rendered from 500 consecutive frames (localization precision ~20 nm) with localization densities encoded as gray values. Scale bar: 10 µm. Magnified center regions are shown as insets (scale bar: 2 µm). b Particle densities in the DY-649 channel observed in different single-molecule experiments. c, d Comparison of the immobile fraction identified by spatiotemporal clustering analysis (c) and the diffusion coefficient within the mobile fraction (d) for HER2 for different treatments. e, f Colocalized HER2 molecules in the mobile (e) and the immobile fraction (f) quantified by dual-color co-tracking. g Correlation of the total fraction of co-localized molecules with the immobile fraction and linear regression (R² = 0.97, Pearson’s r = 0.99, p = 3.5 × 10⁻⁵). Different colors (cyan, control or monovalent; red, homo-bivalent; green, biparaitopic) in b–g indicate the three distinct groups of agents as referenced in the main text. The DARPin are shown schematically in Supplementary Fig. 1. Significance values for b–f are provided in Supplementary Fig. 4.
for most HER2 molecules even after treatment with 6L1G, with HER2 essentially "dropping" from one confinement zone into the next (Supplementary Movie 4 and Supplementary Fig. 9e). Single-molecule trajectories of transiently immobilized particles revealed residual, highly confined diffusion (Supplementary Movies 5–7 and Supplementary Fig. 9e), suggesting transient trapping in nanoscopic membrane domains with a diameter of ~35 nm. Upon treatment with 6L1G, characteristic successive transient immobilization of HER2 was observed (Fig. 7c, Supplementary Movie 4, and Supplementary Fig. 9e). Taken together, these results suggest that the efficient "lockdown" of HER2 by bip-DARPins does not involve irreversible immobilization but rather can be ascribed to a strongly increased propensity for being trapped within nanoscopic confinement zones.

Discussion

Blocking the activity of cell surface receptor tyrosine kinases by mAbs and other binders has emerged as a highly successful strategy in cancer therapy, and frequently these agents block ligand-binding-mediated activation. Molecular mechanisms effective for inhibiting constitutive activation, however, have largely remained unclear. We have previously introduced bip-DARPins that bind HER2 via two distinct epitopes, which yielded a surprisingly effective inhibition of downstream signaling pathways, leading to apoptosis in addicted cell lines and in vivo. To pinpoint the mechanism responsible for HER2 inhibition, we have investigated here the properties of the HER2–bipDARPin complexes in the plasma membrane. We were particularly intrigued by the possibility that bipDARPins may inhibit HER2 at the plasma membrane by the formation of daisy chains (cf. Fig. 1d). While chemical crosslinking confirmed that active bipDARPins crosslink HER2 into oligomers, the 3–4 copies of HER2 we found in these oligomers was surprisingly low. To resolve this conundrum, we focused on the spatiotemporal organization and dynamics of HER2 in the plasma membrane of live cells and its changes upon treatment with different binders. Remarkably, even in untreated cells, a substantial fraction of HER2 (40–60%) was immobile within the time and length scale of our FRAP experiments. Similar levels of immobile fractions have been previously observed in FRAP experiments with HER2, epidermal growth factor receptor (EGFR), and ErbB3 in a variety of cell lines, suggesting that this may be a fundamental feature of ErbB family receptors. In line with previous reports, depolymerizing the cortical actin cytoskeleton did not lead to any significant changes in mobility (Fig. 5c), suggesting that the HER2 immobilization is not caused by the membrane skeleton. SMT, however, revealed...
high local mobility of HER2 and residual diffusion even in the presence of bipDARPins. Rather, we identified transient immobilization even in the absence of any binder as a characteristic feature of HER2 diffusion in the plasma membrane. Such short-term immobilization has been related to receptor trapping in nanoscopic plasma membrane sub-compartments\textsuperscript{34–36}, which have been reported for several cell surface receptors including the ErbB family members EGFR\textsuperscript{32,37} and HER2\textsuperscript{32}. Our results suggest that such transient immobilization of HER2 may result into apparent local confinement when observed on the much larger length scale probed by FRAP.

Treatment with DARPin constructs for treatment were expressed and purified as previously described\textsuperscript{11,13}. PZB (Perjeta\textsuperscript{®}) was a kind gift from Professor Uwe Zangemeister-Wittke and TZB (Herceptin\textsuperscript{®}) was obtained from

Characteristics point to transient partitioning into membrane nanodomains rather than simply the oligomerization-induced decreased hopping probability across the picket fence of the cortical actin cytoskeleton (membrane skeleton (MSK)) as the fundamental basis of heterogeneous diffusion in the plasma membrane (Fig. 8a)\textsuperscript{38–40}. Nanodomains in the mammalian plasma membrane have been related to cooperative interactions of lipids and proteins\textsuperscript{36}. Thus, rather than partitioning, HER2 dimerization and oligomerization could increase the propensity of nanodomain formation. Our observations are in line with the concept of a hierarchical organization of the plasma membrane, with lipids and proteins transiently segregating into nanodomains within the framework of the MSK\textsuperscript{34,41}.

While the molecular basis responsible for HER2-specific nanodomain partitioning/formation remains speculative, the intrinsic propensity of HER2 for transient arrest in the absence of crosslinking agents is evident. The drastic increase in partitioning propensity with increasing oligomerization level is in line with the cooperativity required for nanodomain formation and partitioning\textsuperscript{41}. Based on MSD analysis, we estimate the size of the nanodomains to be below 50 nm, with residual mobility within the nanodomain (or even of the entire nanodomain). These zones do not seem to be related to endocytosis, as no increased endocytosis has been observed for bipDARPin-treated HER2\textsuperscript{10}. Furthermore, partitioning and transient arrest, which are unaffected by extensive treatment with potent inhibitors of the HER2 kinase, do not appear to be related to signaling, as previously described for other receptors\textsuperscript{37,42,43}.

On the contrary, our data implicate the possibility that transient arrest of HER2 obstructs downstream signaling, as we find a correlation between diffusional lockdown (i.e., uptake into nanodomains) and neutralization of signaling activity. However, mixtures of TZB and PZB (the approved combination for clinical therapy) or of homo-bivalent DARPin constructs\textsuperscript{6L46 and GL4G} yielded strong lockdown of HER2 diffusion to a similar level as bipDARPins. Yet, the clinical benchmark combination of TZB and PZB neither potently neutralizes downstream signaling nor causes apoptosis in the absence of cellular cytotoxicity, as has been shown previously\textsuperscript{10,44,45}. Therefore, lockdown is necessary but not sufficient for inducing apoptosis.

This highlights that the specific binding geometry of 6L1G is required for complete kinase inhibition\textsuperscript{10,46}, via formation of signaling-inactive oligomers (Fig. 8b), which also preclude interaction with EGFR and HER3\textsuperscript{10}. As we find that immobilization also occurs on cell lines resistant to HER2-targeted treatment\textsuperscript{25}, diffusion arrest is a feature of a novel mechanism of action, in which loss of mobility precedes—and is a prerequisite for—signaling inhibition\textsuperscript{10}. In therapeutic applications, it is attractive to increase plasma exposure either through conjugation to polymers to increase the molecular size above the renal filtration threshold or exploiting FcRn-mediated salvage pathways through albumin-binding domain or Fc region-containing molecules\textsuperscript{47}. Of note, the mechanism of action described here is compatible with both approaches: PEGylation slightly affects the in vitro potency but not the efficacy of bipDARPins\textsuperscript{10}, and the biparatopic targeting mechanism could be successfully transferred to IgG-based antibody constructs\textsuperscript{48}. Taken together, these insights suggest that deliberate daisy chaining of HER2 potently downregulates its activity by a combination of conformational and diffusional arrest and thus may represent a generic strategy to achieve pan-receptor inhibition.

Methods

Protein production and labeling. DARPin constructs for treatment were expressed and purified as previously described\textsuperscript{11,13}. PZB (Perjeta\textsuperscript{®}) was a kind gift from Professor Uwe Zangemeister-Wittke and TZB (Herceptin\textsuperscript{®}) was obtained from
Kantonsapotheke Zürich. Affibody Zis2 (ref. 22) containing a C-terminal cysteine was coupled to maleimide-containing dyes, either Oregon Green 488 (Thermo Fisher, cat. no. O6034) or DY-647P1 (Dyomics, cat. no. 647FP1-03) as previously described 39.

Generation of inducible HT-HER2 cell line. Stable cell lines for inducible expression of the HT-HER2 fusion were generated using the Flp-In TREx system (Thermo Fisher, cat. no. K650001). First, the human HER2 cDNA (Mammalian Gene Collection, Genbank accession number BC156753.1) without the signal peptide was amplified by PCR. The amplicons were subcloned into a derivative of the pcDNA5/FRT/TO-derived plasmid containing HaloTag-H2ER2 and the Flp recombinase vector pOe44 using TransIT-293x (Merck, cat. no. 213859), 0.02 mM leupeptin (Serva, cat. no. 1867), 0.01 mM pepstatin A (Serva, cat. no. 52682), and 0.02 mM marimastat (Calbiochem, cat. no. 52682) in complete medium was used to select for stably transfected clones. After isolation using cloning discs (Sigma-Aldrich, cat. no. Z374431), single clones were expanded and analyzed by flow cytometry for inducible expression of HaloTag-H2ER2 to obtain the HEK-TREx HT-HER2 cell line.

Crosslinking analysis. Semi-continuous BT474 cells were detached by Accutase treatment (Thermo Fisher, cat. no. AT104) for 5 min at 37 °C, washed with RPMI1604 medium containing 10% (v/v) fetal calf serum, and afterwards washed three times with 25 mM 4-(2-hydroxyethyl)-1-piperazinethanesulfonic acid (HEPES) at pH = 7.5 containing 130 mM NaCl. Cells were counted on a CASY cell counter (model “TT,” Omni Life Science), and 2 × 10^5 cells per sample were transferred to 200 μl HEPS buffer containing 10 nM of each DARPin or IgG. Cells were incubated for 1 h at 37 °C. Afterwards, cells were placed on ice, washed twice with 1 ml HEPS buffer and resuspended in 200 μl HEPES containing 2.5 mM bis[soxalocinimidyl]sulfate (BSI) (Pierce, cat. no. 21580). Crosslinking was performed for 1 h at room temperature; afterwards the reaction was quenched by washing with 3 × 50 mM tris(hydroxymethyl)aminomethane (Tris)–1% Triton X-100 and protease and phosphatase inhibitors at 4 °C. The following protease inhibitors were used at the indicated final concentrations: 1 mM phenylmethylsulfonyl fluoride (PMSF) and 100 μM leupeptin (Serva, cat. no. 52682), and 0.02 mM marimastat (Calbiochem, cat. no. 444289). The following phosphatase inhibitors were obtained from Sigma Aldrich and used at the indicated final concentrations: 1 mM sodium orthovanadate (cat. no. 220590), 1 mM sodium metavanadate (cat. no. 72060), 10 mM sodium fluoride (cat. no. 331058), 30 mM sodium phosphate (cat. no. G9422), and 50 mM sodium fluoride (cat. no. 71519). Lysates were kept on ice, cleared by centrifugation for 10 min at 20,000 × g at 4 °C, and supernatants were prepared for sodium dodecyl sulfate (SDS)–polyacrylamide gel electrophoresis (PAGE) in non-reducing SDS sample buffer (Invitrogen) and stored at −80 °C for 3 months. Samples were separated on gradient gels (Invitrogen) and afterwards blotted at 100 V for 70 min onto activated PVDF-FL membrane (Millipore, cat. no. IPFL00010) using NuPAGE transfer buffer (cat. no. NP00616, Invitrogen) containing 10% MeOH.

The following antibodies were used for western blot analysis: anti-ErbB2 from mouse (Santa Cruz, clone 485, cat. no. 790-2991), anti-mouse from goat conjugated to IRDye800 (Rockland, cat. no. 610-732-124), anti-rabbit from goat conjugated to Alexa Fluor 680 (Invitrogen, cat. no. A21076), anti-GAPDH from mouse (Santa Cruz, clone 6C3, cat. no. sc-32233), anti-ErbB2 from rabbit mAb (Abcam, cat. clone C-18, cat. no. sc-284), and anti-human IgG (heavy and light chain) from goat (Jackson Immunoresearch Laboratories). Rabbit polyclonal DARPin antibody (Sigma-Aldrich, cat. no. G9422), and 50 mM sodium fluoride (cat. no. 71519). Lysates were kept on ice, cleared by centrifugation for 10 min at 20,000 × g at 4 °C, and supernatants were prepared for sodium dodecyl sulfate (SDS)–polyacrylamide gel electrophoresis (PAGE) in non-reducing SDS sample buffer (Invitrogen) and stored at −80 °C for 3 months. Samples were separated on gradient gels (Invitrogen) and afterwards blotted at 100 V for 70 min onto activated PVDF-FL membrane (Millipore, cat. no. IPFL00010) using NuPAGE transfer buffer (cat. no. NP00616, Invitrogen) containing 10% MeOH.

Fluorescence recovery after photobleaching. Experiments were performed at room temperature, unless noted otherwise. We bleached a circular spot with 2 μm diameter, typically located in the equatorial plane 19. Acquisition and bleaching settings were optimized such that bleaching during acquisition was minimized (typically <5% of the initial fluorescence intensity was lost during the whole acquisition as determined by pre-bleach measurements) and to enable bleaching in a single step. If, for a single cell, >40% of initial fluorescence was bleached within one single step (e.g., due to low level of labeled receptor for this cell, or a bleach spot not precisely centered on the membrane, or drift), this cell was excluded from analysis. Because cell death can be accompanied by a dramatic loss of lateral HER2 mobility (Supplementary Fig. 10), we confirmed at the end of all experiments that cultures were essentially unaltered.

Image data including metadata were extracted from the proprietary Leica image file format (.lif) container, as put out by the instrument, with Fiji 51 using the Bioformats importer 23. We used a custom ImageJ script (available from the authors), which extracts the bleaching region of interest (ROI) coordinates from the metadata, then allows manual positioning of a control ROI at a distant spot on the same cell (to correct for acquisition photobleaching), and exports the mean intensity values of bleach and control ROI. Data were scaled on a per-cell basis according to \((P(t) − F(0)) / (\langle P(t) − F(0) \rangle)\), where \(P(t)\) is the mean fluorescence in the channel of interest after the bleach started at time \(t\), \(F\) is the signal immediately before the bleach started, \(\langle \cdot \rangle\) denotes the mean over all pixels of a ROI defined after the bleach, and \(\langle P(t) − F(0) \rangle\) is the (spatial and temporal) mean fluorescence before the bleach step. Assuming a constant offset, the low background fluorescence cancels out in the linear transformation applied for normalization 23. Afterwards, we corrected for the minor drift remaining after optimized settings by averaging single exponential fits to the mean fluorescence intensity of the control ROI 23. Unless noted otherwise, plotted lines indicate mean values of at least five cells per condition.

Single-molecule localization microscopy. For single-molecule experiments, an N-terminal fusion of the SNAP-tag to HER2 (SNAP-HER2), constructed analogously to the HT-HER2 fusion (see above), was used for transient expression in HeLa cells. To this end, HeLa cells were seeded in dishes (6 cm diameter). On the next day, 2.5 μg of plasmid in a volume of 2.5 μl was mixed with CaCl2, added to sterile water to yield a 1 μg μl−1 solution. The mixture was obtained at the total volume of 500 μl. The resulting solution was slowly added to 500 μl of 2× HBSS solution (280 mM NaCl, 50 mM HEPES, 1.5 mM Na2HPO4, pH = 6.95–7.05, sterile-filtered) with agitation and incubated for 10 min at room temperature. Afterwards, the suspension was evenly added to the cell dish, and the cells cultivated at 37 °C in 5% CO2, 5% atmosphere, then detached and plated overnight onto No. 1.5 coverslips that were pre-treated with a 1 mg ml−1 PLL–PEG–RGD solution, which was synthesized as described previously 24, to minimize non-specific binding of dye molecules to the coverslip surface 28. SNAP-HER2 was labeled by incubation with a mixture of 80 nm SNAP Surface 649 (DY-649P1, NEB) and 8 nm SNAP Surface 549 (DY-549P1, NEB) in 1 μl phenol red-free medium (DMEM) for 10 min at 37 °C, and the cells were afterwards washed twice with Dulbecco’s PBS. These concentrations of SNAP substrates ensured similar labeling efficiencies in both channels. All imaging experiments were carried out at room temperature (to minimize endocytosis of HER2). The sample was kept on ice during the imaging. Cells were scanned every 1 µs with an oxygen scavenger and a redox-active photoprotectant (0.5 mg/ml glucose oxidase (Sigma-Aldrich), 0.04 mg/ml catalase (Roche), 5% w/v glucose, 1 μl ascorbic acid, and...
1 μM methyl viologen) to reduce blinking and photobleaching of the fluorophores52. Treatments (DARPins and antibodies) were added to a final concentration of 100 nM. Dual-color single-molecule imaging was performed using TIFREM with an inverted microscope (Olympus IX71) equipped with a triple-line total internal reflection illumination condenser (Olympus) and a back-illuminated electronic multiplying charge-coupled device (EMCCD) camera (iXon DU897D, 512 × 512 pixels, Andor Technology) operated at −80 °C. A 150x magnification objective with a numerical aperture of 1.45 (UAPON 150/1.45 TIRFM, Olympus) was employed for sample illumination and fluorescence collection. For dual-color acquisition, DY-549P1 was excited by a 561 nm diode-pumped solid-state laser (CL-561-500 mW and DY-649P1 was excited by a 642 nm laser diode (LuxX 642-140, Omicron) at 0.65 mW (power output after passage of the objective). Fluorescence was detected using a spectral image splitter (DualView; Optical Insight) with a 640 DCXR dichroic beam splitter (Chroma Technology Corporation) in combination with a 585/40 band-pass filter (Semrock) for detection of DY-549P1 and a 690/70 band-pass filter (Chroma Technology Corp) for detection of DY-649P1, projecting the respective spectral channels on the upper and lower half of the EMCCD chip. Time series of 2000 images were recorded with a time resolution of 32 ms per frame.

**Single-molecule analyses.** Single-molecule localization was achieved using the multiple-target tracing algorithm (MTT)59. Briefly, individual signals were first identified with a fixed false positive rate of 10^-6 using pixel-wise hypothesis-testing against the local background noise (9 × 9 pixel evaluation box) and then localized with sub-pixel precision (typically 20 nm) approximating the microscope’s point-spread function with a two-dimensional Gaussian profile with predetermined fixed radius. Spatiotemporal super-resolution images were rendered by placing all localization within the first 500 frames on an upsampled pixel grid blurred with the average localization precision as described in detail previously37. Diffusion properties were determined only from the DY-649 channel because particle densities were more homogeneous. Prior to tracking, the data set was scanned for transient immobilization events (>540 ms or 20 frames) within a 120 nm search radius around each localization using an adapted version of the density-based spatial clustering of applications with noise principle59 as described earlier39. Individual positions of mobile particles were subsequently connected into trajectories (allowing for a three-frame-long observation gap) based on their locally characteristic displacement statistics calculated along each molecule trajectory96. MSDs of the tracked (>32 tracks or 10 frames) were fitted according to MSD = 4Dr − 4/3Dtr + 4εr, where r is the time lag (duration of displacement), Dr is the camera exposure time, and εr is the localization precision96, weighted by their expected inverse variances in order to obtain the instantaneous diffusion coefficient D (r > 160 ms or 5 frames). Crosslinked receptors were identified by dual-color single-molecule co-tracking as described in detail previously39. Briefly, spectral channels were first aligned spatially with sub-pixel accuracy based on a prior calibration measurement with multicolor fluorescent beads (TetraSpeck microspheres, 100 nm, Invitrogen). The fraction of mobile points was subsequently probed for co-localization (consecutive co-localization within 150 nm for at least 320 ms or 10 frames) (immobile as well as dual-labeled) as calculated for each frame respective to the observed number of total localizations and reported as their average over time. Alternation between mobile and immobile periods within individual trajectories was quantified by confinement analysis, as described previously90. Confinement was produced with a time resolution of 320 ms (10 frames) along individual trajectories (>3.2 s or 100 frames) against unrestricted motion of the HER2 monomer with an expected diffusion coefficient of 0.19 μm² s⁻¹ (OLAG). A controlled threshold to discriminate confined versus unbounded periods of diffusion was established based on simulation of free Brownian motion with a fixed false positive rate of 10^-6 using pixel-wise hypothesis-testing against the local background noise. Transition probabilities were calculated from the obtained state sequence (https://www.cs.ubc.ca/~murphyk/Software/HMM/hmm.html).

In all box plots showing data derived from single-molecule localization and tracking experiments, the center line indicates the median, the box limits indicate upper and lower quartiles, the whiskers show the 1.5x interquartile range.

**Reporting summary.** Further information on research design is available in the Nature Research Reporting Summary linked to this article.

**Data availability.** Source data underlying Figs. 2–7 are provided in Supplementary Data 1. The extensive raw data sets generated during the current study are available from the corresponding author upon reasonable request. All derived data and analyses supporting the findings of this study is included in this manuscript and its supplementary information files.

**Code availability.** Unless previously published and referenced in the text, all software is available from the authors upon reasonable request.

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Author contributions J.C.S., C.P.R., B.S., J.P., and A.P. designed research; J.C.S., C.P.R., J.S.B., M.S., and I.K. performed experiments; J.C.S., C.P.R., M.S., and I.K. analyzed the data; J.C.S., J.P., and A.P. wrote the manuscript with contributions from all authors.

Competing interests The authors I.C.S., M.S., and A.P. filed the following patents relating to the described bispecific antibodies: Bispecific HER2 Ligands for Cancer Therapy [US20191274181 A1; published]; Inventors: Andreas Plückthun, Florian Kast, Martin Schwil, Annemarie Honegger, Rastislav Tamaskovic, Christian Jost; Current Assignee: University of Zurich Her2-binding tetrameric polypeptides [EP3373374; published]; Inventors: Florian Kast, Martin Schwil, Annemarie Honegger, Jakob Stüber, Rastislav Tamaskovic, Andreas Plückthun; Current Assignee: University of Zurich. The University of Zurich has licensed out the invention to Innoven Biologics, Inc. The remaining authors declare no competing interests.

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Supplementary material

Apoptosis-inducing anti-HER2 agents operate through oligomerization-induced receptor immobilization

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### Supplementary figures

| Class                                      | Epitope(s)                        | Constructs | (abbreviation) |
|-------------------------------------------|-----------------------------------|------------|----------------|
| Single DARPin                             | HER2 ECD I                        | 9.26       | (EL1)          |
|                                            | HER2 ECD IV                       | 9.29       | (GL1G)         |
| Monoparatopic (bivalent binding)          | HER2 ECD I                        | 9.26-(Gly-Ser4)4-9.26 | (6L1G) |
|                                            | HER2 ECD IV                       | G3-(Gly-Ser4)4-G3       | (GL4G)         |
| Active biparatopic (bivalent binding, strong antitumor effect) | HER2 ECD I                        | 9.26-(Gly-Ser4)4-9.26 | (6L1G)         |
|                                            | HER2 ECD IV                       | 9.29-(Gly-Ser4)1-G3     | (9L1G)         |
| Control fusions                           | Non                              | off7-(Gly-Ser4)4-off7   | (OL4O)          |
|                                            | HER2 ECD IV (monovalent)          | off7-(Gly-Ser4)4/1-G3   | (OL4/1G)        |

Supplementary Figure 1: Overview of DARPin constructs used in this study. Abbreviation: ECD, extracellular subdomain.
Supplementary Figure 2: (a) Histograms of the flow cytometry data used in generation of Supplementary Table 1. Total HER2 expression (endogenous native HER2 and HaloTag-HER2 fusion, where present) was detected using the affibody ZHER2 conjugated...
to DY-647P1. For the HEK293_Halo-HER2 cell line, expression was induced by doxycycline addition where indicated. (b) Gating strategy applied in flow cytometry experiments shown in (a) and Supplementary Table 1. Intact cells were first identified as particles in the appropriate size range, which are inaccessible to a permeability marker (DAPI). A plot of the forward scatter peak area versus the peak height was then used to gate for single cells.
Supplementary Figure 3: Single-molecule confocal spectroscopy in a HEK293_Halo-HER2 cell line suggests that bipDARPins induce HER2 immobilization. (a) HT-HER2 was labeled with a mixture of HaloTag ligands carrying Alexa Fluor 647 (HTL-AF647) or Cy3 (HTL-Cy3). At 5 µW laser power, a pronounced initial bleaching phase was followed by a steady state with distinct single-molecule bursts (binning: 100 ms). (b)
After addition of biologically active, apoptosis-inducing DARPin 9L1G, the burst count in the steady state is significantly reduced compared to non-treated cells or cells treated with monovalent control-DARPin OL1G (see Supplementary Figure 1), which can only bind a single HER2 molecule. The first 100 s, where the background signal is not constant due to bleaching, were excluded from burst detection (binning: 100 ms for all samples). (c) Results of automated burst counting after treatments as in (b). Error bars represent the SEM.
Supplementary Figure 4: Statistical significance for data shown in Fig. 6b-f. Pair-wise significance testing (Welch one-sided t-test with unequal sample size and variance) of localization density of HER2-fSNAP labelled with DY-649 (a), immobile fraction (b), diffusion coefficient of the mobile phase estimated by mean squared displacement (MSD) analysis (c), fraction of points being both mobile and co-localized in both spectral channels (d) and fraction of points being both immobile and co-localized in both spectral channels (e). The significance matrix is read left-to-right color-coded by the reduction risk (RR = –lg(p-Value): ns, RR < 1.3; *, RR ≥ 1.3; **, RR ≥ 2 and ***, RR > 3).
Supplementary Figure 5. Diffusion properties of HER2 evaluated by mean squared displacement (MSD) analysis. Pooled MSDs of the mobile periods (and the immobile periods in the inset) from single-molecule trajectories of HER2 treated with different agents. Different colors indicate the three distinct groups of agents (cf. Figure 6) as referenced in the main text.
Supplementary Figure 6. Diffusional arrest by 6L1G is equivalent to crosslinking by the combination of monovalent but biparatopic 6L46 and GL4G. (a) Representative SMLM super-resolution images rendered from 150 consecutive frames with localization densities encoded as grey values. Scale bar: 5 µm. (b) Particle densities in the DY-649 channel observed in different single-molecule experiments. (c, d) Comparison of the immobile fraction identified by spatiotemporal clustering analysis (c) and the diffusion coefficient within the mobile fraction (d) for HER2 for different treatments. (e, f) Colocalized HER2 molecules in the mobile (e) and the immobile fraction (f). The DARPins are shown schematically in Supplementary Figure 1. Significance values for panels b-g are provided in Supplementary Figure 7.
Supplementary Figure 7. Statistical significance for data shown in Supplementary Figure 6b-f. Pair-wise significance testing (Welch one-sided t-test with unequal sample size and variance) of localization density of HER2-fSNAP labelled with DY-649 (a), immobile fraction (b), diffusion coefficient of the mobile phase estimated by mean squared displacement (MSD) analysis (c), fraction of points being both mobile and co-localized in both spectral channels (d) and fraction of points being both immobile and co-localized in both spectral channels (e). The significance matrix is read left-to-right color-coded by the reduction risk (RR = –\text{lg}(p\text{-Value}): \text{ns}, \text{RR} < 1.3; *, \text{RR} \geq 1.3; **, \text{RR} \geq 2 \text{ and } ***, \text{RR} > 3).
Supplementary Figure 8: Statistical significance for data shown in Fig. 7d-f. Pair-wise significance testing (Welch one-sided t-test with unequal sample size and variance) for the (a) identified confined fraction (b) the transitioning probability into confined motion and (c) the reverse transition probability out of confined into free motion. The significance matrix is read left-to-right color-coded by the reduction risk (RR = −lg(p-Value): ns, RR < 1.3; *, RR ≥ 1.3; **, RR ≥ 2 and ***, RR > 3).
Supplementary Figure 9. Spatiotemporal dynamics of transient arrest of HER2 diffusion. (a, b) Half-lives calculated from the estimated probabilities for transitioning from free to arrested diffusion (a) and, vice versa, transitioning from confined to free motion (b), as obtained from confinement analysis. (c, d) Respective pair-wise significance testing (Welch one-sided t-test with unequal sample size and variance) for the mobile half-life (c) and the arrested half-life (d). The significance matrix is read left-to-right color-coded by the reduction risk (RR = −lg(p-Value)): ns, RR < 1.3; *, RR ≥ 1.3; **, RR ≥ 2 and ***, RR > 3). (e) Detected immobilization events color-coded by time for
monovalent OL4G and biparatopic 6L1G. The white arrow points at immobilization events scattered across space over time as detected by spatiotemporal clustering. Temporal color-coding of the trajectories according to the legend in the bottom (in milliseconds). Scale bar: 1 µm. The overall particle density was very similar in both experiments.
Supplementary Figure 10: Metabolic inhibition (ATP depletion) is unsuitable for single-molecule tracking experiments because it leads to rapid loss of receptor mobility if combined with photoprotection cocktails. Cells were exposed to a photoprotection cocktail combined with metabolic inhibitors NaN$_3$ and 2-deoxy-D-glucose in PBS to achieve ATP depletion. The time course of immobilization, measuring 30 frames of 0.020 s every 30 s, was quantified using the DBSCAN algorithm.
## Supplementary tables

Supplementary Table 1: Estimation of total HER2 expression (HaloTag fusion and wt HER2) on various cell lines.

| Cell line                  | Rel. mean surface expression | Estimated\(^2\) mean number of receptors per cell | Estimated\(^3\) mean cell surface density [µm\(^{-2}\)] |
|----------------------------|-------------------------------|--------------------------------------------------|--------------------------------------------------|
| **Detection with anti-HER2 affibody\(^4\)**                     |                               |                                                  |                                                  |
| BT-474                    | 100.0%                        | 870,000                                          | 693                                              |
| HEK293_Halo-HER2, induced  | 100.8%                        | 877,297                                          | 698                                              |
| HEK293_Halo-HER2, non-induced | 8.1% ± 0.9%                 | 70,323 ± 8,204                                   | 56 ± 7                                           |
| HeLa                      | 1.1% ± 0.3%                   | 9,726 ± 2,962                                    | 8 ± 2                                            |
| **Detection with anti-HER2 DARPin\(^5\)**                       |                               |                                                  |                                                  |
| BT-474                    | 100.0%                        | 870000                                           | 693                                              |
| HEK293_Halo-HER2, induced  | 108.3%                        | 942416                                           | 750                                              |
| HEK293_Halo-HER2, non-induced | 6.9%                        | 60237                                            | 48                                               |
| HeLa                      | 1.8%                          | 15445                                            | 12                                               |

\(^{1}\) Histograms of the flow cytometry data for the HER2 affibody are shown in Supplementary Figure 2; the histograms for BT474 and HeLa cells measured using the anti-HER2 DARPin-GFP fusion have been previously published\(^4\).

\(^{2}\) Calculated from the known receptor number for BT-474 cells\(^5\).

\(^{3}\) Estimating a cell as a sphere with a radius of 10 µm.

\(^{4}\) Measured by using the anti-HER2 affibody ZHER2 (ref. 6).

\(^{5}\) Measured by using an anti-HER2 DARPin-GFP fusion.
Supplementary Table 2: Diffusion and interaction of HER2 as quantified by SMLM.

|                     | un- | OL4G | GL4G | 6L46 | PZB | TZB | 6L1G |
|---------------------|-----|------|------|------|-----|-----|------|
|                     | (n=3) | (n=7) | (n=6) | (n=8) | (n=5) | (n=3) | (n=5) |
| Loc. Density        | Median | 0.55 | 0.35 | 0.63 | 0.44 | 0.52 | 0.56 | 0.40 |
|                     | IQR  | 0.02 | 0.19 | 0.14 | 0.28 | 0.48 | 0.17 | 0.15 |
|                     | Mean | 0.55 | 0.43 | 0.59 | 0.46 | 0.58 | 0.60 | 0.45 |
|                     | SEM  | 0.01 | 0.08 | 0.04 | 0.06 | 0.13 | 0.07 | 0.07 |
| Imm. Fraction       | Median | 5.76 | 4.16 | 18.47 | 15.82 | 11.87 | 15.69 | 60.13 |
|                     | IQR  | 2.75 | 4.21 | 15.53 | 6.47 | 10.58 | 5.06 | 15.50 |
|                     | Mean | 5.64 | 6.03 | 18.84 | 15.01 | 15.19 | 15.10 | 56.92 |
|                     | SEM  | 1.06 | 1.92 | 3.14 | 1.75 | 4.30 | 1.88 | 5.10 |
| Diff. Coeff         | Median | 0.22 | 0.19 | 0.14 | 0.12 | 0.12 | 0.14 | 0.06 |
|                     | IQR  | 0.02 | 0.03 | 0.02 | 0.02 | 0.02 | 0.03 | 0.01 |
|                     | Mean | 0.21 | 0.20 | 0.14 | 0.12 | 0.13 | 0.14 | 0.06 |
|                     | SEM  | 0.01 | 0.01 | 0.01 | 0.00 | 0.01 | 0.01 | 0.00 |
| Mobile Dual-lab.    | Median | 0.02 | 0.00 | 0.34 | 0.69 | 0.31 | 0.25 | 1.04 |
| Fraction [%]        | IQR  | 0.07 | 0.03 | 0.45 | 1.01 | 0.79 | 0.27 | 1.04 |
|                     | Mean | 0.04 | 0.01 | 0.37 | 0.91 | 0.47 | 0.25 | 1.23 |
|                     | SEM  | 0.03 | 0.01 | 0.11 | 0.26 | 0.23 | 0.08 | 0.32 |
| Immobile Dual-lab.  | Median | 0.00 | 0.00 | 0.28 | 0.04 | 0.07 | 0.37 | 5.90 |
| Fraction [%]        | IQR  | 0.02 | 0.00 | 0.56 | 0.20 | 0.55 | 0.86 | 2.99 |
|                     | Mean | 0.01 | 0.06 | 0.38 | 0.12 | 0.43 | 0.43 | 5.44 |
|                     | SEM  | 0.01 | 0.06 | 0.13 | 0.06 | 0.38 | 0.38 | 1.15 |
| Confined Frac-      | Median | 4.35 | 3.13 | 17.34 | 11.79 | 14.67 | 7.75 | 57.35 |
| tion [%]            | IQR  | 1.11 | 2.65 | 7.02 | 6.54 | 4.47 | 3.33 | 5.62 |
|                     | Mean | 3.95 | 3.59 | 18.41 | 13.33 | 14.71 | 6.58 | 60.43 |
|                     | SEM  | 0.47 | 1.03 | 2.49 | 1.79 | 1.36 | 1.41 | 2.78 |
| Conf. Trans.        | Median | 0.75 | 0.82 | 2.27 | 2.85 | 2.69 | 1.66 | 10.75 |
| Prob. [%]           | IQR  | 1.13 | 0.44 | 0.88 | 0.88 | 0.72 | 0.57 | 3.92 |
|                     | Mean | 1.01 | 0.94 | 2.25 | 2.88 | 2.65 | 1.66 | 11.08 |
|                     | SEM  | 0.45 | 0.09 | 0.32 | 0.24 | 0.28 | 0.22 | 0.92 |
| Mob. Trans.         | Median | 5.96 | 10.09 | 3.92 | 8.55 | 7.29 | 6.71 | 3.56 |
| Prob. [%]           | IQR  | 1.58 | 6.68 | 1.97 | 3.81 | 2.67 | 1.03 | 1.84 |
|                     | Mean | 6.38 | 11.29 | 4.72 | 9.82 | 8.30 | 6.65 | 3.57 |
|                     | SEM  | 0.64 | 1.66 | 0.83 | 1.22 | 1.47 | 0.40 | 0.51 |
| Mobility Half-Time  | Median | 2.93 | 2.70 | 0.99 | 0.77 | 0.81 | 1.33 | 0.20 |
| [s]                 | IQR  | 3.40 | 1.05 | 0.40 | 0.27 | 0.26 | 0.48 | 0.07 |
|                     | Mean | 3.26 | 2.48 | 1.10 | 0.80 | 0.88 | 1.37 | 0.19 |
|                     | SEM  | 1.32 | 0.22 | 0.19 | 0.07 | 0.12 | 0.18 | 0.02 |
| Confinement         | Median | 0.36 | 0.21 | 0.56 | 0.25 | 0.29 | 0.32 | 0.61 |
| Half-Time [s]       | IQR  | 0.08 | 0.13 | 0.23 | 0.09 | 0.08 | 0.05 | 0.35 |
|                     | Mean | 0.34 | 0.21 | 0.52 | 0.23 | 0.28 | 0.32 | 0.68 |
|                     | SEM  | 0.03 | 0.07 | 0.07 | 0.02 | 0.04 | 0.02 | 0.12 |

IQR, interquartile range; SEM, standard error of the mean. n denotes the number of cells analyzed per condition.
Supplementary methods

**Confocal single-molecule fluorescence spectroscopy**

Confocal single-molecule fluorescence measurements were conducted on a customized MicroTime 200 (PicoQuant) based using an Olympus IX71 microscope body as previously described\(^7\). In brief, fluorescent dyes were excited alternatingly\(^6\) by a 20-MHz supercontinuum laser (SC-450-4, Fianium, with a 520/15 band-pass filter, Chroma Technology) and a 635-nm pulsed laser (PicoQuant) at 5 µW of excitation power each, as measured at the back aperture of the objective. The light was focused into the sample by an UplanApo 60×/1.20-W objective (Olympus). SPCM-AQR-15 single-photon avalanche diodes (PerkinElmer) were used to detect photons from each channel, and their arrival times were recorded in four channels of a HydraHarp 400 counting module (PicoQuant) with a resolution of 16 ps. A piezo stage combination (P-733.2 and PIFOC, Physik Instrumente GmbH), to which the objective was mounted, enabled 3D scans. After performing an initial x-y scan to obtain cell outlines, time traces were recorded at a fixed point in the apical (upper) cell membrane for 180 s. Data processing and analysis (binning: 100 ms for all samples) were done using Fretica, a Wolfram Symbolic Transfer Protocol add-on for Mathematica (Wolfram Research) (https://schuler.bioc.uzh.ch/programs/). The first 100 s, where the background signal is not constant due to bleaching, were excluded from burst detection. A binning time of 1 ms, a burst threshold of 10 photons per bin, and 20 and 10\(^6\) photons as lower and higher bounds for the total number of photons in a burst, respectively, were used for burst identification.

**Flow cytometry**

Subconfluent cells were harvested using trypsin. \(1\times10^6\) cells (as determined using a CASY TT cell counter, OLS OMNI Life Science) were washed once by resuspension in 1 ml Dulbecco’s phosphate buffered saline (DPBS) and subsequent centrifugation at 800 × g for 1 min. Cells were then resuspended in 1 ml DPBS supplemented with 1% bovine serum albumin and 50 mM sodium azide and (PBSBA), which furthermore contained 500 ng ml\(^{-1}\) of 4’,6-diamidino-2-phenylindole (DAPI). All samples were then split up into two aliquots of 500 µl each. To one of the aliquots, 6 µM of non-labeled affibody ZHER2 was added, and the samples incubated for 20 min at room temperature. Subsequently, ZHER2 conjugated to DY-647-P1 was added to a concentration of 40 nM, and the samples incubated for further 20 min. Finally, the samples were washed three times in PBSBA as described above and transferred to FACS tubes (BD).
prior to measurement on a LSR Fortessa II (BD) flow cytometer. Data were analyzed in FlowJo 10.7.1 (FlowJo, LLC).
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