Biased Agonism as a Mechanism for Differential Signaling by Chemokine Receptors

Sudarshan Rajagopalan, Daniel L. Bassoni, James J. Campbell, Norma P. Gerard, Craig Gerard, and Tom S. Wehrman

From the Departments of Medicine, Duke University Medical Center, Durham, North Carolina 27710, DiscoveRx Corporation, Fremont, California 94538, Department of Dermatology, Brigham and Women’s Hospital, Boston, Massachusetts 02115, and Division of Respiratory Diseases, Department of Pediatrics, Children’s Hospital, Boston, Massachusetts 02115

Chemokines display considerable promiscuity with multiple ligands and receptors shared in common, a phenomenon that is thought to underlie their biochemical “redundancy.” Their receptors are part of a larger seven-transmembrane superfamily, commonly referred to as G protein-coupled receptors, which have been demonstrated to be able to signal with different efficacies to their multiple downstream signaling pathways, a phenomenon referred to as biased agonism. Biased agonism has been primarily reported as a phenomenon of synthetic ligands, and the biologic prevalence and importance of such signaling are unclear. Here, to assess the presence of biased agonism that may underlie differential signaling by chemokines targeting the same receptor, we performed a detailed pharmacologic analysis of a set of chemokine receptors with multiple endogenous ligands using assays for G protein signaling, β-arrestin recruitment, and receptor internalization. We found that chemokines targeting the same receptor can display marked differences in their efficacies for G protein- or β-arrestin-mediated signaling or receptor internalization. This ligand bias correlates with changes in leukocyte migration, consistent with different mechanisms underlying the signaling downstream of these receptors induced by their ligands. These findings demonstrate that biased agonism is a common and likely evolutionarily conserved biological mechanism for generating qualitatively distinct patterns of signaling via the same receptor in response to different endogenous ligands.

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‡ †1 ‡ § †2

The abbreviations used are: 7TMR, seven-transmembrane receptor; **Barr**, β-arrestin; GPCR, G protein-coupled receptor; EA, enzyme acceptor; ED, enzyme donor; PK, ProLink; CCL, CC ligand; CCR, CC receptor; CXCL, CXC ligand; CXCR, CXC receptor.
**Experimental Procedures**

**Pharmacologic Assay Systems—PathHunter® GPCR Arrestin,** PathHunter GPCR Active Internalization, and cAMP Hunter™ cell lines (DiscoveRx Corp., Fremont, CA) were used to test the function of the chemokine GPCR targets.

All of the assays used in this study are based on enzyme fragment complementation technology (8). The basis of enzyme fragment complementation technology centers on two fragments of β-galactosidase (β-gal) enzyme: the enzyme acceptor (EA), which lacks residues 11–41, and the enzyme donor (ED), whose structure includes those missing residues. When EA and ED are mixed, they bind to form an active β-gal enzyme that hydrolyzes substrate to yield a detectable luminescent signal (9). This technology has been described in detail previously (8–10).

To link β-gal activity to cellular cAMP levels, three reagents are used in the HitHunter™ assay (11, 12): an ED conjugated with cAMP (ED-cAMP), a separate antibody that binds to cAMP (ED), and the EA fragment. In the absence of cAMP, the cAMP antibody is bound to ED-cAMP, preventing formation of an active enzyme complex. Free cAMP generated upon adenylyl cyclase activation by Gs results in competitive displacement of ED-cAMP from the cAMP antibody; this ED-cAMP then complements with EA to form an active enzyme. The greater the concentration of free cAMP, the more antibody that is bound to it, thus leaving ED-cAMP to form an active β-gal enzyme complex and hydrolyze the substrate to produce a luminescent signal. Detailed protocols for this assay result in highly reproducible and quantitative data as described previously (12).

For βarr recruitment, the PathHunter β-gal complementation system from DiscoveRx Corp. was used (13). In this assay, the 7TMR of interest is fused to the ED peptide, termed ProLink (PK), which weakly binds a complementary fragment, EA, which is fused to the C terminus of the arrestin protein. Activation of the PK-tagged 7TMR results in recruitment of the βarr-EA fusion and results in complementation of the enzyme. Again, the enzymatic activity is detected with luminescence. The βarr-EA fusion in PathHunter parental cells is designed to be in excess such that the PK-fused 7TMRs are the limiting factor; this results in an assay with no receptor reserve and where potency will closely track with ligand binding. A detailed protocol for performing this experiment has been published previously (13).

Receptor internalization was quantified using the GPCR Active Internalization assay from DiscoveRx Corp. that is based on previously described technology that monitors total receptor internalization (14). In this assay, a cell line is engineered to coexpress an untagged GPCR, an EA-tagged βarr, and a PK tag linked to endofin, which is localized to endosomes. Activation of the untagged GPCR induces βarr recruitment to and internalization of the receptor-βarr-EA complex in PK-tagged endosomes. Therefore, this assay measures recruitment of receptor-βarr complexes to endosomes.

Assay conditions were executed as follows. 20 μl of working cell suspension (5000 cells/well for internalization and arrestin; 10,000 cells/well for cAMP measurements) were added to white, tissue culture treated 384-well assay plates and incubated overnight at 37 °C in 5% CO₂. Ligands were reconstituted in PBS + 0.1% BSA prior to use. Chemokine ligands were used from DiscoveRx Corp. or R&D Systems (Minneapolis, MN). Agonists were serially diluted 1:3 in Opti-MEM + 1% FBS to generate 11-point titrations of ligand (plus vehicle-only (−) control). PathHunter Arrestin cell lines were exposed to compound for 1.5 h at 37 °C in 5% CO₂; PathHunter Active Internalization cell lines were exposed to compound for 3 h at 37 °C in 5% CO₂. PathHunter chemiluminescence detection reagents (DiscoveRx Corp., catalog number 93-0001) were added to cells according to the manufacturer’s instructions. For cAMP assays, on the day of the assay, cell plate media was removed from the plate and replaced with 15 μl of an antibody solution consisting of 2 parts Hanks’ balanced salt solution, 10 μg HEPES, and 1 part cAMP X+ antibody reagent. Agonists were serially diluted 1:3 in Hanks’ balanced salt solution, 10 μg HEPES, and forskolin to generate 11-point titrations of ligand (plus vehicle-only (−) control) with a constant concentration of forskolin. Cells were exposed to compound for 0.5 h at 37 °C in 5% CO₂. HitHunter cAMP X+ detection reagents were added to according to the manufacturer’s instructions (DiscoveRx Corp., catalog number 90-0075). The signal for all assays was detected using an EnVision 2100 multilabel reader (PerkinElmer Life Sciences) in luminescence mode. Raw relative luminescence units were plotted using GraphPad Prism (sigmoidal dose response, variable slope) software (San Diego, CA).

**Bias Analysis—** Equimolar plots (“bias plots”) and bias factors were calculated as described previously (15). The theoretical rationale for the equation used to calculate ligand bias has been described previously (15). The equation used to calculate bias factor is derived from the method of Furchgott (16) for comparison of equiactive signaling and is related to intrinsic relative activities proposed by Ehert and co-workers (17) (see supplemental materials from Rajagopal et al. (15) for a detailed discussion). Briefly, we first account for different levels of amplification inherent to each pathway downstream of the stimulus of AR* by including an amplification factor $a_{path}$, resulting in $s = a_{path} \times [AR^*]$. The fraction of receptors in a signaling-competent conformation is given by Equation 1.

$$\frac{[AR^*]}{[R_0]} = \frac{K_a[A]}{K_D + (1 + K^*)[A]}$$  

(Eq. 1)

When the responses of the two different pathways, e.g. G protein and βarr, are equal, the underlying stimuli are equal.

$$a_1 \times \frac{K_a[A_1]}{K_D + (1 + K^*)[A_1]} = a_2 \times \frac{K^*[A_2]}{K_D + (1 + K^*_2)[A_2]}$$  

(Eq. 2)

After some straightforward manipulations, 1/[A₂] can be expressed in terms of 1/[A₁] as follows.

$$\frac{1}{[A_2]} = \frac{a_2}{a_1} K^*_2 \times \frac{1}{[A_1]} + \frac{1}{K_D} \left( \frac{a_2}{a_1} K^*_2 (1 + K^*_1) - (1 + K^*_2) \right)$$  

(Eq. 3)

In this linear relationship, $m = (a_2/a_1) \times (K^*_2/K^*_1)$ is the slope of the line. If we then compare the slopes from one ligand ($m_{lig}$) with the reference balanced agonist ($m_{ref}$), the amplification terms cancel each other out, and we obtain Equations 4 and 5.
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where $\beta$ is the bias factor and estimates the molecular efficacy of pathway 1 versus pathway 2 on a logarithmic scale; e.g., a bias factor of 1 for $G_i/\beta$ Garr means that the ligand is 10 times better at generating the active receptor conformation for $G_i$ signaling compared with $\beta$arr signaling (compared with the reference balanced agonist).

If we derive this same relationship in terms of a pharmacologic fit with the Hill coefficient set to 1 with no basal activity, we obtain a similar expression after some straightforward manipulations as shown below.

$$R_1 = \frac{E_{\text{max},1}[A]}{[A] + EC_{50,1}} = \frac{E_{\text{max},1}}{1 + \frac{EC_{50,1}}{[A]}} = \frac{E_{\text{max},2}}{1 + \frac{EC_{50,2}}{[A]}} = R_2$$ (Eq. 6)

$$E_{\text{max},1} \left(1 + \frac{EC_{50,2}}{[A_2]} \right) = E_{\text{max},2} \left(1 + \frac{EC_{50,1}}{[A_1]} \right)$$ (Eq. 7)

$$\frac{1}{[A_1]} = \frac{1}{E_{\text{max},2}EC_{50,1}} \left(1 + \frac{EC_{50,2}}{[A_2]} \right) - \frac{E_{\text{max},2}}{E_{\text{max},1}}$$ (Eq. 8)

$$\frac{1}{[A_1]} = \frac{(E_{\text{max},1}EC_{50,2})}{EC_{50,1}E_{\text{max},2}} \left(1 + \frac{E_{\text{max},1} - E_{\text{max},2}}{E_{\text{max},1}EC_{50,1}} \right)$$

$$= \frac{RA_{12}}{1} + \frac{(E_{\text{max},1} - E_{\text{max},2})}{E_{\text{max},1}EC_{50,1}}$$ (Eq. 9)

where $RA_{12}$ is the intrinsic relative activity proposed by Ehlerdt (18). Therefore, if the data can be fit well by a logistic expression with a Hill coefficient equal to 1, then the intrinsic relative activity is equal to the ratio of equilibrium constants for receptor activation. Then we obtain Equation 10.

$$\beta = \log \left( \frac{RA_{12,\text{lig}}}{RA_{12,\text{ref}}} \right) = \log \left( \frac{E_{\text{max},1}EC_{50,2}}{EC_{50,1}E_{\text{max},2}}_{\text{lig}} \times \frac{E_{\text{max},2}EC_{50,1}}{EC_{50,2}E_{\text{max},1}}_{\text{ref}} \right)$$ (Eq. 10)

This form is equivalent to one that directly compares the effective signaling, or relative efficacy, of a ligand through each pathway relative to a reference balanced agonist. To better account for experimental errors, significance for bias was assessed not on whether the bias factor was statistically different from zero but rather on whether it was statistically different from the “balanced” agonist that had been chosen arbitrarily.

Ligand bias is not an absolute term but rather a relative term in comparing signaling under different conditions or between different ligands. As most receptors have a single endogenous agonist, that ligand can be chosen as the reference balanced agonist. However, in the setting of multiple endogenous agonists, the choice of the reference balanced agonist is necessarily arbitrary. Calculations and assessment of significance were performed in GraphPad Prism 5.1.

Cell Transfection—HEK293 cells were maintained in Eagle’s minimum essential medium supplemented with 10% FBS and 1% penicillin/streptomycin. For DNA transfection, FuGENE transfection reagent (Roche Applied Science) was added at a ratio of 5 µl to 1 µg of plasmid DNA to a solution of plasmid DNA in 500 µl of serum-free Eagle’s minimum essential medium. This transfection mixture was incubated for 30–45 min prior to addition to HEK293 cells of 50% confluence.

Confocal Microscopy—For confocal microscopy, HEK293 cells were split into 35-mm glass bottom dishes (MatTek, Ashland, MA) and transfected with constructs encoding 1 µg of receptor and 0.2 µg of $\beta$arr2-GFP. Forty-eight hours after transfection, cells were starved for at least 5 h in serum-free medium prior to treatment with ligand. For live cell imaging, cells were maintained at 37 °C with a heating plate while confocal microscopy was performed. Chemokines were added to a final concentration of 100 nM, and images were taken from 0 to 30 min. For immunostaining, cells were treated for 30 min with 100 nM ligand followed by aspiration of serum and fixation with 4% formaldehyde for 30 min. Samples were then washed three times with PBS followed by permeabilization and blocking with 0.1% Triton X-100 in 2% BSA in PBS for 1 h. Cells were then incubated overnight with primary antibody targeting the receptor at 1:200 dilution in 2% BSA in PBS. Samples were then washed three times with BSA/PBS prior to 1-h incubation with secondary antibody at 1:500 dilution and with Hoechst 33258 to stain DNA. Samples were then washed with BSA/PBS and visualized.

Ex Vivo Chemotaxis Assay for Human Peripheral Blood Lymphocytes—Blood was drawn into EDTA tubes, and RBCs were sedimented with dextran T-500. RBC-depleted suspension was overlaid on Ficoll-Paque Plus (GE Healthcare) and spun at 400 × g for 45 min at room temperature. Interface cells (peripheral blood mononuclear cells) were recovered and washed in a large volume of migration medium (RPMI 1640 medium (Invitrogen) with 10% bovine calf serum (Invitrogen) pre-equilibrated overnight in a 37 °C incubator with 5% CO2). Peripheral blood mononuclear cells were resuspended in migration medium and incubated in a T-175 flask for 45 min at 37 °C in 5% CO2. Nonadherent cells (primarily lymphocytes) were recovered for the chemotaxis assay.

Corning Costar 24-well Transwell plates were used for ex vivo migration. Chemokine dilutions in 600 µl of migration medium were added to the inner 8 wells of each 24-well plate. Recombinant human CCR10 ligands CCL27 and CCL28 were obtained from R&D Systems, and recombinant human CXCR4 ligands CXCL10, CXCL11, and CXCL11 were obtained from Peprotech, Inc. (Rocky Hill, NJ). Transwell inserts containing tissue culture treated polycarbonate membranes with 5-µm pore size were gently placed on top of the chemokine dilution, being careful not to trap bubbles. A lymphocyte suspension (100 µl containing $5 \times 10^5$ lymphocytes) was added to each Transwell rapidly after the Transwell was placed in the well. Four wells were set up for each concentration of each chemokine. Plates were incubated at 37 °C in 5% CO2 for 90 min, and then Transwells were removed and discarded.
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**TABLE 1**

| Chemokine receptor-ligand combinations tested in this study |
|------------------------------------------------------------|
| CCR1, CCL3, CCL4, CCL5, CCL8, CCL14, CCL15, CCL23, CCL28, CXCL1, CXCL2, CXCL3, CXCL4, CXCL5, CXCL6, CXCL9, CXCL10, CXCL11 |

Polystyrene microspheres (15 μm; Polysciences, Warrington, PA) were added to each well (5 × 10^4/well) for subsequent quantitation of migrated cells, and the contents of each well were transferred to centrifuge tubes for flow cytometry staining. Cells migrated to CCR10 ligands were stained using a three-color protocol consisting of anti-human CD4, CD34, and CD27. Cells migrated to CXCR3 ligands were stained using a four-color protocol consisting of anti-human CD3, CD8, CD27, and CD45RA. Flow cytometry data were collected using a FACSCanto flow cytometer (BD Biosciences) and analyzed using FlowJo 8.6.6 (Tree Star, Inc., Stanford, CA). The percentage of migration of each cell type was determined by calculating the ratio of beads to each cell type in the starting population versus the same cell type in the migrated population.

**RESULTS**

**Signaling by Chemokine Receptors**—To test the hypothesis that different ligands for the same chemokine receptor were acting as biased ligands, we obtained detailed concentration-response data on G protein signaling, βar signaling, and receptor internalization for a set of representative chemokine receptors. There are two major classes of chemokine receptors based on their chemokine ligands. CC ligands (CCL), which bind to CC receptors (CCR), have a motif with two adjacent cysteines, whereas CXC ligands (CXCL), which bind to CXC receptors (CXCR), have a single amino acid in between the two conserved cysteines. We chose three CCRs (1, 5, and 10) and three CXCRs (1–3) and analyzed their responses to a number of their multiple endogenous agonists (Table 1). G protein signaling was assessed by inhibition of forskolin-stimulated cAMP (a Gi/o response) using an antibody-based assay (12). A complementary approach to identifying biased ligands is to calculate the level of ligand bias as encapsulated by a bias factor. Bias factors are a method of quantifying ligand bias on a logarithmic scale; e.g., a bias factor of 1 between G protein- and βarr-mediated signaling signifies 10-fold more effective signaling through G proteins than βarrs (15). Limitation of this calculation is in cases of very low signaling through one pathway, which typically results in a poor fit with relatively high errors and thus a poorly determined bias factor. In such cases, a biased ligand can be identified by a bias plot. These complementary approaches allow a comprehensive analysis of ligand bias and identification of biased agonists.

The qualitative bias plot analysis suggested the presence of ligand bias in a number of chemokine receptors (highlighted by dashed black circles): CXCR2 (Fig. 2F), CXCR3 (Fig. 2G), CCR1 (Fig. 2B), and CCR10 (Fig. 2D). At CXCR2 (Fig. 2F), CXCL6 (blue) signals more effectively through G proteins over βar or internalization compared with the other ligands, consistent with it being a G protein-biased agonist. At CXCR3 (Fig. 2G), CXCL9 (magenta) appears to be relatively βarr-biased, whereas CXCL11 (brown) is biased toward internalization. At CCR1 (Fig. 2B), there is significant spread between a number of agonists in their responses to G proteins and βarrs, consistent with some level of G protein and βarr bias between those ligands. An interesting pattern is present in the comparison of βarr recruitment with internalization with CCL15 (green) and CCL23 (blue), demonstrating increased internalization compared with βarr recruitment, whereas CCL3 (magenta) and CCL5 (orange) demonstrate relatively decreased internalization compared with similar levels of βarr recruitment. At CCR10 (Fig. 2D), CCL27 (magenta) is clearly identified as a G protein-biased agonist compared with CCL28 (orange). Notably, at CXCR1 (Fig. 2E), the suggestion of increased internalization by CXCL8 compared with other signaling responses from simple pharma-
FIGURE 1. Multiple assays for assessment of chemokine signaling. Concentration-response data for G protein signaling (left column; cAMP signal demonstrating inhibition of forskolin-stimulated cAMP by Gαi/o), βarr recruitment (middle column), and receptor internalization (right column) for CCR1 (A), CCR5 (B), CCR10 (C), CCR1 (D), CXCR2 (E), and CXCR3 (F) are shown. Shown are means ± S.E. from at least three experiments and fits with logistic dose-response functions with Hill coefficient equal to 1 (Table 2). Error bars represent S.E.

TABLE 2
Pharmacologic parameters of signaling by chemokine receptors

Concentration-response data were fit with simple logistic equations with Hill coefficients equal to 1, allowing determination of maximal response (Emax) and EC50 for G protein signaling, βarr recruitment, and receptor internalization. Bias factors (β) were calculated as described under "Experimental Procedures." Bias factors that were statistically significantly different from the arbitrarily defined balanced agonist with a bias factor of 0 (p < 0.05 by unpaired t test) are in bold. Data from at least three experiments were used for the fits. S.E., standard error; N.D., not determined.
cologic parameters alone does not hold; rather, CXCL1 (magenta) and CXCL6 (orange) are balanced partial agonists compared with the full, balanced agonist CXCL8.

A quantitative analysis of ligand bias was then performed based on the pharmacologic fits from Table 2, allowing a calculation of bias factors (Table 2 and Fig. 3). Bias factors could be calculated for the majority of ligand-receptor combinations, although in a few situations they could not be determined because of low signaling in one of the pathways (Table 2 and Fig. 3). At half of the receptors tested (CCR1, CCR10, and CXCR3), there were statistically and likely physiologically significant levels of ligand bias observed. At CCR1 (Fig. 3A), CCL5 and CCL23 were identified as G protein-biased compared with CCL3 with CCL5, CCL15, and CCL23 displaying some levels of bias for or against internalization relative to H9252 recruitment. At CCR5 (Fig. 3B), CCL8 and met-CCL5 both show statistically significant H9252 bias compared with the reference agonist CCL3 although with bias factors less than 1. At CCR10 (Fig. 3C), because of the lack of H9252 recruitment by CCL28, most bias factors could not be calculated. At CXCR1 (Fig. 3D), no statistically significant levels of bias were observed, whereas at CXCR2 (Fig. 3E), some ligands displayed statistically significant levels of bias but all less than 1. At CXCR3 (Fig. 3F), there were no significant H9252- or G protein-biased agonists, but CXCL11 displayed significantly higher levels of internalization than CXCL9 or CXCL10.

**Ligand-dependent Differences in H9252 Recruitment and Receptor Internalization**—For those receptor-ligand systems (CCR1, CCR10, and CXCR3) that demonstrated evidence of significant ligand bias (i.e. an absolute value of any bias factor >1 was our cutoff for significant ligand bias), confocal microscopy was used to determine H9252 redistribution, receptor internalization, and H9252-receptor colocalization in response to ligand stimulation at a single, high concentration of ligand (100 nM) in HEK293 cells that were transiently transfected with H9252-GFP and receptor. As described below, this orthogonal assay confirmed the results of the luminescence-based H9252 redistribution and receptor internalization assays.

Both CCR1 (Fig. 4A and supplemental Fig. 1) and CXCR3 (Fig. 4C and supplemental Fig. 1) displayed constitutive activity with “class A” patterns of H9252 distribution with puncta at the membrane in cells transfected with both receptor and H9252, whereas H9252 cotransfected with CCR10 (Fig. 4B and supplemental Fig. 1) was uniformly distributed in the cytoplasm, consistent with minimal constitutive activity. Upon ligand stimulation, these receptors all showed some level of H9252 redistribution. At CCR1 (Fig. 4A and supplemental Fig. 1), stimulation with CCL3, -5, and -23 all resulted in H9252 redistribution from the class A pattern to a “class B” pattern of H9252 localization into endosomes with internalized receptor. Stimulation with CCL14 or CCL15 did not result in significant H9252 redistribution or receptor internalization. Notably, only CCL15 and CCL23 treatment resulted in significant receptor internalization with considerably less receptor at the plasma membrane, consistent with the luminescence-based receptor internalization assay. We also confirmed the results of the luminescence-
based receptor assay in an alternate clone that again demonstrated higher levels of receptor internalization induced by CCL23 compared with CCL3 (Fig. 5).

At CCR10 (Fig. 4B and supplemental Fig. 1), stimulation with CCL27 resulted in a class B pattern, whereas that with CCL28 resulted in a weak class A pattern, consistent with its weak response in the recruitment assay (only a 2-fold-response with an EC$_{50}$ of 300 nM). This difference in βar internalization was also reflected by significant receptor internalization with CCL27 that was absent in cells stimulated with CCL28. These findings were again consistent with the results of the luminescence-based receptor assays.

At CXCR3 (Fig. 4C and supplemental Fig. 1), CXCL9 and CXCL10 stimulation resulted in a class A pattern, whereas CXCL11 stimulation resulted in a class B pattern, consistent with its increased signal in the βar recruitment assay (Fig. 1F). CXCL9- and CXCL10-treated cells displayed more surface-expressed receptor than CXCL11-treated cells, whereas all samples displayed some internalized receptor, consistent with the known constitutive internalization of the receptor (20). Thus, with biased chemokines targeting the same receptor, we observed distinct patterns of βar redistribution and receptor internalization.

**Ligand Bias Is Associated with Distinct Physiologic Outcomes**—We then tested whether differences in bias between different ligands for the same receptor were associated with different chemotactic profiles in human peripheral blood mononuclear cells. Because the ligands tested for CCR1 were shared with a number of other receptors (1), we did not test it any further. CCR10 demonstrated a high degree of ligand bias between its two ligands, CCL27 and CCL28, both of which are capable of reaching maximal G protein signaling as assessed by inhibition of cAMP formation but only one of which, CCL27, is capable of

*FIGURE 3. Chemokine bias factors.* Bias factors between different pathways for each ligand were calculated as described; shown are mean ± S.E. from Table 2. Bias factors that were statistically significantly different from the balanced agonist are in **bold**. In the example of the G/arrestin bias factor, a positive bias factor denotes bias toward G, signaling, whereas a negative bias factor denotes bias toward βar recruitment. *N.D.,* not determined. *Error bars* represent S.E. See text for discussion.

*FIGURE 4. Biased ligands display distinct patterns of βar recruitment and receptor internalization.* At CCR1 (A) and CXCR3 (C), ligand stimulation results in a change (*Live cells*) from a class A pattern prior to ligand stimulation (*Pre*), consistent with weak constitutive activity, to differential class B patterns with βar internalization after ligand stimulation (*Post*). At CCR10 (B), there is little constitutive activity, and stimulation of the receptor results in a class A pattern. These are associated with distinct patterns (*Fixed cells*) of βar recruitment, receptor internalization, and colocalization (βar2-GFP; green; receptor, red; DNA, blue). See text for full discussion.
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Recruiting βarr and internalizing the receptor (Fig. 1C). CCL28 was clearly identified as a G protein-biased agonist in a qualitative assessment (Fig. 2) and was biased to such an extent that its bias factor could not be calculated (Fig. 3). This bias was confirmed in an assessment of βarr redistribution and receptor internalization (Fig. 4). Notably, although CCL28 is a lower potency agonist than CCL27 for G protein signaling, it displays similar potency with higher efficacy for monocyte migration compared with the balanced agonist CCL27.

At CXCR3, we tested the effects of its three ligands, CXCL9, CXCL10, and CXCL11, on peripheral blood mononuclear cell migration (Fig. 6B). These ligands did not display the same level of marked bias that CCL28 displayed at CCR10; rather, the ligands for CXCR3 displayed more complex behavior. CXCL11 appeared to be biased toward internalization (Figs. 2 and 3) and was also a more efficacious agonist in all pathways tested compared with CXCL9 and CXCL10 (Fig. 1). This activity is most consistent with the pattern of βarr recruitment by the different ligands, suggesting that at CXCR3 βarrs may play a role in mediating chemotaxis (21) and a smaller role in receptor desensitization because the most potent βarr binder was also the most potent chemotactic stimulus.

DISCUSSION

Biased agonism has been noted as a property of 7TMRs for over two decades (4) with one of the first descriptions being the functionally selective responses of the acetylcholine receptor to pilocarpine and carbachol (22). In general, biased agonists targeting 7TMRs have been generated synthetically and display distinct physiologic profiles from endogenous, balanced agonists, suggesting a role for these compounds as novel drugs (23, 24). However, it has been unclear whether biased agonism is an “accident” of 7TMR complexity that has only been exploited by synthetic drugs or whether it is a property that is utilized by endogenous systems as an added layer for specificity in signaling (25). One of the best characterized examples of endogenous ligand bias is at CCR7, a chemokine receptor with two ligands, CCL19 and CCL21, which both activate G protein signaling (26) but only one of which, CCL19, results in higher levels of βarr recruitment (26), GRK3-dependent βarr internalization, and receptor desensitization (27), which presumably results in a distinct physiological response as these two ligands play different roles in the immune system while acting through the same receptor. At CXCR4, there are distinct signaling patterns associated with monomeric or dimeric CXCL12 that can either promote or inhibit chemotaxis (28). Here we have demonstrated that in systems with multiple endogenous ligands biased agonism is a general, and likely conserved, mechanism for generating qualitatively distinct patterns of signaling via the same receptor.

Viewing previous studies through the paradigm of biased agonism yields a consistent model for receptor activation: different chemokines are capable of binding different sites of the receptor, thereby allosterically activating different signaling pathways. For example, CXCR3 ligands have been shown previously to bind at distinct sites of the receptor (29) that are also associated with signaling that requires different portions of the receptor for chemotaxis, calcium mobilization, and receptor internalization (30). Notably, our finding that βarr recruitment did not correlate with receptor internalization is consistent with the previous observation that CXCR3 internalization can be independent of clathrin and the βarrs (20), suggesting that at CXCR3 βarrs may act primarily as a positive regulator of signaling as opposed to an alternate pathway of constitutive receptor internalization and degradation (20). We did find, unlike an earlier study (31), that CXCL9 did stimulate βarr recruitment, although there was no apparent βarr bias observed at this receptor. The biology and pharmacology of CCR10 (32) are not as well characterized as that of CXCR3, and so the functional significance of the extreme bias between its two ligands is less clear. These findings clearly buttress the evolving model of multiple active states of the receptor associated with distinct signaling profiles (33).

Biased agonism therefore allows a single chemokine receptor to generate distinct physiologic responses in response to differ-
ent chemokines, thereby adding a layer of complexity to chemokine receptor signaling to augment the diversity attained with spatiotemporal expression. These distinct chemokine signaling profiles likely contribute significantly to the distinct phenotypes of knock-out mice to different ligands for the same receptor (34). Presumed biochemical redundancy has been considered to be a major hurdle in chemokine drug development, and although that may be true for antagonists, our findings suggest that biased agonists targeting these receptors would have distinct physiologic effects. It also suggests that post-translational processing of chemokines, such as cleavage of N-terminal residues (35), may not just change the relative agonism of a molecule but also its relative bias. Thus, the presence of endogenously biased agonists has far ranging impact on our understanding of chemokine pharmacology and biology.

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