Deciphering and modulating G protein signalling in C. elegans using the DREADD technology

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G-protein signalling is an evolutionary conserved concept highlighting its fundamental impact on developmental and functional processes. Studies on the effects of G protein signals on tissues as well as an entire organism are often conducted in Caenorhabditis elegans. To understand and control dynamics and kinetics of the processes involved, pharmacological modulation of specific G protein pathways would be advantageous, but is difficult due to a lack in accessibility and regulation. To provide this option, we designed G protein-coupled receptor-based designer receptors (DREADDs) for C. elegans. Initially described in mammalian systems, these modified muscarinic acetylcholine receptors are activated by the inert drug clozapine-N-oxide, but not by their endogenous agonists. We report a novel C. elegans-specific DREADD, functionally expressed and specifically activating G_q-protein signalling in vitro and in vivo which we used for modulating mating behaviour. Therefore, this novel designer receptor demonstrates the possibility to pharmacologically control physiological functions in C. elegans.

Designer receptors exclusively activated by designer drugs (DREADDs) have been developed as a tool to study and to specifically manipulate G-protein signalling in vivo. This technology for the selective control of signalling cascades has been found to be invaluable not only to investigate the impact of specific G-protein signalling on distinct physiological processes but also to control cellular functions in a cell-type specific manner. Besides DREADDs as pharmacogenetic tools also optogenetic methods have been generated. Both approaches are non-invasive and based on modified receptors which can be specifically activated – DREADDs exclusively by synthetic ligands and optogenetic tools only by light.

The first generation of DREADDs was based on muscarinic acetylcholine receptors which have been altered by two point mutations within transmembrane domains 3 and 5 to inhibit activation by the endogenous agonist acetylcholine (ACH) but allowing stimulation by the inert compound clozapine-N-oxide (CNO). Further modification allowed for specific activation of G_q, G_s, or G_i signalling pathways, respectively. These DREADDs are called rM3Dq, rM3Ds, and hM4Di by convention. They have been extensively pharmacologically characterized and widely used to specifically activate and inactivate for instance neuron populations in vivo by employing the G_q- and the G_i-specific DREADDs, respectively. DREADDs have also been involved to study neuronal impact on fear memory, Parkinson disease, Down syndrome, and the role of glial cells in vivo. In the periphery, the DREADD technology has been applied to study metabolic functions in islet β-cells and hepatocytes.

Despite the power and simplicity in the use of DREADDs many physiological processes and functions associated with specific signalling cascades are not easy to delineate in mammals due to the complexity of the organisms. The application in less complex systems would be highly advantageous to overcome these limitations. Recently, the DREADD technology was transferred to Drosophila melangoster showing that mammalian DREADDs can be efficiently expressed in the fruitfly and modulate physiological functions linked to G-protein pathways. The use of mammalian DREADDs in other organisms also raises the interesting question whether organism-specific DREADDs can be generated likewise. The nematode Caenorhabditis elegans, a very simple model organism frequently employed for the investigation of various biological questions, is highly suitable for the dissection of the impact of signalling cascades. So far only optogenetics have been successfully utilised as a
tool for the manipulation of some signalling cascades in the nematode12–14. However, G protein-signalling pathways are not well addressed by non-genetic tools.

Here, we show that the DREADD technology can be applied in C. elegans using a newly designed DREADD based on the nematode muscarinic receptor GAR-3b. In-depth pharmacological characterisation in vitro revealed that the DREADD can be activated by Oxo M and CCh to induce spicule protraction involving the Gq protein-signalling cascade. This effect is independent of the presence of a hermaphrodite. (C) In the absence of GAR-3 neither Oxo M nor CCh trigger spicule protraction. (D) DREADDs activating the Gq protein-signalling cascade specifically trigger spicule protraction upon stimulation with CNO, but not with Oxo M or CCh. Such DREADDs can be rM3Dq or a C. elegans-specific DREADD (cegar-3Dq) in gar-3-deficient males. (E) Protraction rate in male nematodes containing mammalian DREADD constructs. Males were incubated with 100 mM Oxo M, 10 mM CCh, 2 mM CNO or H2O as negative control. Subsequently, spicule protraction was scored and protraction rates calculated in respect to the total male count. Wild-type worms (pha-1(e2123); him-5(e1490)), gar-3 (pha-1(e2123); him-5(e1490) gar-3(gk305)) and transgenic gar-3; Ex[gar-3(+)] males served as controls. Data are shown as mean ± SD. ***p < 0.001; n ≥ 250. Indicated below each set of columns is the schematic model related to the respective data.

Figure 1. The mammalian rM3Dq DREADD and rM3R are not functional in spicule protraction, as a model for DREADD activation in C. elegans. (A–D) Schematic representation of spicule protraction and theoretical model of GAR-3 and DREADD activation in this context, that all DREADD characterisations are based on (see Figs 1E and 4F). (A) GAR-3 mediates a Gq-protein signal which is involved in spicule protraction of the male nematode during mating culminating in insertion of the spicule into the hermaphrodite's vulva. (B) GAR-3 can be activated by Oxo M and CCh to induce spicule protraction involving the Gq protein-signalling cascade. This effect is independent of the presence of a hermaphrodite. (C) In the absence of GAR-3 neither Oxo M nor CCh trigger spicule protraction. (D) DREADDs activating the Gq protein-signalling cascade specifically trigger spicule protraction upon stimulation with CNO, but not with Oxo M or CCh. Such DREADDs can be rM3Dq or a C. elegans-specific DREADD (cegar-3Dq) in gar-3-deficient males. (E) Protraction rate in male nematodes containing mammalian DREADD constructs. Males were incubated with 100 mM Oxo M, 10 mM CCh, 2 mM CNO or H2O as negative control. Subsequently, spicule protraction was scored and protraction rates calculated in respect to the total male count. Wild-type worms (pha-1(e2123); him-5(e1490)), gar-3 (pha-1(e2123); him-5(e1490) gar-3(gk305)) and transgenic gar-3; Ex[gar-3(+)] males served as controls. Data are shown as mean ± SD. ***p < 0.001; n ≥ 250. Indicated below each set of columns is the schematic model related to the respective data.

Results

Mammalian DREADDs are not active in C. elegans. One prerequisite for the use of DREADDs in C. elegans is that the synthetic ligand CNO utilised to activate these receptors does not have any adverse effects on nematodes. To elucidate the influence the compound has on C. elegans fertility, development, viability, and the neuronal system as well as certain aspects of behaviour we treated wild-type nematodes in liquid culture with varying concentrations of CNO and assayed brood size, individuals reaching adulthood, lifespan, locomotion, pharyngeal pumping, egg laying and sensitivity to aldicarb. However, none of the parameters was affected (Supplementary Fig. 1) indicating that the compound does not have any major side effects on C. elegans.

As mammalian DREADDs have been shown to be a useful tool for analyses in Drosophila, suggesting that they are able to activate distinct G-protein cascades in vertebrates and invertebrates likewise15, we tested these receptors in C. elegans. To investigate the functionality of mammalian DREADDs in the nematode, systems are required in which the physiological implications of Gq, Gs, and Gi protein-mediated pathways, respectively, are well understood. Protraction of copulatory spicules from the tail of the male nematode is dependent on a Gq signalling cascade15,16 and thus, offers one suitable read-out. Spicule protraction occurs during mating when the spicule is inserted into the hermaphrodite’s vulva (Fig. 1A)15,17. The C. elegans homolog of the muscarinic acetylcholine receptor M3, GAR-3, is a G protein-coupled receptor (GPCR) involved in controlling this process. GAR-3 has been shown to activate a Gq cascade similar to its mammalian homolog18–20 and even triggers G-protein signalling in mammalian cells20 indicating that this
neither cholera toxin nor pertussin toxin, a strong inhibitor of Gi-protein signalling 31, altered signalling on performed with pertussis and cholera toxin to test for coupling to other G protein-signalling pathways. Tab. 3). For further validation of the G protein-coupling properties of GAR-3b, DMR measurements were ties in properly expressing or processing the receptors.

GAR-3b (Fig. 2A), but only Y3.33 is conserved in C. elegans with an HA-epitope to allow for detection using commercially available antibodies. Therefore, we chose GAR-3b for generation of a C. elegans-specific DREADD and – analogously to mammalian DREADDs – mutated amino acids Y3.33 (Y146C) and A5.46 (A237G). The DREADD was N-terminally tagged with the muscarinic acetylcholine receptors 21,23,24, whereas GAR-3 is pharmacologically very similar to mammalian M3R and activates the Gq signalling cascade upon stimulation with ACh or CCh 19–21. Further, alignment analyses revealed an overall amino acid sequence identity of Gq-3 to M3R of 32.8%. Two splice variants, GAR-3a and GAR-3b, have been described with GAR-3b being the predominantly occurring form. The two isoforms only differ in 26 amino acid length within the 3rd intracellular loop (ICL) 21. Also, amino acids Y3.33 and A5.46 (universal amino acid numbering system 21), which are widely conserved among muscarinic acetylcholine receptors, are also present in the C. elegans GAR-3b (Fig. 2A), but only Y3.33 is conserved in C. elegans GAR-2 (Fig. 2B). Therefore, we chose GAR-3b for generation of a C. elegans-specific DREADD. Since mammalian DREADDs did not seem to be properly expressed in C. elegans, we set out to generate a nematode-specific DREADD for modulation of G protein-signalling pathways in the worm. We sought to base this modified receptor on a muscarinic acetylcholine receptor similar to the mammalian DREADDs. Three G protein-linked acetylcholine receptor genes are known in C. elegans (gar-1, gar-2, gar-3) with GAR-2 being described to bind Gi/Gs proteins and GAR-3 being a Gq-coupling receptor 21,22. However, GAR-1 and GAR-2 differ in their pharmacological properties from mammalian muscarinic acetylcholine receptors 21,23,24, whereas GAR-3 is pharmacologically very similar to mammalian M3R and activates the Gq signalling cascade upon stimulation with ACh or CCh 19–21. Further, alignment analyses revealed an overall amino acid sequence identity of Gq-3 to M3R of 32.8%. Two splice variants, GAR-3a and GAR-3b, have been described with GAR-3b being the predominantly occurring form. The two isoforms only differ in 26 amino acid length within the 3rd intracellular loop (ICL) 21. Also, amino acids Y3.33 and A5.46 (universal amino acid numbering system 21), which are widely conserved among muscarinic acetylcholine receptors, are also present in the C. elegans GAR-3b (Fig. 2A), but only Y3.33 is conserved in C. elegans GAR-2 (Fig. 2B). Therefore, we chose GAR-3b for generation of a C. elegans-specific DREADD – analogously to mammalian DREADDs – mutated amino acids Y3.33 (Y146C) and A5.46 (A237G). The DREADD was N-terminally tagged with an HA-epitope to allow for detection using commercially available antibodies.

Pharmacological characterisation of ceGAR-3Dq. The modified GAR-3b was pharmacologically characterized in respect to agonist and coupling specificity. Due to the high conservation of the three major G proteins Gq, Gi, and Gs among metazoan species analyses of functional G protein-coupling of receptors from C. elegans or other invertebrate species can be performed using mammalian systems 27–29 with the results being transferable to the original system. Thus, we used transiently transfected COS-7 cells, as DREADD cell surface expression was highest compared to CHO-K1 and HEK-293GT cells (Supplementary Fig. 3A). Utilising label-free dynamic mass redistribution (DMR) experiments, we demonstrated that GAR-3b is only activated by CCh (Fig. 3A) whereas the receptor mutant (ceGAR-3Dq) is indeed a DREADD solely activated by the inert drug CNO but not by the muscarinic agonist CCh (Fig. 3B). CNO activated the DREADD in a concentration-dependent manner with an EC50 value of 3.5 μM, whereas the muscarinic agonist CCh did not have an effect on receptor activity (Fig. 3C). To analyse the coupling specificity, further experiments involved detection of second messengers such as inositol phosphate (IP) as a read out for receptor activation. Consistent with the DMR measurements IP accumulation assays showed an increase in IP formation upon CCh stimulation of GAR-3b but not upon CNO treatment (Fig. 3D). In contrast, DREADD-mediated IP formation occurs only after stimulation with CNO. Further, measurement of calcium release also demonstrates the concentration-dependent activation of the DREADD-receptor upon CNO treatment (Fig. 3E), whereas GAR-3b is only activated by CCh (Fig. 3F). Neither of these compounds has any effect on mock-transfected cells demonstrating the receptor-specificity (Supplementary Fig. 3B).

This is consistent with previous studies where GAR-3-transfected CHO-K1 cells were shown to accumulate inositol phosphates and release Ca2+ upon CCh treatment 19,21,23,30. Further, these elevated Ca2+ levels have been shown to stimulate cAMP production independent of Gq-protein-signalling 20. Also, it was demonstrated that GAR-3 is linked to Gq–protein coupling in cell culture experiments 20,25,29. Thus, we tested for potential Gq- or Gi-protein coupling. Firstly, we applied second messenger assays to measure cAMP accumulation with and without pre-incubation of the adenylyl cyclase inhibitor forskolin. Neither stimulation of GAR-3b nor of ceGAR-3Dq resulted in a change of cAMP accumulation (Supplementary Fig. 4A,B). However, it has to be noted that expression levels of GAR-3b and ceGAR-3Dq were rather low. Therefore, the lack in identifying Ca2+-mediated cAMP formation might be due to a lack of sensitivity. In fact, in a more sensitive CRE-based reporter gene assay an increase of cAMP was observed (Supplementary Fig. 4C,D). The EC50 values are similar to those obtained in DMR measurements (CNO at ceGAR-3Dq: 1.6 μM; CCh at GAR-3b: 4.9 μM) (Supplementary Tab. 3). For further validation of the G protein-coupling properties of GAR-3b, DMR measurements were performed with pertussis and cholera toxin to test for coupling to other G protein-signalling pathways. Neither cholera toxin nor pertussin toxin, a strong inhibitor of Gi–protein signalling 31, altered signalling on CCh-stimulated GAR-3b transfected cells suggesting no involvement of toxin-sensitive Gq- or Gi–protein pathways (Supplementary Fig. 4E,G). These results indicate exclusive Gq–protein coupling of the DREADD and according to the suggested nomenclature the DREADD was therefore named ceGAR-3Dq.3.
Gq-protein signalling triggered by ceGAR-3Dq modulates spicule protraction behaviour in male nematodes. We next tested the applicability of the GAR-3-based DREADD ceGAR-3Dq in vivo. To ensure that the receptor is functional we first assessed expression of ceGAR-3Dq::yfp based on genomic gar-3 driven by the gar-3 promoter which is indistinguishable from gar-3::yfp, albeit appearing to be generally weaker (Fig. 4A–E). Subsequently, the capability of the DREADD to activate a Gq-protein-signalling cascade when stimulated with CNO was investigated by quantifying spicule protraction movements (see Fig. 1D). Transgenic gar-3 (gk305); Ex[cegar-3Dq] nematodes displayed no spicule protraction upon treatment with Oxo M or CCh (Oxo M 17.9 ± 11.5%; CCh 13.6 ± 4.8%) similar to the effect seen in untreated males (16.8 ± 2.1%; Figs 1B–D and 4F, Supplementary Tab. 1B), indicating that the DREADD is not activated by GAR-3 muscarinic agonists. However, spicule protraction was triggered upon stimulation with the synthetic DREADD agonist CNO (37.4 ± 6.3%) (Figs 1D and 4F, Supplementary Tab. 1B). These data are consistent with the pharmacological analyses in vitro showing that ceGAR-3Dq is a DREADD. To optimise the observed effect of ceGAR-3Dq-activated Gq-protein signalling on spicule protraction, a dose-response curve using increasing concentrations of CNO was conducted (Fig. 4G). However, due to the limited solubility of the compound, a maximal concentration of 2 mM CNO was tested. Unfortunately, stimulation did not reach saturation and thus, we were unable to determine an EC50 value. Approximately 40% of males protracted their spicules at this concentration. Activation of the DREADD and thus, spicule protraction was already detected 2 minutes after start of CNO treatment. Treatment duration did not have any influence on the extent of spicule protraction (Fig. 4H).

We next asked whether DREADD activation can be temporally controlled. Upon removal of gar-3(gk305); Ex [cegar-3Dq] nematodes from CNO the spicule protraction rate decreased over time (Fig. 4I). Approximately 30 minutes post drug withdrawal spicule protraction reached the same level as observed in untreated controls, suggesting that DREADD activation is reversible. These results indicate that ceGAR-3Dq can be specifically activated by CNO in vivo and is able to mediate a Gq-protein signalling cascade. This activation can be temporally controlled.

Figure 2. Amino acid alignment of transmembrane domains 3 and 5 of muscarinic acetylcholine receptor orthologs. (A) Sequences of mAChR type 3 from 13 different species were retrieved from publically available databases such as NCBI/GenBank. Alignment was performed using the Clustal W alignment. Shown is the amino acid sequence of transmembrane domain 3 and 5. Conserved residues are shaded in blue. Highlighted in red are the highly conserved tyrosine Y3.33 (Y146) and alanine A5.46 (A237) which are mutated in mammalian DREADDs. (B) Analogously, alignment of mAChR type 2 was performed. While Y3.33 is also conserved in C. elegans GAR-2, amino acid 5.46 differs from other muscarinic orthologs.
Discussion

GPCRs are the largest family of cell surface receptors and the signals they mediate are involved in nearly all physiological functions. Thus, gaining an in-depth understanding of their signalling modes and the ability to
Figure 4. Cegar-3Dq is functional upon stimulation with CNO in *C. elegans*. (A–D) Expression and protein localisation of cegar-3Dq::yfp is indistinguishable from gar-3::yfp, both driven by the gar-3 promoter. Expression was detected in the pharynx (A, B; 1./3. row: DIC, 2./4. row: fluorescent image) and in the male tail (C, D; 1./3. row: DIC, 2./4. row: fluorescent image). Localisation of the DREADD appears to be similar to the one of GAR-3. Scale bars = 10 μm. (E) Analyses of expression levels by quantification of intensities reveals that cegar-3Dq::yfp expression is significantly weaker than gar-3 expression. For quantification, images of the pharynx from
specifically modulate the signalling pathways has been intensively studied. However, determining the impact of a single type of receptor and its signalling pathways in a distinct cell type or tissue is virtually impossible as a single receptor is usually present in more than one cell type and endogenous agonists activate more than one receptor31. For the same reasons it is problematic to modulate distinct signalling cascades in a specific cell type in order to alter or control cellular functions. To overcome those limitations pharmacogenetic techniques in the form of designer receptors have been developed. These so-called DREADDs are only activated by the inert compound CNO but not by the endogenous ligand ACh and thus, are a valuable tool to specifically activate G protein-signalling pathways in vivo. Combining this technique with the use of simple model organisms would render a powerful tool for the investigation and control of cell-type specific signalling.

In the present study, we generated a DREADD for the nematode C. elegans based on the G protein-linked acetylcholine receptor gar-3. This DREADD couples to the Gα protein-signalling cascade similar to GAR-3b and is activated exclusively by CNO. As a proof of principle, we provide functional data that the DREADD can be employed for spatiotemporal control of signalling in the nematode.

The DREADD technology is highly suitable for the use in the model organism C. elegans and genetically amenable so that DREADD constructs can be easily introduced. Further, we have shown that the synthetic compound CNO which activates DREADDs can be administered by feeding or soaking nematodes without having any major detrimental effects.

Although expression of selected mammalian GPCRs in C. elegans has been shown to be generally possible for some receptors34, introduction of mammalian DREADDs rM3Dq, rM3Dv, and hM4Di as well as rM3R in the nematode did not render any functional receptors, despite their high sequence and functional conservation among various species. Our data indicate that these receptors are possibly not properly expressed in C. elegans. Several factors may contribute to this fact. As transgenic lines containing the receptors were obtained, a potential toxic effect of the constructs or their gene products can be ruled out. However, processing or membrane targeting of the mammalian proteins might be an issue. Alternatively, it is conceivable that certain co-factors or interaction partners required for functional expression are not present in C. elegans. Even though the codons of the mammalian cDNA sequences were reviewed prior to construct generation and no overly problematic ones were identified, it cannot be excluded that non-optimal codon usage might account for this effect. It is, however, conceivable that these receptors are transcribed but not processed properly in nematode cells.

As the technology of mammalian DREADDs was not easily transferable to C. elegans, we sought to generate a nematode-specific one and have chosen the M3R homolog GAR-3. While the ceGAR-3Dq transgene used for the in vivo studies is based on the genomic locus of gar-3, cDNA of the predominantly occurring GAR-3b isoform was used for its initial pharmacological characterisation. This isoform does not differ from GAR-3a in its pharmacological properties, which is also similar to rat M3R where large parts of the 3rd intracellular loop can be deleted without changing receptor functionality35. In both cases we inserted two point mutations according to mammalian DREADDs in transmembrane domains 3 and 5 of the receptor (Y146C/A237G in GAR-3) for generation of the novel DREADD. DMR measurements demonstrated that the modified GAR-3b is indeed a DREADD, only activated by CNO but not by the muscarinic agonist CCh (Fig. 3). The EC50 value for CNO activation determined in DMR measurements is 3.5 μM (pEC50: 5.46 ± 0.43) which is about 100-fold higher than for the mammalian DREADD rM3Dq determined in yeast or cell culture experiments34,35.

This lower potency of CNO at the C. elegans DREADD is probably due to the lower potency of agonists at GAR-3b compared to rM3R (Supplementary Tab. 3). In vitro analyses of coupling properties of GAR-3b and ceGAR-3Dq revealed a robust Gαi-mediated signalling with a Ca2+-dependent increase in cAMP37. Similar results have been obtained in previous studies38.

Although it cannot be easily proven that the GAR-3-based DREADD exclusively activates Gαi signalling cascades especially as there are 21 Gα subunits in C. elegans with 17 being uncharacterized in regards to their signalling abilities as no clear homologs exist38,39, our functional assays point towards ceGAR-3Dq activating predominantly Gαi signalling.

In vivo analyses revealed that the GAR-3b-derived DREADD ceGAR-3Dq is able to mediate a Gαi-protein signalling cascade upon stimulation specifically with CNO but not with CCh or Oxo M. Activation of different cegar-3Dq::yfp expressing strains were taken and analysed in comparison with images from nematodes carrying gar-3::yfp. (F) Protraction rate in male nematodes containing the cegar-3Dq construct. Males were stimulated with 100 mM Oko M, 10 mM CCh, 2 mM CNO or H2O as negative control. Subsequently, spicule protraction was scored. Wild-type worms (pha-1(e2123); him-5(e1490); gar-3(pha-1(e2123); him-5(e1490) gar-3(gk305)) and transgenic gar-3; Ex[gar-3(+)] males served as controls. Data are shown as mean ± SD. n.s. not significant; **p < 0.01; ***p < 0.001; n ≥ 300. (G) Spicule protraction rates are dependent on the concentration of CNO. Male nematodes were incubated with the indicated concentrations of CNO and spicule protraction was scored subsequently. Nematodes deficient for gar-3 (pha-1(e2123); him-5(e1490) gar-3(gk305)) served as negative control. Data are shown as mean ± SD, **p < 0.01; ***p < 0.001; n ≥ 200. (H) Treatment duration does not have an effect on spicule protraction rate after 3 minutes. Male nematodes were incubated with 2 mM CNO and spicule protraction was scored. As negative control, nematodes deficient for gar-3 (pha-1(e2123); him-5(e1490) gar-3(gk305)) were used. Data are shown as mean ± SD, **p < 0.01; ***p < 0.001; n ≥ 200. (I) The effect of DREADD activation is reversible. Transgenic gar-3; Ex[gar-3(+)] males were stimulated with 2 mM CNO. After 3 minutes the spicule protraction rate was determined (time point termed “0 minutes after CNO withdrawal”) and nematodes were withdrawn from the drug by transferring them into M9. Subsequently, spicule protraction was measured at distinct time points. Data are shown as mean ± SD, n ≥ 150.
ceGAR-3Dq led to spicule protraction in males lacking gar-3 showing that the DREADD can complement the GAR-3-signalling cascade when activated. The detected effect is approximately 40% while 80% are achieved when re-introducing the gar-3 genomic sequence into gar-3-deficient nematodes and activating the receptor with CCh or Oxo M. Several factors may account for this reduced efficacy. Firstly, activation of both receptors occurs upon stimulation with different compounds. Although the EC₅₀ values for both are within a similar range in vitro, nematodes were incubated in lower concentrations of CNO compared to Oxo M or CCh due to its limited solubility. Moreover, it is well possible that different drug uptake levels or limited drug accessibility of CNO can explain the observed differences in vivo. Obviously, this potential limitation in drug accessibility needs to be kept in mind when using DREADDs for applications in different types of tissue in C. elegans. Secondly, lower expression levels of the DREADD compared to gar-3 (Fig. 3) might also contribute to ceGAR-3Dq having a smaller effect on spicule protraction rate than GAR-3. However, despite these obvious limitations ceGAR-3Dq is useful for modulating G₉-protein-signalling in C. elegans for certain applications.

The use of this novel ceGAR-3Dq in particular and DREADDs in general in the model organism C. elegans is highly advantageous for various applications. It can be employed for studying the impact of G-protein pathways in distinct cell-types or organs on the whole organism, an approach for which mammalian systems sometimes are too complex. Many cell-specific promoters have been described and offer a platform to express DREADDs in specific tissues. Further, it is highly suitable to manipulate the activity of certain cell types with neuroscience being one potential field of application. The neuronal network of C. elegans offers an ideal system for studies on neuronal networks and connectomes as it is well characterised. A hermaphrodite contains 302 neurons, which form about 7,000 chemical synapses. Probing specific populations of neurons and stimulating or inhibiting their activity will help understanding neuronal networks and subsequently, for instance animal behaviour. So far, optogenetic tools have been employed to address these topics. DREADDs are a powerful alternative to this technique as they allow for a dose-dependent modulation of signalling activity.

In summary, our results show that DREADDs are a functional tool in the nematode C. elegans. The newly designed nematode-specific DREADD ceGAR-3Dq based on the G protein-coupled acetylcholine receptor GAR-3 activates a G₉ protein-signalling cascade and is able to modulate related pathways in vivo. This receptor is exclusively stimulated by the inert compound CNO and thus, does not interfere with any physiological receptors. This pharmacogenetic toolbox now established in C. elegans offers a plethora of areas of application. To fully explore the entire experimental potential of DREADDs in C. elegans future analyses will need to focus on the design of nematode DREADDs specific for activating other G protein cascades such as G₉ and G₁ protein-signalling pathways.

Methods

Materials. Carbamylcholine chloride (carbachol, CCh), Oxotremorine M (Oxo M), forskolin, 3-isobutyl-1-methyloxanthine (IBMX), aldicarb, and standard chemicals were purchased from Sigma Aldrich. Clozapine-N-Oxide (CNO) was obtained from the National Institutes of Health (Bethesda, MD) as part of the Rapid Access to Investigative Drug Program funded by the National Institute of Neurological Disorders and Stroke. Substances applied were dissolved in the media or buffer the according experiment was performed in.

Generation of plasmids and transgenes. GAR-3b constructs for in vitro assays were generated by cloning gar-3b cDNA into the mammalian expression vector pcDNA5/FRT and contained a N-terminal HA-tag which does not alter functional properties of GPCRs (pSP109). cegar-3Dq was engineered by introducing the two point mutations Y146C/A237G into pSP109 (pSP112). For generation of constructs for in vivo analyses recombineering was employed, gar-3:yyfp (genomic) inserted behind the gar-3 promoter (pYL9) was a kind gift from Dr. Rene Garcia, Texas A&M University. The construct containing the C. elegans-specific DREADD cegar-3Dq downstream the gar-3 promoter (pSP110) was generated by inserting the two point mutations (Y146C/A237G) into the genomic sequence of gar-3 in the vector pYL9 (kind gift from Dr. Rene Garcia, Texas A&M University). Constructs comprising gar-3 promoter-driven mammalian DREADDs (pSP106, pSP108, pSP114) and rat M3R (pSP104) were cloned by exchanging the cDNA of the respective receptor in pYL9. For details see Supplementary Methods.

Cell culture. CHO-K1 and COS-7 cells were purchased from Leibniz Institute DSMZ – German Collection of Microorganisms and Cell Culture GmbH (CHO-K1: No. ACC 110, COS-7: No. ACC 60), HEK-GT cells were obtained from ThermoFisher Scientific (GripTite 293 MSR Cell Line, No. R79507). All cells were grown in F12 media or Dulbecco’s modified Eagle medium (DMEM) supplemented with 10% fetal bovine serum, 100 U/ml penicillin, and 100 μg/ml streptomycin at 37 °C in a humidified incubator with 7% CO₂. Lipofectamine 2000 (Life Technologies) was used for transient transfection according to the manufacturer’s recommendations.

Dynamic mass redistribution. Dynamic mass redistribution measurements are a label free assay system to study GPCR properties in living cells. Thus, COS-7 cells were seeded into T25 flask 16 to 24 hours prior to transfection with 3 μg plasmid DNA and 7.5 μl Lipofectamine. The next day, 6,000 cells were split into one well of an Epic 384-Well Fibronectin-Coated Microplate (Corning, Kaiserslautern, Germany). DMR measurements were carried out at a Corning Epic label-free detection platform. CCh and CNO were dissolved in HBSS buffer at different concentrations. For equilibration, cells incubated with HBSS buffer for 2 hours, followed by a baseline recording for 5 minutes. After adding CCh and CNO DMR was recorded for up to 75 minutes. To determine G protein-coupling specificity cells were incubated either with pertussis (100 ng/ml) or cholela toxin (1 μg/ml) overnight prior to DMR recordings.
**IP accumulation assay.** To measure inositol phosphate (IP) formation COS-7 cells were split into 12-well plates (1.5 × 10^5 cells/well) and transfected with a total amount of 0.5 μg of plasmid DNA and 1.5 μl Lipofectamine per well. Subsequently, cells were incubated with 2 μCl/ml of [myo-3H]-inositol (18.6 Ci/mmol, Perkin Elmer) for 18 hours. Thereafter, cells were washed once with serum-free DMEM containing 10 mM LiCl followed by incubation with test compounds for 30 minutes at 37 °C. Intracellular IP levels were determined by anion-exchange chromatography as described previously.45

**Fluorometric Calcium measurements.** COS-7 cells were split into T25-flasks (1.2 × 10^6) and transfected with 3 μg plasmid DNA and 7.5 μl Lipofectamine the next day. 48 hours post transfection, cells were detached using Versene and labelled in DMEM with 4 μM Fluo-4 AM (Molecular probes) for 30 minutes at 37 °C. Free dye was removed by centrifugation, and cell suspensions were re-suspended into HBSS buffer (132 mM NaCl, 10 mM HEPES, 6 mM KCl, 5.5 mM glucose, and 1 mM MgCl₂ adjusted to pH 7.4 with NaOH). The cell suspension was dispensed into black, clear-bottom 384 microwell plates (Corning, Kaiserslautern, Germany) with 45,000 cells per well. Fluorescence measurements were performed in a two-step protocol using a fluorescence imaging plate reader and a robotic liquid handling station (Freedom Evo 150, Tecan, Männedorf, Switzerland). Fluorescence intensity was corrected for the background and normalized to the initial intensities (F/F₀).

**C. elegans strains.** C. elegans variety Bristol, N2 and grown at 22 °C. The allele gar-3(gk305) was generated by the C. elegans gene knockout consortium. The strain pha-1(e2123); him-5(e1490); gar-3(gk305) was previously described (kind gift from Dr. Rene Garcia, Texas A&M University) and kept at 15 °C. The transgenes aprEx-183[gar-3;ypf(pYL9)]pha-1(+)pBSK, aprEx184[cegar-3;3Dq::yfp(pSP110)pha-1(+)pBSK], aprEx186[RM3R::yfp(pSP104)pha-1(+)pBSK], aprEx187[RM3Dq::yfp(pSP106)pha-1(+)pBSK] aprEx188[RM3D::yfp(pSP108)pha-1(+)pBSK] and aprEx189[hraM4DC::yfp(pSP114)pha-1(+)pBSK] were generated for this study (for details see Supplementary Methods) and cultivated at 25 °C.

**Spicule protraction assay.** 24 hours prior to conducting the assay, wild-type L4 males were put separately onto NGM agar plates containing Escherichia coli OP50. Subsequently, males were mounted onto a 2% agarose pad and a 100 μl of 2 mM CNO, 100 mM Oxo M, 10 mM CCh or H₂O, respectively, were applied. Males were scored for spicule protraction using a Leica M165FC microscope.

**Microscopy.** For analysis of transgene expression adult males were mounted in M9 containing 250 μM levasimole onto a 2% agarose pad. Differential interference contrast (DIC) and confocal fluorescent images were collected with an Olympus Fluoview FV1000 setup. Fluorescence signals were quantified by intensity analysis using ImageJ software.

**Statistical analyses.** Statistical significance of assay data from spicule protraction assays was determined using a Fisher’s exact test for each genotype and condition. For statistical analyses of brood size, individuals reaching adulthood, lifespan, locomotion, pharyngeal pumping, egg laying, sensitivity to aldicarb, and fluorescent intensity calculations a student’s t-test was performed.

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S.P. and D.T. conceived the project, designed the experiments, wrote the manuscript, and prepared the figures. S.P., F.F., C.B. and J.W. carried out C. elegans experiments. D.T. carried out the pharmacological experiments. T.S. contributed to the manuscript. All authors reviewed the manuscript.

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