KLHL12 Promotes Non-Lysine Ubiquitination of the Dopamine Receptors D\textsubscript{4,2} and D\textsubscript{4,4}, but Not of the ADHD-Associated D\textsubscript{4,7} Variant

Kamila Skieterska, Pieter Rondou\textsuperscript{a}, Béatrice Lintermans\textsuperscript{a}, Kathleen Van Craenenbroeck\textsuperscript{*}

Laboratory of GPCR Expression and Signal Transduction (L-GEST), Ghent University, Ghent, Belgium

\textsuperscript{a} Current address: Center for Medical Genetics Ghent, Ghent University, Ghent, Belgium

\textsuperscript{*} kathleen.vancraenenbroeck@ugent.be

Abstract

Dopamine D\textsubscript{4} Receptor Polymorphism

The dopamine D\textsubscript{4} receptor has an important polymorphism in its third intracellular loop that is intensively studied and has been associated with several abnormal conditions, among others, attention deficit hyperactivity disorder.

KLHL12 Promotes Ubiquitination of the Dopamine D\textsubscript{4} Receptor on Non-Lysine Residues

In previous studies we have shown that KLHL12, a BTB-Kelch protein, specifically interacts with the polymorphic repeats of the dopamine D\textsubscript{4} receptor and enhances its ubiquitination, which, however, has no influence on receptor degradation. In this study we provide evidence that KLHL12 promotes ubiquitination of the dopamine D\textsubscript{4} receptor on non-lysine residues. By using lysine-deficient receptor mutants and chemical approaches we concluded that ubiquitination on cysteine, serine and/or threonine is possible.

Differential Ubiquitination of the Dopamine D\textsubscript{4} Receptor Polymorphic Variants

Additionally, we show that the dopamine D\textsubscript{4,7} receptor variant, which is associated with a predisposition to develop attention deficient hyperactivity disorder, is differentially ubiquitinated compared to the other common receptor variants D\textsubscript{4,2} and D\textsubscript{4,4}. Together, our study suggests that GPCR ubiquitination is a complex and variable process.

Introduction

Dopamine is an important neurotransmitter in the mammalian brain that controls many basic processes, such as reward-motivated behavior, emotion, movement, sexual behavior and endocrine regulation, by binding to the specific receptors at the postsynaptic and presynaptic
Dopamine receptors (DR) belong to the superfamily of G protein-coupled receptors (GPCR) and are divided into two classes: dopamine D1-like receptors (including D1R and D5R) that bind to Gs proteins and activate adenylyl cyclase and dopamine D2-like receptors (including D2R, D3R and D4R) that inhibit adenylyl cyclase via Gi/o proteins [1–3].

The dopamine D4 receptor (D4R) has an important polymorphism in its third intracellular loop (IC3) where a 16-amino acid sequence is repeated 2 to 11 times resulting in different receptor variants referred to as D4.2R to D4.11R [4]. Although many variants exist, three of them are the most frequent in the human population, respectively D4.4R (64%), D4.7R (21%) and D4.2R (8%) [5]. The functional role of this polymorphism is not well characterized but several studies suggest association of the 7-repeat variant (D4.4R) with a predisposition to develop attention deficit hyperactivity disorder (ADHD) [6, 7]. In our previous study we have identified by yeast two-hybrid screening a BTB-Kelch protein, KLHL12, which specifically interacts with the polymorphic repeats of the D4R and functions as adaptor in the Cullin3-based E3 ligase complex promoting ubiquitination of the receptor [8].

Ubiquitination is a posttranslational modification involving a covalent attachment of a 76-amino acid polypeptide, ubiquitin, predominantly to lysine residues in the substrate protein. Ubiquitin can be attached to a single lysine residue resulting in monoubiquitination or to multiple lysines in the substrate protein resulting in multi-monoubiquitination. Ubiquitin itself contains seven lysine residues and all of them can serve as ubiquitin binding sites. Addition of an extra ubiquitin to the previous substrate-bound ubiquitin molecule leads to the formation of polyubiquitin chains of different configuration [9, 10].

Unconventional ubiquitination is the phenomenon suggested for the first time by Berejtschopf et al. in 1998 [11] and refers to ubiquitin binding to non-lysine residues. Until now, besides on lysine, ubiquitination on the N-terminus of the polypeptide backbone [11–13], on cysteine [14–16], on serine and threonine [17–19] residues has been described. Different types of ubiquitination can lead to different functional effects. The best characterized role of ubiquitination is to mark its target proteins for proteasomal degradation, which is mainly mediated by lysine 48-linked polyubiquitination, and lysosomal degradation. Additionally, ubiquitination can also be involved in many other processes, such as protein trafficking, DNA repair, inflammation and protein translation [9, 20]. The unconventional ubiquitination seems to have similar functions as typical lysine-mediated ubiquitination and works mainly as a signal for protein degradation [14, 17–19, 21].

Another interesting feature of conventional and unconventional ubiquitination is their redundancy. When the most preferred ubiquitin binding site is not available, ubiquitin can be attached to another lysine or non-lysine residue in the substrate protein [17–19, 21]. Finally, ubiquitination is also a reversible process in which ubiquitin can be removed from the substrate protein by deubiquitinating enzymes.

Ubiquitination also plays an important role in GPCR regulation. Two major types of GPCR ubiquitination can be identified. The first one, agonist-dependent ubiquitination of plasma membrane receptors, functions as a sorting signal for lysosomal degradation. This mechanism is important for signal termination of receptors after prolonged agonist stimulation or irreversibly activated receptors [22]. In this case, stimulated receptors are targeted to lysosomes via the highly conserved endosomal-sorting complex required for transport (ESCRT) pathway [22–28].

The second major type of GPCR ubiquitination is the agonist-independent or constitutive ubiquitination. Basally ubiquitinated receptors can be present both at the plasma membrane and in the endoplasmic reticulum (ER). Ubiquitination of GPCRs during synthesis in the ER functions mainly as a quality control system in which misfolded receptors are ubiquitinated and targeted for degradation via the ER-associated degradation (ERAD) pathway [29].
However, there are examples described of properly folded GPCRs e.g. Adenosine receptor A2A that are directed from the ER to the proteasome and after deubiquitination are transported to the cell surface [30].

We have investigated before the role of KLHL12-mediated ubiquitination in D4R degradation but no influence of KLHL12 on the expression level of D4R or on the half-life of the receptor were observed, allowing us to conclude that this specific ubiquitination does not lead to receptor degradation [31].

In this current work we investigate which specific residues in the sequence of the D4R are ubiquitinated and provide evidence for the ubiquitination on non-lysine residues. This poorly studied phenomenon has so far only once been suggested for a GPCR [32]. Moreover, we also show for the first time that the D4.7-R variant, which is associated with a predisposition to develop ADHD, is differentially ubiquitinated compared to the other common receptor polymorphic variants, namely D4.2R and D4.4R, suggesting the possibility of different downstream signaling effects.

Materials and Methods

Plasmids and antibodies

Plasmids encoding HA D4.2R, HA D4.4R, HA D4.7R, HA D4.6R, Flag Ubiquitin, Etag KLHL12, Cmyc Cullin3, were described before [8]. The D4.2R contains only four lysine (K) residues in its whole primary structure, namely at sites 229 (in the third intracellular loop before the polymorphic region), 304, 311 (both in the third intracellular loop after the polymorphism) and 381 (in the C-terminal tail). The HA D4.2R construct was used as a template for the application of single-nucleotide mutations (encoding arginine instead of lysine) using the QuickChange® Site-Directed Mutagenesis Kit from Stratagene (La Jolla, CA, USA).

The four individual K-R mutants were made separately, and are further denoted as D4.2R K229R, K304R, K311R, and K381R. The K304R mutant was used as a template for the construction of the quadruple mutant D4.2R K229,304,311,381R (further denoted as HA D4.2 4KRR).

HA β2AR was a kind gift from Prof. Dr. R. Lefkowitz (Duke University, Durham, NC). HA CXCR4 was purchased from pCDNA.org.

Primary antibodies used were mouse monoclonal anti-HA (clone16B12; Covance Research Products, cat. no. MMS-101-P-1000), rabbit polyclonal anti-Etag (Abcam, cat. no. ab3397-250), rabbit polyclonal anti-c-myc (Sigma, cat. no. C3956), rabbit polyclonal anti-HA (GeneTex, cat. no. GTX29110), rat monoclonal anti-HA (clone 3F10; Roche, cat. no. 11867423001), horseradish peroxidase (HRP)-conjugated mouse monoclonal anti-Flag M2 (Sigma, cat. no. A8592), rabbit polyclonal phospho-p44/42 MAPK (Cell Signaling, cat. no. 9101L) and mouse monoclonal p44/42 MAPK (clone L34F12; Cell Signaling, cat. no. 4696S).

Secondary antibodies used were goat anti-rabbit IRDye680 LT (cat. no. 926–68021), goat anti-mouse IRDye680 LT (cat. no. 926–68020), goat anti-mouse IRDye680 RD (cat. no. 926–68070), goat anti-rabbit IRDye800 (cat. no. 926–32211), goat anti-mouse IRDye800 (cat. no. 926–32210), goat anti-rat IRDye680 LT (cat. no. 926–68029) and were all purchased from LI-COR Biosciences.

Cell culture and transfection

HEK293T cells were cultured in Dulbecco’s modified Eagle’s medium (DMEM; Invitrogen), supplemented with 10% fetal calf serum, penicillin (100 U/ml), and streptomycin (100 µg/ml) in a controlled environment (37°C, 98% humidity, 5% CO2). HEK293T cells were transfected using the Polyethylenimine (PEI) method as described before [8]. A total amount of 10 µg of DNA was used for transfection of cells in one 10 cm dish.
HEK293S cells parental were a kind gift of Prof. Dr. Nico Callewaert [33]. Cells were stably transfected with calcium phosphate method with pHA D4.2R or pHA D4.2 4KRR and were grown in DMEM/F12 (Gibco, Invitrogen) supplemented with 10% fetal calf serum, penicillin (100 U/ml), streptomycin (100 μg/ml), and 0.5 mg/ml G418 (Geneticin, Gibco) in a controlled environment (37°C, 98% humidity, 5% CO2).

**Co-immunoprecipitation**

Forty eight hours after transfection cells were washed twice with cold phosphate-buffered saline (PBS), harvested and the cell pellet was frozen at −70°C for at least 1 h before lysis. Cell lysates were subjected to immunoblot analysis, or to immunoprecipitation (IP) followed by immunoblotting (IB), as described before [34].

**Ubiquitination assay—sequential double immunoprecipitation**

Ubiquitination assay was described before [35]. Briefly, cells were lysed in radioimmunoprecipitation buffer (RIPA) supplemented additionally with the inhibitor of deubiquitinating enzymes (10 mM), N-ethylmaleimide (NEM). For the first IP 2 μg of anti-HA (16B12) antibody was used and samples were incubated overnight with antibody and agarose beads. Next, three wash steps with the lysis buffer were performed and proteins bound to the beads (receptor and interacting partners) were eluted under denaturing conditions, and a quarter of the eluate was used to confirm the first IP. The rest of the eluates, containing denatured proteins, were diluted with lysis buffer and subjected to a second IP to remove receptor-interacting proteins from the first IP and specifically isolate the protein of interest. Finally, the eluates from the second IP were subjected to immunoblotting for the detection of ubiquitinated receptor.

**Chemical cleavage of ubiquitin with sodium hydroxide and dithiothreitol**

A modified protocol of sequential double immunoprecipitation was used to detect the influence of highly reducing conditions (dithiothreitol-DTT) or high pH (NaOH) on the ubiquitination status of the receptor. High pH treatment was performed by resuspending immunoprecipitates in 0.5% SDS, boiling for 3 min, cooling to room temperature, adding NaOH to a final concentration of 50 mM, and then incubating the samples at 32°C for 1 h. Mock-treated samples were incubated with PBS. Samples were then neutralized via the addition of 0.5 M Tris/HCl, pH 6.8. 5x Laemmli sample buffer (5% SDS; 50% glycerol; 0.2% bromophenol blue; 65 mM Tris/HCl, pH 6.8) without reducing agents was added, and samples were boiled for 3 min. Next, eluates were transferred to the new tubes, diluted with lysis buffer and a second round of IP was performed as described before.

Reducing treatments were performed by resuspending immunoprecipitates in 5x Laemmli sample buffer supplemented with 100 mM DTT followed by boiling for 3 min. Next, eluates were transferred to the new tubes, diluted with the lysis buffer and a second round of IP was performed.

All protein samples were separated via SDS-PAGE, transferred to the nitrocellulose membrane and subjected to immunodetection with primary and secondary antibodies.

**Detection of p44/42 mitogen-activated protein kinase (MAPK) phosphorylation by in cell western**

HEK293S cells stably transfected with HA D4.2R or HA D4.2 4KRR were grown on poly-D-lysine coated 96-well plates until 80% confluency. After overnight incubation in serum free medium, cells were treated with dopamine in concentration ranging from $10^{-8.5}$ to $10^{-4.5}$ M for 3 min at
37°C. Incubation was stopped by removing the culture medium, followed by addition of fixing solution (3.7% formaldehyde in PBS) for 20 min at room temperature (RT). Next, cells were permeabilized by washing 4 times for 5 min with Triton washing solution (0.1% Triton X-100 in PBS). Subsequently, cells were blocked with a blocking buffer (LI-COR Biosciences) for 90 min and finally, cells were incubated overnight at 4°C with two primary antibodies: rabbit phospho-p44/42 MAPK and mouse p44/42 MAPK diluted in the blocking buffer. The next day, the plate was washed 4 times for 5 min at RT with Tween washing solution (0.1% Tween-20 in PBS) and incubated with fluorescently labeled secondary antibodies: goat anti-rabbitIRDye800 and goat anti-mouse IRDye680 RD for 1 hour at RT. After final washing with Tween washing solution, fluorescent signal was detected with the Odyssey Infrared Imaging system. In the analysis, background values of the secondary antibodies are subtracted and the phospho-p44/42 MAPK signal is normalized against the total p44/42 MAPK signal.

**Immunofluorescence microscopy**

HEK293S cells stably expressing HA D4.2Ro or HA D4.2 4KRR were seeded in wells with coverslips. Twenty four hours after seeding, cells were fixed with fixing solution (3.7% formaldehyde in PBS) for 20 min at RT. After washing, cells were permeabilized with 0.1% Triton X-100 in blocking buffer (3% newborn calf serum, 1% bovine serum albumin in PBS) for 20 min at RT. Subsequently, cells were incubated for one hour with blocking buffer, followed by incubation for one hour with rabbit anti-HA diluted in blocking buffer (1:1000). Next, secondary antibody anti-rabbit Alexa Fluor 488 diluted in blocking buffer (1:500) was applied and incubation was performed in the dark for 1 hour. Cell nuclei were stained with DAPI (1 μg/ml). After each step, mentioned above, coverslips were washed three times with PBS. Mounting was performed with Mowiol and samples were analysed using the Axiocam 200 microscope (Zeiss, Thornwood, NY).

**Statistical analyses**

To determine significance, results were compared with control conditions by means of t test (for two groups) or by One-way analysis of variance (ANOVA) with Bonferroni post-test (for more than two groups). All statistical analyses were performed using Prism software (version 5; GraphPad Software). P < 0.05 at the 95% confidence level was considered significant.

**Results**

The dopamine D4 receptor is ubiquitinated on non-lysine residues

In previous studies we have shown that KLHL12, a BTB-Kelch protein, functions as an adaptor in the Cullin3-based E3 ubiquitin ligase complex and promotes ubiquitination of the D4R

In this study, we aim to characterize the ubiquitination pattern of the D4R into more detail.

The D4.2R contains only four lysine (K) residues in its whole primary structure, namely at sites 229 (in the IC3 before the polymorphic region), 304, 311 (both in the IC3 after the polymorphism) and 381 (in the C-terminal tail) that can potentially be ubiquitinated (Fig 1). In order to elucidate which lysine residue(s) could serve as acceptor site(s) for ubiquitin-conjugation, four receptor mutants, each with one of the four lysine residues mutated to arginine (R), were constructed using site-directed mutagenesis. Although the structure of an arginine residue resembles the structure of lysine, the arginine cannot serve as an acceptor site for ubiquitin-conjugation. Therefore, receptor ubiquitination is thought to be impaired upon mutation of the preferred ubiquitin-conjugation site. In parallel, we constructed a receptor mutant in which all four lysine residues were mutated (D4.2 4KRR) and this mutant thus should be completely
unable to undergo ubiquitination. First, the expression profile of these mutants and their ability to interact with KLHL12 was tested. Upon transient transfection of the HA-tagged receptor constructs and Etag KLHL12, lysates were tested for expression of all receptors and further subjected to immunoprecipitation with anti-HA. From these experiments, it can be concluded that all K-R receptor mutants are clearly expressed, and are able to interact with KLHL12 (S1 Fig).

Subsequently, ubiquitination of all individual mutant receptors was tested by ubiquitination assays in HEK293T cells (Fig 2). Surprisingly, comparable ubiquitination levels could be observed for all four single K-R mutant receptors (lanes 4, 6, 8, 10). KLHL12 promoted ubiquitination of the WT HA D₄R (lane 3) but also of all four single K-R mutant receptors (1 and 2 IP: lanes 5, 7, 9, 11). These observations could reflect the possibility that upon mutation of one lysine residue, another lysine residue serves as an alternative acceptor site for ubiquitin-conjugation. Therefore, these results suggest redundancy of the lysine residues in the D₄R to serve as attachment sites for basal or KLHL12-promoted ubiquitination.

In the next step, the membrane expression and functionality of the quadruple HA D₄ 4KR,R mutant was verified by examining its ability to phosphorylate MAP kinase in response to agonist stimulation (S2 Fig). In a second stage, the ubiquitination level of this mutant receptor was investigated. In these experiments, a very strong increase in ubiquitination of the mutant.
Fig 2. Ubiquitination of all four single K-R mutants of D₄R is increased by KLHL12. HEK293T cells were transiently transfected as indicated. 48 h post-transfection, cells were harvested and lysed. 5% of the lysate was used for IB to visualize HA-tagged WT D₄R and all single K-R mutants, (Flag Ub)n-proteins, and Etag KLHL12, respectively (bottom panels). The rest of the lysate was subjected to a sequential double IP with anti-HA (16B12). Specific purification of the HA D₄R WT and all K-R mutants was confirmed after the first (middle panels) and second IP (upper panels) upon IB with anti-HA (16B12, 1:2000), whereas receptor ubiquitination was revealed upon IB with anti-Flag-HRP (1:2000). * D₄R specific signal. ** Association of two heavy chains of mouse anti-HA antibody (each 50 kDa). The results shown are representative of three independent experiments.

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The dopamine D_{4}R receptor was again detected in the ubiquitination assay upon overexpression of Etag KLHL12 (Fig 3) which suggests that D_{4}R can be ubiquitinated on other residues than lysine.

Up to now, besides on lysine, ubiquitination on the N-terminus [11–13], on cysteine [14–16], on serine and threonine [17–19] was described. Because of its extracellular location, KLHL12-mediated ubiquitination of the N-terminus of D_{4}R is unlikely. Therefore, in further investigations we focused only on the possible ubiquitination of the residues which are located in the intracellular parts of the receptor. Fig 1 shows the estimated localization of all possible ubiquitination sites in the intracellular part of the D_{4}R.

**KLHL12 can promote ubiquitination on cysteine residues of D_{4}R**

As shown in Fig 1 there are two conserved cysteine residues in each polymorphic repeat in the third intracellular loop and an additional two are present at the C-terminus of D_{4}R. To investigate a possible attachment of ubiquitin to cysteine we decided to use a specific chemical approach, in which we rely on the fact that the thioester bond which is formed between ubiquitin and cysteine is susceptible to strong reducing conditions (for example high concentration of DTT). For this assay, HA-tagged WT D_{4}R or mutant D_{4}R 4KRR were immunoprecipitated from transiently transfected HEK293T cells. The resulting IP pellets containing ubiquitinated receptors were treated with 100 mM DTT, which has been shown to cleave Ub chains attached to cysteine residues [15, 16, 36]. Next, eluates were diluted with lysis buffer and a second IP was performed, as before, to prevent detection of a ubiquitination signal originating from D_{4}R interacting proteins instead of the receptor itself (Fig 4). Treatment with the strong reducing agent DTT after performing the ubiquitination assay resulted in a very strong decrease in ubiquitination signal of WT D_{4}R and of the lysine-deficient D_{4}R 4KRR mutant. When Etag KLHL12 was overexpressed, the detected ubiquitination signal was much stronger also in the samples treated with DTT (e.g. compare in Fig 4 lane 11 and lane 13 for WT D_{4}R and compare lane 5 and 7 for the mutant D_{4}R 4KRR), but DTT treatment again decreased the ubiquitination signal very efficiently. These results indicate that KLHL12 can promote cysteine-linked D_{4}R ubiquitination. The ubiquitination signal is still visible after DTT treatment in case of D_{4}R 4KRR when Etag KLHL12 is overexpressed and therefore, these results suggest that there are also other residues, besides cysteine and lysine, that can be ubiquitinated in the D_{4}R.

**KLHL12 promotes ubiquitination on serine and/or threonine residues of D_{4}R**

When ubiquitin is attached to serine or threonine an oxyester bond is formed. This kind of bond can be destroyed by treatment with a strong base (50–100 mM NaOH) [36, 37]. To examine the possibility of ubiquitination of the D_{4}R on serine and threonine residues, IP eluates were treated with 50 mM NaOH for 1h at 32°C as described before [37] (Fig 5A). In the same experiment we also examined the influence of highly reducing conditions on the same samples to remove ubiquitin from cysteine residues (as described above). This facilitated the comparison of the results of the two different chemical treatments, indicative for serine/threonine and cysteine ubiquitination, respectively. After the first IP samples were diluted with lysis buffer and a second IP was performed similar as before.

As shown in Fig 5A treatment with NaOH led to a significant decrease in basal ubiquitination as well as a strong decrease in KLHL12-enhanced ubiquitination signal in case of both WT D_{4}R and D_{4}R 4KRR mutant. In this experiment the influence of DTT can also be clearly observed. Independent ubiquitination assays upon treatment with NaOH and DTT were performed at least four times for each of the treatments and we consistently observed a clear decrease in ubiquitination signal. However, often the receptor level after the second IP was not
equal in all of the samples and therefore, we decided to quantify the detected ubiquitination signal and normalize it against the receptor levels to identify trends in obtained data (Fig 5B–5E). The results presented in Fig 5A suggest that NaOH is more potent than DTT in removing ubiquitin moieties from D4R, although the performed quantification from two to seven independent experiments does not support this observation. However, and most importantly, all these results together suggest that ubiquitination of cysteine, serine and/or threonine residues in the D4R is possible. Moreover, from this quantification we can conclude that the decrease in ubiquitination signal caused by the chemical treatments is bigger in case of KLHL12-enhanced ubiquitination especially for the D4.2 4KRR (compare bars representing percentage of maximal ubiquitination signal detected upon treatment in Fig 5B and 5C for WT D4.2R and Fig 5D and 5E for D4.2 4KRR). Another interesting observation that can be made based on experiments involving chemical treatments is the overall lower ubiquitination level of D4.2 4KRR mutant compared to the WT D4.2R (Fig 5F). This suggests that lysines which are present in WT receptor can also be ubiquitinated and represent a fraction of the total ubiquitination signal, but it is not possible to conclude whether they are the preferred ubiquitin-binding sites in the KLHL12-promoted ubiquitination. As suggested by Wang et al. [38] before making a conclusion about non-lysine ubiquitination based on chemical treatment experiments one should be aware that too long alkaline treatment can affect also amide bonds formed between lysine and ubiquitin. Therefore, an additional control experiment was performed in which the influence of alkaline treatment on known lysine-linked ubiquitination samples was tested. We have examined the effect of NaOH treatment on ubiquitination of two other GPCRs for which

Fig 3. Lysine lacking D4.2 4KRR mutant is ubiquitinated and KLHL12 enhances its ubiquitination. HEK293T cells were transiently transfected as indicated. 48 h post-transfection, cells were harvested and lysed. 5% of the lysate was used for IB to visualize HA D4R, (Flag Ub)n-proteins, and Etag KLHL12, respectively (left panels). The rest of the lysates were subjected to a sequential double IP with anti-HA (16B12). Specific purification of the receptor after the second IP was confirmed upon IB with rat anti-HA (1:2000), whereas receptor ubiquitination was revealed upon IB with anti-Flag-HRP (right panels, 1:2000). The results shown are representative of four independent experiments.

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ubiquitination on lysine residues was described, namely the chemokine CXCR4 receptor [26, 39] and the β2-adrenoceptor (β2AR) [28]. S3 Fig clearly demonstrates that NaOH strongly affects basal and KLHL12-enhanced ubiquitination of D4R but seems to have no effect on ubiquitination of the other two receptors. This indicates that the NaOH treatment conditions used in our experiments are not strong enough to affect typical isopeptide bonds which are formed between ubiquitin and lysine residues in substrate proteins.

KLHL12 promotes differential ubiquitination of the D4R polymorphic variants

As mentioned in the introduction D4R has an important polymorphism in the third intracellular loop where a 16-amino acid sequence is repeated from 2 to 11 times. The most common variants in the human population are D4.4R (64%), D4.7R (21%) and D4.2R (8%), respectively [4]. Not much is known about functional differences between different polymorphic variants but several studies suggest an association between the D4.4R allele and ADHD [6].

Previously, we have demonstrated that KLHL12 binds specifically to the polymorphic region of D4R and promotes ubiquitination of the receptor. For these reasons, we were interested to explore whether KLHL12 differentially binds to the most frequent polymorphic variants and to study the effect of KLHL12 on the ubiquitination of these receptor variants.

First, we have examined using co-immunoprecipitation if KLHL12 binds to all common D4R polymorphic variants, namely D4.2R, D4.4R and D4.7R. As a negative control we used an
Fig 5. Treatment with NaOH decreases ubiquitination of D₄R WT and D₄R 4KR mutant. A) HEK293T cells were transiently transfected as indicated. 48 h post-transfection, cells were harvested and lysed. 5% of the lysates were used for IB to visualize (Flag Ub)ₙ-proteins, HA D₄R, and Etag KLHL12, respectively (left panels). The rest of the lysates were subjected to IP with anti-HA (16B12). After IP proteins were eluted in 0.5% SDS at 95°C. Next, lysates were incubated with or without addition of 50 mM NaOH for 1 h at 32°C, mock treated samples were incubated with PBS. Additionally, samples were treated with a highly reducing agent (5 min, 95°C, 100 mM DTT). After dilution with RIPA lysis buffer a second round of IP was performed with anti-HA antibody.
an artificial receptor variant from which the polymorphic region was removed (D₄.₀R). Fig 6 clearly shows that KLHL12 binds to all common polymorphic variants D₄.₂R, D₄.₄R and D₄.₇R and as expected no interaction with D₄.₀R was detected.

Next, ubiquitination assays were performed in HEK293T cells overexpressing different HA-tagged D₄R polymorphic variants with or without co-expression of Etag KLHL12. A strong increase in ubiquitination signal was observed in case of D₄.₂R and D₄.₄R variants when Etag KLHL12 was overexpressed. Surprisingly, we could hardly detect an increase in ubiquitination of the D₄.₇R variant (Fig 7).

The ubiquitination process requires integrated action of many proteins at the same time and therefore, we have also examined if the D₄.₇R variant is still capable to form a complex with E3 ligase, Cullin3. The results from the co-immunoprecipitation studies indicate that D₄.₇R is able to form a complex with Cullin3 (Fig 8), although we cannot rule out the possibility that this interaction could be mediated by adaptor proteins other than KLHL12.

Discussion

Previously, we have shown that KLHL12 interacts with the polymorphic repeats of the D₄R and enhances its ubiquitination [8] but this has no influence on receptor degradation [31]. In the present study we investigated which specific residues of the D₄R are ubiquitinated by the KLHL12-Cullin3 E3 ligase complex. In the whole sequence of D₄R there are only four lysine residues and all of them can potentially serve as ubiquitin binding sites as they are localized in the intracellular domains of the receptor. Using a site-directed mutagenesis approach each lysine was separately mutated to arginine. Next, sequential double IP experiments demonstrated that all four mutants are ubiquitinated and overexpression of KLHL12 increases their ubiquitination level (Fig 2). This suggests that ubiquitination occurs on multiple lysines or that the ubiquitination system is flexible and when the most preferred ubiquitin binding site is unavailable it can ubiquitinate another available lysine. Next, we created the quadruple mutant HA D₄.₂ R₄KR in which all four lysines were mutated to arginines. This mutant should be unable to undergo lysine-mediated ubiquitination. Surprisingly, we could still detect basal ubiquitination of this mutated receptor and upon co-expression of KLHL12 an increase in its ubiquitination was visualized (Fig 3). This finding encouraged us to search for other possible ubiquitination sites in the D₄R. Ubiquitination on non-lysine residues is still a poorly studied phenomenon and until now ubiquitination on the N-terminus [11–13, 40], cysteine [14–16], serine and threonine [17–19] residues was described, but it has only been suggested once for a GPCR unfortunately without characterizing the precise amino acid residue [32]. In our study we investigated the possibility of ubiquitin binding to cysteine, serine and threonine residues within the intracellular domains of the D₄R. One of the most common approaches to verify ubiquitination on a non-lysine residue is a chemical cleavage of ubiquitin [38]. Ubiquitin attached to a lysine residue forms an isopeptide bond which is very stable, whereas a thioester bond connecting ubiquitin with cysteine can be destroyed during treatment with a highly specific purification of the receptor after the second IP was confirmed upon IB with rat anti-HA (1:2000), whereas receptor ubiquitination was revealed upon IB with anti-Flag-HRP (right panels, 1:2000). N: samples treated with 50 mM NaOH, D: samples treated with 100 mM DTT. Experiment with two different chemical treatments was performed three times. Additionally, two more experiments were performed with NaOH treatment alone. B-E) Quantification of the ubiquitination signal detected in four to seven independent experiments was performed using Fiji software. Values were normalized to the receptor level in the corresponding lanes. Graph bars show percentage of maximal ubiquitination signal (mean ± SEM) where maximum is equal to 100% and represents the signal detected in samples which did not undergo any chemical treatment (NT- non-treated) in each independent experiment. Compared with NT: *** P < 0.001; ** P < 0.01; * P < 0.05. F) Quantification of the ubiquitination signal detected in five to seven independent experiments was performed using Fiji software. Values were normalized to the receptor level in the corresponding lanes. Graph bars show percentage of ubiquitination signal detected for WT D₄.₈R (mean ± SEM). Compared with the ubiquitination signal detected for WT D₄.₈R without or with overexpression of KLHL12, respectively: *** P < 0.001; ** P < 0.01. doi:10.1371/journal.pone.0145654.g005
reducing agent e.g. a high concentration of DTT [15, 16, 36]. Our results clearly indicate that upon treatment with a very high concentration of DTT the ubiquitination signal detected after the second IP is significantly reduced for both WT D4.2R and D4.2 4KRR, suggesting ubiquitination on cysteine residues (Fig 4).

Next, we examined the possibility of ubiquitination on serine and/or threonine residues. When ubiquitin is attached to a serine or threonine residue an oxyester bond is formed which is susceptible for cleavage in a strong alkaline environment [36, 37]. We have performed a ubiquitination assay in which, after the first IP, the samples were treated with NaOH and after the second IP a strong decrease in ubiquitination signal was observed (Fig 5A). These results thus suggest that D4R ubiquitination on cysteine, serine and/or threonine residues can be promoted by KLHL12. It is, however, not possible to say which residues are the most preferred for KLHL12-induced ubiquitination as quantification of the results from multiple independent experiments indicates a comparable decrease in ubiquitination levels for both types of treatments (Fig 5B–5E). The decrease is more explicit when Etag KLHL12 is co-expressed which suggests that during KLHL12-promoted ubiquitination more non-lysine residues in the D4R are ubiquitinated compared to the basally ubiquitinated receptor. This decrease is even more striking in case of the D4.2 4KRR mutant in which no lysine residues are available. This indicates that in the WT receptor KLHL12 promotes ubiquitination on lysine and non-lysine residues and in the D4.2 4KRR mutant, in which lysines are not available, only non-lysine residues are ubiquitinated. The fact that in all experiments we detect an overall lower ubiquitination signal from D4.2 4KRR (Fig 5F) compared to WT D4.2R further supports the hypothesis about the

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**Fig 6.** All common D4R polymorphic variants interact with KLHL12 while D4.0R, lacking repeats in the third intracellular loop, does not interact. HEK293T cells were transiently transfected as indicated. 48 h post-transfection, cells were harvested and lysed. Part of the lysate was used for IB to verify expression of HA-tagged D4R variants and Etag KLHL12 (left panels). The rest of the lysate was subjected to IP with anti-HA (16B12). Specific purification of the receptor after IP was confirmed upon IB with rat anti-HA (1:2000). Interaction of Etag KLHL12 with the D4R polymorphic variants was verified by IB with anti-Etag (1:2000). Data shown are representative of three independent experiments. Expected molecular weights of different D4R variants: D4.0R (44–50 kDa), D4.2R (47–53kDa), D4.4R (51–57 kDa), D4.7R (56–62 kDa).

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Fig 7. KLHL12 strongly enhances ubiquitination of D₄₂R and D₄₇R but hardly of D₄₇R. A) HEK293T cells transiently transfected as indicated were lysed 48 h post-transfection. 5% of the lysate was used for IB to verify expression of HA-tagged D₄R polymorphic variants, (Flag Ub)n-proteins and Etag KLHL12 (left panels). The rest of the lysate was used for double sequential IP with anti-HA (16B12). Specific purification of the receptors after the second IP was confirmed upon IB with mouse anti-HA (16B12, 1:2000), whereas receptor ubiquitination was revealed upon IB with anti-Flag-HRP (right panels, 1:2000). * Association of two heavy chains of mouse anti-HA antibody (each 50 kDa). Data shown are representative of similar results from five independent
involvement of lysine residues in KLHL12-mediated ubiquitination of D₄R. To exclude the possibility that the alkaline treatment affects amide bonds which are normally formed between ubiquitin and lysine residues we have also checked the influence of NaOH on ubiquitination of other GPCRs for which lysines were described as ubiquitin binding sites. We have used the CXCR4 receptor for which three lysines in the C-terminal tail were shown to be critical for receptor ubiquitination [26]. As a second receptor we used the β₂AR for which lysines in the third intracellular loop and in the C-terminal tail were presented to be important for ubiquitination and subsequent lysosomal degradation [28]. In this control experiment (S3 Fig), we have seen again a very strong effect of NaOH treatment on basal and KLHL12-induced ubiquitination of D₄R and no effect on CXCR4 and β₂AR ubiquitination. This result allows us to conclude that the conditions used for alkaline treatment are not affecting amide bonds which are formed during the typical lysine-linked ubiquitination process.

Most papers that study ubiquitination on non-lysine residues identified several different residues as possible ubiquitination sites. Some of them are most preferred by the ubiquitination machinery but when those are not available ubiquitin can bind to other amino acids. Such situation was described for example for NS-1 nonsecreted immunoglobulin light chain (NS-1 LC) which is predominantly ubiquitinated on serine/threonine residues but when these amino acids were mutated then lysine residues could serve as ubiquitin-binding sites. Mutation of all three types of amino acids (serines, threonines and lysines) was required to efficiently reduce ubiquitination and

**B) Quantification of the ubiquitination signal detected in three to five independent experiments was performed using Fiji software. Values were normalized to the receptor level in the corresponding lanes. Graph bars show ubiquitination signal (mean ± SEM, fold basal) where basal ubiquitination represents the signal detected in samples without overexpression of KLHL12, for the specific D₄R variant, in each independent experiment. Compared with the D₄X R without KLHL12 overexpression: ***, P < 0.01; *, P < 0.05.**

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Fig 8. D₄R forms a complex with Cullin3. HEK293T cells were transiently transfected as indicated. 48 h post-transfection, cells were harvested and lysed. 5% of the lysate was used for IB to verify expression of HA-tagged receptors, Cmyc Cullin3 and Etag KLHL12 (left panels). The rest of the lysate was subjected to IP with anti-HA (16B12). Specific purification of the receptor after IP was confirmed upon IB with rat anti-HA (1:2000). Interaction of Etag KLHL12 with D₄R was verified by IB with anti-Etag (1:2900) and the presence of Cmyc Cullin3 was visualized with anti-Cmyc (right panels, 1:1000).

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stabilize the NS-1 LC [37]. Another example represents the T-cell antigen receptor α-chain (TCRα) for which mutation of two conserved serine residues to alanines in the cytosolic tail inhibited significantly receptor ubiquitination. However, when serine residues were replaced with lysine, cysteine or threonine residues the protein was still ubiquitinated [17]. Our findings seem to be in agreement with this previously described phenomenon showing that the ubiquitination machinery is a very flexible system and can easily change from one to another ubiquitination site if necessary.

To obtain a direct proof of ubiquitination on a non-lysine residue we decided to perform mass spectrometry (MS) analysis. Detection of ubiquitination by MS is possible due to the diglycine remnant that remains attached to the ubiquitinated residue after tryptic digestion [41]. We have performed MS/MS analysis using a Fourier transform mass spectrometer but no ubiquitination was detected. Unfortunately, during tryptic digestion D₄R is cut to single amino acid or very large fragments making it very hard to analyze. The computational prediction of cleavage of D₄R with other enzymes, commonly used to prepare samples for MS analysis, did not suggest any good alternative to trypsine that could resolve this problem. According to our knowledge non-conventional ubiquitination was never proved with MS analysis probably due to the labile nature of thio-/oxy-ester bonds which are formed between ubiquitin and non-lysine residues. So far, only one study described the use of a novel peptide-based SILAC method to identify non-conventional ubiquitination of T-cell receptor α [42]. The authors could identify modified peptide which did not contain lysine in its sequence, however, they were unable to pinpoint the specific residue that undergoes ubiquitination.

Therefore, it is not very surprising that the detection of ubiquitination of D₄R with MS/MS was not successful.

However, and most importantly, all the obtained data allow us to conclude that KLHL12 can promote ubiquitination of D₄R on non-lysine residues and that ubiquitination on cysteine, serine and/or threonine is possible. The functional role of this non-lysine ubiquitination is hard to study because it would require the creation of a receptor with all possible ubiquitination sites mutated. Mutation of all intracellular cysteines, serines and threonines increases the chances of disrupted protein folding and it can also interfere with other posttranslational modifications. GPCRs are tightly regulated by phosphorylation which mainly occurs on serine and threonine residues and also by palmitoylation on cysteine residues (conserved cysteines located on the C-terminus of many GPCR) and also D₄R shows these posttranslational modifications. Additionally, this can also inhibit binding of some interacting proteins, e.g. β-arrestins for which often phosphorylation of the receptor is required and which also play a critical role in the regulation of GPCR signaling. Therefore, mutation of all of these residues increases the possibility of creating of a mutant with completely different signaling properties compared to the wild type receptor. In most of the examples of non-lysine ubiquitination, which were described up to now, this modification seems to serve as a signal for protein degradation [14, 17–19, 21]. Only in the case of Pex5p receptor (cytosolic receptor for peroxisome matrix proteins) ubiquitination of conserved cysteine¹⁰ was shown to be involved in regulation of the recycling of the receptor form the peroxisomes to cytosol [15]. To the best of our knowledge, ubiquitination of the non-lysine residue was suggested only once for a GPCR. The group of Shenoy has shown that β₂AR is probably ubiquitinated on a non-lysine residue in response to the treatment with an antagonist, carvedilol, as they could still detect ubiquitination of the β₂AR in which lysines were mutated. Interestingly, the same mutant was not ubiquitinated upon agonist, isoprotanol, treatment. Carvedilol-promoted ubiquitination is mediated by a completely different E3 ubiquitin ligase which is not involved in the, already well defined, lysine-mediated ubiquitination of β₂AR. However, both types of ubiquitination lead to endocytosis and lysosomal sorting of the receptor [32].
In the second part of our study we investigated the ubiquitination status of the three most common D₄R polymorphic variants. First, the interaction of KLHL12 with these variants was examined. After IP of HA-tagged receptors co-precipitation of Etag KLHL12 with D₄.2R, D₄.4R and D₄.7R was detected (Fig 6). As a negative control we used D₄.0R which is an artificial receptor variant lacking the polymorphic repeats. Next, a double sequential IP was performed to investigate the ubiquitination status of the different polymorphic variants. Ubiquitination of D₄.2R and D₄.4R was clearly increased when Etag KLHL12 was co-expressed, but surprisingly we hardly observed increased D₄.7R ubiquitination (Fig 7A and 7B), although this receptor variant is still able to form a complex with KLHL12 and Cullin3 (Fig 8). The differential ubiquitination pattern of the D₄.7R, compared to the other D₄R variants, is highly interesting as until now there is hardly evidence for pharmacological differences between this variant and the other receptor variants [43–45], although behavioral research showed that the D₄.7R is linked with a predisposition to develop ADHD [6], and also its association with increased sexual behavior, alcohol and cigarette craving was suggested. The functional role of differential KLHL12-mediated ubiquitination of the D₄R is still under investigation. This finding opens a lot of new questions for future research on the possible role of ubiquitination in receptor signaling and suggests that ubiquitination can regulate GPCRs in a much more complicated way than previously expected.

Supporting Information

S1 Fig. All K-R D₄R mutants interact with KLHL12. HEK293T cells were transiently transfected as indicated. 48 h post-transfection, cells were harvested and lysed. 5% of the lysate was used for IB to visualize HA-tagged D₄ receptors and Etag KLHL12. The rest of the lysate was subjected to IP with anti-HA (16B12). Specific purification of the receptors after IP was confirmed upon IB with anti-HA (1:2000), whereas co-precipitation of Etag KLHL12 was verified by IB with anti-Etag (1:2000). * Association of two heavy chains of mouse anti-HA antibody (each 50 kDa).

S2 Fig. D₄.2 4KRR is expressed on the membrane and its stimulation leads to the phosphorylation of p44/42 MAPK. A) HEK293S cells stably expressing HA D₄.2R and HA D₄.2 4KRR were treated with dopamine in concentration ranging from 10⁻⁸.⁵ to 10⁻⁴.⁵ M. An in cell western assay was performed and phosphorylated p44/42 MAP kinase was detected with rabbit anti-phospho p44/42 MAPK (1:1000). Values were normalized against the signal detected with mouse anti-p44/42 MAPK (1:800). B) HEK293S cells stably expressing HA D₄.2R and HA D₄.2 4KRR were seeded in wells with coverslips. Cells were stained using primary antibody rabbit anti-HA (1:1000) and secondary antibody anti-rabbit Alexa Fluor 488 (1:500). Cell nuclei were colored with DAPI.

S3 Fig. Treatment with NaOH decreases ubiquitination of D₄.2R but has no influence on ubiquitination of CXCR4 receptor or β₂-adrenoceptor (β₂AR). HEK293T cells were transiently transfected as indicated. 48 h post-transfection, cells were harvested and lysed. 5% of the lysate was used for IB to visualize HA-tagged receptors, (Flag Ub)n-proteins, and Etag KLHL12, respectively (left panels). The rest of the lysate was subjected to IP with anti-HA (16B12). After IP proteins were eluted in 0.5% SDS at 95°C. Next, lysates were incubated with or without addition of 50 mM NaOH for 1 h at 32°C, mock treated samples were incubated with PBS. After dilution with RIPA lysis buffer a second round of IP was performed with anti-HA antibody. Specific purification of the receptors after the second IP was confirmed upon IB.
with rabbit anti-HA (1:2000), whereas receptor ubiquitination was revealed upon IB with anti-Flag-HRP (right panel, 1:2000). N: samples treated with 50 mM NaOH.

(AIF)

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Author Contributions
Conceived and designed the experiments: KS PR KVC. Performed the experiments: KS PR BL. Analyzed the data: KS PR KVC. Wrote the paper: KS PR KVC.

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