Gαolf Mutation allows Parsing the Role of cAMP- and Extracellular Signal-Regulated Kinase-dependent Signaling in L-DOPA-Induced Dyskinesia

• **Abbreviated title** (45<50): PKA and ERK signaling in L-DOPA-induced dyskinesia

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ABSTRACT

Although L-DOPA remains the reference treatment of Parkinson’s disease, its long-term beneficial effects are hindered by L-DOPA-induced dyskinesia (LID). In the dopamine-denervated striatum, L-DOPA activates dopamine D1 receptor (D1R) signaling, including cAMP-dependent protein kinase (PKA) and extracellular signal-regulated kinase (ERK), two responses associated with LID. However, the cause of PKA and ERK activation, their respective contribution to LID, and their relationship are not known. In striatal neurons, D1R activates adenylyl-cyclase through Gαolf, a protein up-regulated following lesion of dopamine neurons in rats and in patients. We report here that increased Gαolf levels in hemiparkinsonian mice are correlated with LID following chronic L-DOPA treatment. To determine the role of this upregulation, we performed unilateral lesion in mice lacking one allele of the Gnαl gene coding for Gαolf (Gnαl +/-). Despite an increase in the lesioned striatum, Gαolf levels remained below those of unlesioned wild type mice. In Gnαl +/- mice, the lesion-induced L-DOPA stimulation of cAMP/PKA-mediated phosphorylation of GluA1 Ser845 and DARPP-32 Thr34 was dramatically reduced, whereas ERK activation was preserved. LID occurrence was similar in Gnαl +/- and +/- mice after a 10-day L-DOPA (20 mg/kg) treatment. Thus, in lesioned animals Gαolf up-regulation is critical for the activation by L-DOPA of D1R-stimulated cAMP/PKA, but not ERK signaling. Although the cAMP/PKA pathway appears required for LID development, our results indicate that its activation is unlikely to be the main source of LID. In contrast, the persistence of L-DOPA-induced ERK activation in Gnαl +/- mice supports its causal role in LID development.

INTRODUCTION

Parkinson’s disease (PD) results from neurodegeneration of nigrostriatal dopamine (DA) neurons. Despite innovative therapies, the DA precursor L-DOPA remains the drug of reference. However, after an initial period of full efficacy, this treatment is complicated by L-DOPA-induced dyskinesia (LID) with an incidence of 30-50% after 5 years and 60% after 10 years (Rascol, 2000; Van Gerpen et al., 2006). Better understanding of the neuronal mechanisms underlying the development of LID is therefore important to identify therapeutic strategies (Cenci, 2007; Jenner, 2008).

DA D1 receptor (D1R) agonists are more efficient than D2R agonists to induce dyskinesia in animal models and PD patients (Calon et al., 1999; Rascol et al., 2001; Rascol et al., 2006; Carta et al., 2008). D1Rs are necessary for LID development (Westin et al., 2007; Darmopil et al., 2009). D1R signaling through cAMP-dependent protein kinase (PKA) and 32-kDa DA- and cAMP-regulated phosphoprotein (DARPP-32) is activated by L-DOPA in 6-OHDA-lesioned rodents (Santini et al., 2007; Lebel et al., 2010) or MPTP-treated monkeys (Santini et al., 2010). Strong D1R-dependent activation of extracellular signal-regulated kinase (ERK) (Gerfen et al., 2002; Pavon et al., 2006; Santini et al., 2007; Westin et al., 2007; Rylander et al., 2009; Santini et al., 2009b) and mTOR signaling (Santini et al., 2009a) is also observed in striatonigral medium-sized spiny neurons (MSNs). Pharmacological or genetic blockade of these pathways supports their necessary role in LID (Santini et al., 2007; Santini et al., 2009a; Bateup et al., 2010; Fasano et al., 2010; Lebel et al., 2010). However, the cause
of their activation in DA-deprived striatum and their relative contribution in LID development are unknown.

Levels of D1R (Shinotoh et al., 1993; Turjanski et al., 1997; Hurley et al., 2001) and major mediators of D1R-signaling, including PKA and DARPP-32 (Girault et al., 1989; Nishino et al., 1993) are unchanged in PD and its animal models. In MSNs D1Rs and adenosine A2A receptors (A2AR) are coupled to adenylyl-cyclase through a heterotrimeric G-protein comprising Goolf, Gβ2, and Gγ7 subunits (Drinnan et al., 1991; Herve et al., 1993; Kull et al., 2000; Zhuang et al., 2000; Corvol et al., 2001; Schwindinger et al., 2003; Schwindinger et al., 2010). Levels of Goolf increase in DA-denervated striatum of rats and PD patients (Herve et al., 1993; Marcotte et al., 1994; Penit-Soria et al., 1997; Corvol et al., 2004; Rangel-Barajas et al., 2010). Since Goolf levels are a limiting factor for receptor signaling (Corvol et al., 2001; Corvol et al., 2007), Goolf up-regulation is a plausible but untested mechanism for the D1R hypersensitivity in LID-developing animals.

The objective of this study was to examine the association of Goolf with LID and evaluate the respective contribution of cAMP- and ERK-mediated pathways in LID. We used a mouse model of hemiparkinsonism treated with L-DOPA (Lundblad et al., 2004) and heterozygous mutant mice lacking one allele of the Goolf gene (Gnal) (Belluscio et al., 1998; Corvol et al., 2007). This approach allowed us to parse the role of these various signaling components in the apparition of LID.

MATERIAL AND METHODS

Animals
We used 8-week-old C57Bl/6J male mice. Mice heterozygous for a null mutation of the gene encoding Goolf (Gnal) kindly provided by L. Belluscio (Belluscio et al., 1998) were backcrossed for eight generations with C57Bl/6J wild type mice and mated with C57Bl/6J mice for producing heterozygous Gnal mutant mice and wild type littermates. Animals less than 30-week old were used for the experiments. The mice were maintained in a 12 h light–dark cycle, in stable conditions of temperature (22°C), with free access to food and water. All the experiments were in accordance with the guidelines of the French Agriculture and Forestry Ministry for handling animals (decree 87-848) and were conducted under the approval of the “Direction Départementale de la Protection des Populations de Paris” (authorization number C-75-828, license B75-05-22).

6-OHDA lesions and post-operative care
Mice were anesthetized with a mixture of xylazine (10 mg/mL) and pentobarbital (25 mg/mL) (Centravet Plancoet, La Milière, France), and mounted in a digitalized stereotactic frame (Stoelting Europe, Dublin, Ireland) equipped with a mouse adaptor. 6-OHDA-HCl (3.0 mg/mL, Sigma-Aldrich, Lyon, France) was dissolved in a solution containing 0.2 g/L ascorbic acid and 9 g/L NaCl. Mice received two unilateral injections (2 x 2 µL) of 6-OHDA into the right striatum at the following coordinates according to the mouse brain atlas of Paxinos and Franklin (Paxinos and Franklin, 2001): AP= +1.2 mm, L= +2.1 mm, DV= -3.2 mm; and AP= +0.6 mm, L= +2.4 mm, DV= -3.2 mm. Each injection was performed with a 36-gauge, 50-mm long stainless steel cannula connected to a syringe pump (MTI Corporation, UC Berkeley, USA) by a polyethylene catheter, at a slow rate of 0.25 µL/min to minimize tissue damage. After the injection, the cannula was left in place for an additional 4 min before being slowly retracted. Sham mice were injected with vehicle only (ascorbic acid in saline).
Post-operative care was needed to minimize the mortality rate. Mice were let on a warm plate during about 24 h after surgery in order to avoid hypothermia. To reduce suffering, mice received subcutaneous (s.c.) injections of a non-steroidal anti-inflammatory drug (flunixin meglumine, 4 mg/kg, Sigma-Aldrich), just after the surgery and twice daily during 2 days post-surgery. All the lesioned mice were examined daily and the weakest animals received injections of sucrose 5% (10 mL/kg, s.c.) and intraperitoneal (i.p.) saline (10 mL/kg) to avoid dehydration. Concentrated milk was provided to all the lesioned animals during 3 days after the operation. Mice were allowed to recover for 4 weeks before behavioral evaluation and drug treatment. Lesions were assessed at the end of experiments by determining the striatal levels of tyrosine hydroxylase (TH) using immunohistochemistry or immunoblotting (see below). Only animals with a TH-depleted area above 80% of the striatum area and/or TH level reduction by more than 80% in the lesioned striatal area as compared to the control side were included in the analyses (i.e. about 85% of surviving lesioned mice).

**L-DOPA and benserazide treatments**

L-DOPA and the peripheral DOPA decarboxylase inhibitor benserazide hydrochloride (Sigma-Aldrich) were dissolved together in physiological saline solution (NaCl, 9 g/L). L-DOPA and benserazide hydrochloride were injected i.p. at a dose of 20 and 12 mg/kg, respectively in a volume of 10 mL/kg of body weight. For acute treatments mice were euthanased 30 min after the first injection of L-DOPA and benserazide. The chronic treatment consisted in a daily single injection of L-DOPA and benserazide during 10 days. Mice were sacrificed on day 11, 30 min after the last injection.

**Behavioral tests**

**Cylinder test.** On day 1 of chronic treatment, the anti-parkinsonian effects of L-DOPA on sensorimotor function were evaluated in the cylinder test (Lundblad et al., 2002). Each lesioned mouse was tested before the beginning of L-DOPA therapy and 1 h after the first injection of L-DOPA plus benserazide. Mice were placed one by one in glass cylinders (10 cm diameter and 15 cm height) and videorecorded for 5 min without previous habituation to the glass cylinder. Once introduced into the cylinder, the mice showed an exploratory behavior reflected by rearing and forepaw contacts on the wall. The number of contacts with the right or left forepaw was counted in blind conditions regarding mouse genotype. To eliminate accidental touches without physiological meaning, we counted only contacts in which the animal supported its body weight on the paw with extended digits. The use of the impaired (left) forepaw was expressed as a percentage of the total number of contacts on the wall.

**Abnormal Involuntary Movements (AIMs).** 6-OHDA-lesioned mice were chronically treated with L-DOPA for 10 days. AIMs were assessed on day 5 (two experiments) and day 10 (all experiments) in blind conditions regarding mouse genotype, using a previously validated scale for scoring LID in mouse (Lundblad et al., 2004; Lundblad et al., 2005). Immediately after L-DOPA administration, mice were placed in separate cages and abnormal movements were assessed for 1 min (monitoring period) every 20 min, over a period of 140 min. Abnormal movements, clearly distinct from natural stereotyped behaviors (i.e., grooming, sniffing, rearing, and gnawing), were classified into four different subtypes: locomotive (tight contralateral turns), axial (contralateral dystonic posture of the neck and upper body), limb (jerky and fluttering movements of the limb contralateral to the side of the lesion), and orofacial (vacuous jaw movements and tongue protrusions) AIMs. Each subtype was scored on a severity scale from 0 to 4: 0, absent; 1, occasional; 2, frequent; 3, continuous; 4,
continuous and not interruptible by external stimuli. The total AIMs score corresponded to the sum of individual scores for each AIM subtype. A composite score was obtained by the addition of scores for axial, limb and orofacial AIMs (ALO score). The ALO score is considered to more closely reflect the human dyskinetic behavior than the locomotive AIMs score (LOC score) (Lundblad et al., 2002). To identify the extreme behaviors highly and weakly dyskinetic mice were separated by a quartile analysis based on total AIMs scores.

**Tissue preparation and immunofluorescence**

Thirty minutes after the last injection of L-DOPA, mice were rapidly anesthetized with pentobarbital (500 mg/kg, i.p., Sanofi-Aventis) and perfused transcardially with 40 g/L paraformaldehyde in phosphate buffered saline (PBS, 0.1 M sodium phosphate, 0.14 M NaCl), pH 7.4. Brains were postfixed overnight in the same solution at 4°C. The following day, 30-µm-thick sections were cut with a Vibratome (Leica, Nussloch, Germany) and stored at -20°C in a buffered solution containing 30% volume/volume (v/v) ethylene glycol, 30% (v/v) glycerol, and 0.1 M sodium phosphate pH 7.4, until they were processed for immunofluorescence. Brain regions were identified using a mouse brain atlas (Paxinos and Franklin, 2001), and sections at +0.86 mm from the bregma (dorsal striatum) were selected and processed as follows.

**Day 1.** Free floating sections were rinsed in Tris-buffered saline (TBS; 0.10 M Tris, 0.14 M NaCl), pH 7.4, incubated for 5 min in TBS containing 3% H2O2 (v/v) and 10% methanol (v/v), and rinsed three times 10 min in TBS. After 20-min incubation in 0.2% Triton X-100 in TBS (v/v), sections were rinsed three times in TBS, blocked with BSA 30 g/L in TBS, and incubated overnight (or longer as indicated) at 4°C with the primary antibodies. For detection of phosphorylated proteins, 0.1 M NaF was included in all buffers and incubation solutions. Antibodies were a mouse monoclonal antibody for tyrosine hydroxylase (TH) (1:500 with an incubation ≥ 2 days; Sigma-Aldrich), a rabbit polyclonal antibody for Gaolf protein (1:500) (Herve et al., 2001), rabbit polyclonal antibodies for phospho-Thr202/Tyr204-ERK1/2 (phospho-ERK, 1:400; Cell Signaling Technology), phospho-Ser10-acetyl-Lys14-histone H3 (pAcH3, 1:500, Upstate Ltd, UK), and phospho-(Ser/Thr) PKA substrate (phospho-PKA substrate, 1:500; Cell Signaling Technology).

**Day 2.** Following incubation with primary antibodies, sections were rinsed three times for 10 min in TBS and incubated for 45 min with goat Cy3-coupled (1:400; Jackson Laboratory, Bar Harbor, ME, USA) or goat A488-coupled (1:400; Invitrogen) secondary antibodies. Finally, sections were rinsed for 10 min twice in TBS and twice in Tris buffer (0.25 M Tris, pH 7.4) before mounting in Vectashield (Vector Laboratories). For immunofluorescence detection with Odyssey-LI-COR infrared fluorescent detection system (see below), suitable secondary antibodies were used: IRDye 700DX conjugated anti-rabbit IgG and IRDye 800CW conjugated anti-mouse IgG (Rockland Immunochemical Inc., Gilbertsville, PA, USA), diluted in TBS (1/1000). After rinsing as above, sections were mounted on glass slides and dried 2 h. For determining the TH-depleted area in the lesioned mice, TH immunoreactivity was revealed by immunohistochemistry using biotinylated antibodies, avidine-biotine-peroxidase complex (Vector Laboratories) and 3,3'-diaminobenzidine (DAB) staining (Valjent et al., 2004).

**Confocal microscopy analysis of brain sections**

Confocal microscopy and image analysis were performed at the Institut du Fer à Moulin imaging facility. Images from each region of interest were obtained with a sequential laser-scanning confocal microscope (SP5; Leica) using the same adjustments for all the sections from one experiment. The number of phospho-ERK-positive cells was counted manually in
375 x 375 µm confocal images taken in the lesioned and unlesioned dorsolateral striatum in blind conditions concerning the mouse genotype. MacBiophotonics Image J was used to count automatically the number of nuclei positive for pAcH3 and phospho-PKA substrate. TH and Gαolf immunofluorescence intensity was quantified in the striatum with MacBiophotonics Image J and the data represented mean gray levels above background value in 375 x 375 µm confocal images. The background level was evaluated by measuring the mean gray values in large fiber bundles passing through the striatum. All measurements were carried out in blind conditions regarding the mouse treatment and genotype.

**LI-COR analysis of brain sections**

In some experiments, TH, Gαolf and pAcH3 immunofluorescence intensity was measured using the Odyssey-LI-COR infrared fluorescent detection system (LI-COR, Lincoln, NE, USA). This system commonly used to quantify immunoblots can also be used to quantify immunofluorescence in regions of interest of tissue sections. All sections were scanned together at 21-µm resolution. The regions of interest were drawn as a circle of 1 mm of diameter in the dorsolateral striatum on the lesioned and unlesioned sides. The mean fluorescence intensity was measured in the regions of interest and the background determined in the cerebral cortex, a non-immunoreactive region. The validity of this approach to evaluate Gαolf levels was tested by determining the relationship between the fluorescence intensity and the amount of striatal tissue. Samples containing increasing amounts of Gαolf were prepared by mixing extracts (10 mg/mL protein in 2%, weight/vol SDS) of striatum and cortex that expresses and does not express Gαolf, respectively. Two µL of these samples were spotted on nitrocellulose membrane (Hybond ECL, GE Healthcare Europe, Velizy-Villacoublay, France). After fixation in a Ponceau S solution (Sigma-Aldrich), the membrane was blocked in BSA (30 g/L in TBS) and incubated in anti-Gαolf primary and IRDye-conjugated anti-rabbit secondary antibodies as described above. The mean fluorescence intensity of each dot was measured using the Odyssey-LI-COR detection system and the background determined as the fluorescence intensity of the dots containing only cortex extract. In parallel, free floating brain sections were incubated in the same solutions as the nitrocellulose membrane and mean fluorescence intensity was measured in various striatal areas as described above.

**Immunoblotting**

Mice were sacrificed by decapitation 30 min after L-DOPA plus benserazide injection. The heads of the animals were immediately immersed in liquid nitrogen for 6 s. The brains were removed and the striata dissected out within 20 s on an ice-cold surface, sonicated in 750 µL of 10 g/L sodium dodecyl sulfate (SDS), and placed at 100°C for 10 min. The effectiveness of this extraction procedure in preventing protein phosphorylation and dephosphorylation, hence ensuring that the level of phosphoproteins measured ex vivo reflects the in vivo situation, has been previously tested (Svenningsson et al., 2000). Aliquots (5 µL) of the homogenate were used for protein determination using a bicinchoninic acid assay kit (Pierce Europe, Oud Beijerland, the Netherlands). Equal amounts of protein (20 µg) were separated by 10% (w/v) polyacrylamide gel electrophoresis in the presence of SDS and transferred overnight to polyvinylidene difluoride membranes (Amersham Pharmacia Biotech, Uppsala, Sweden) (Towbin et al., 1979). The membranes were immunoblotted using phospho-Ser845 (Cell Signaling Technology, Beverly, MA) and phosphorylation state-independent (Upstate Biotechnology) glutamate AMPA receptor subunit 1 (GluA1) antibodies, phospho-Thr34 rabbit polyclonal and phosphorylation state-independent mouse monoclonal DARPP-32 antibodies (both gifts of P. Greengard, the Rockefeller University, New York), and TH
monoclonal mouse antibodies (Chemicon, Temecula, CA). Secondary antibodies (1/5000) were IRDye 700DX conjugated anti-rabbit IgG and IRDye 800CW conjugated anti-mouse IgG (Rockland Immunochemical Inc., Gilbertsville, PA, USA). Their binding was quantified using an Odyssey-LI-COR infrared fluorescent detection system (LI-COR, Lincoln, NE, USA). The levels of phosphoproteins were normalized for the amount of the corresponding total protein detected in the sample.

**Statistics**

Correlations between variables were estimated using multiple regression analysis. Multiple comparisons were analyzed using two-way ANOVA followed by Bonferroni post hoc tests for specific comparisons. For 2-group comparisons two-tailed paired or unpaired Student t-test was used.

**RESULTS**

**Gaolf is up-regulated in the 6-OHDA-lesioned dorsolateral mouse striatum**

Increased Gaolf protein levels have been reported in the dorsal striatum of 6-OHDA-lesioned rats (Herve et al., 1993; Marcotte et al., 1994; Penit-Soria et al., 1997; Corvol et al., 2004; Rangel-Barajas et al.) and in the putamen of PD patients (Corvol et al., 2004). In contrast, Gaolf levels after DA depletion had never been explored in the hemiparkinsonian mouse model. In a first experiment we investigated the levels of Gaolf in the dorsal striatum of 6-OHDA-lesioned mice. Twelve mice (55% of the operated mice) survived and were well lesioned [see Methods for criteria]. We evaluated the lesion-induced changes in Gaolf and other markers in striatal sections using two different methods of immunofluorescence measurement with either the Odyssey-LI-COR system (Fig. 1A-D) or confocal microscopy (Fig. 1E, F).

The striatal 6-OHDA injections induced a dramatic decrease in TH immunofluorescence detected with Odyssey-LI-COR (Fig. 1A). TH immunoreactivity consistently disappeared in the most dorsolateral part of the striatum, although the lesion extension in the medioventral part varied from one animal to another. To detect the precise extension of D1R hypersensitivity within the striatum and to measure the levels of Gaolf specifically in this region, we used the L-DOPA-induced increase in phospho-Ser10, acetyl-Lys14 histone H3 (pAcH3), a prominent response directly linked to D1R hypersensitivity (Santini et al., 2009b; Cenci and Konradi, 2010). Lesioned mice were treated with L-DOPA (20 mg/kg) and benserazide (12 mg/kg) 30 min before the sacrifice, a treatment which does not alter Gaolf levels (Corvol et al., 2004 and Alcacer and Hervé, unpublished observations), and the distribution of pAcH3-positive cells was determined in the striatum (Fig. 1B). An intense pAcH3 signal was detected in response to L-DOPA treatment in the dorsolateral part of the TH-depleted striatum (Fig. 1B). Interestingly the area of strong pAcH3-immunoreactivity was consistently more restricted than the TH-depleted zone (Fig. 1A, B).

In order to validate the use of Odyssey-LI-COR for evaluating Gaolf levels in tissue, we compared immunofluorescence signals measured in striatal sections with those obtained with various concentrations of striatal homogenates in dot blot (Fig. 1C). In dot blot, fluorescence intensity was linearly related with the concentration of striatal tissue (and thus of Gaolf) on a wide scale. Using this approach, Gaolf immunofluorescence was found to increase in the dorsolateral part of the striatum where TH immunofluorescence was absent and pAcH3
present, as compared to the contralateral side (Fig. 1D). Very similar results were obtained using confocal microscopy (Fig. 1E and F). Since quantification of images from confocal microscope provided results similar to those with Odyssey-LI-COR, we used confocal imaging in the rest of the study. Thus, our results revealed a significant increase in Gαolf immunoreactivity within the DA-deficient region of the striatum of 6-OHDA-lesioned mice, coincident with the area of increased D1R signaling.

**Striatal Gαolf increase correlates with dyskinetic behavior**

Since Gαolf immunoreactivity was higher in the mouse dorsolateral striatum after the lesion, we examined whether this increase was maintained after chronic treatment with L-DOPA and whether it was associated with the development of LID. To address this question 6-OHDA-lesioned mice received a 10-day treatment with L-DOPA (20 mg/kg per day) and benserazide (12 mg/kg) referred to as “L-DOPA” below. In a series of six experiments, 65 mice (68% of the operated mice) survived the operation and displayed a decrease in striatal TH levels greater than 80%. One group received a single injection of L-DOPA (n = 21), another group was chronically treated with L-DOPA during 10 days (n = 44). An additional control group of 6 sham-lesioned mice also received L-DOPA chronically. Thirty minutes after the last injection of L-DOPA, all the animals were sacrificed to compare the Gαolf levels in their 6-OHDA-injected and intact striata by confocal immunofluorescence analysis. For comparison between different animals and experiments we used the immunofluorescence on the lesioned side/immunofluorescence on the unlesioned side ratio (Fig. 2A). As expected, in the sham-operated mice, the Gαolf immunofluorescence intensities were similar in the vehicle-injected and intact striata (Fig. 2A). In contrast, striatal Gαolf immunofluorescence was significantly higher in the 6-OHDA-lesioned striatum than in the contralateral striatum in mice that had received one or repeated injections of L-DOPA (Fig. 2A).

Since among chronically L-DOPA-treated mice some animals developed intense LID but not others, we examined whether the levels of Gαolf differed between these two groups. To perform this analysis, we separated the highly dyskinetic mice (i.e. those having total AIMs scores in the quartile of highest values, higher than 33.5, n = 11) and the weakly dyskinetic mice (total AIMs score in the quartile of lowest values, lower than 12.5, n = 11), among a total number of 44 animals in 3 distinct experiments. The mice in both groups displayed more than 80% reduction in TH in the dorsolateral striatum (data not shown). In lesioned mice, we did not detect any significant correlation between TH depletion and AIM score (data not shown). In the quartile of highly dyskinetic mice, Gαolf immunofluorescence was higher in the lesioned striatum than in the contralateral unlesioned striatum (Fig. 2B). In contrast, in the quartile of weakly dyskinetic mice Gαolf immunofluorescence was similar on the two sides (Fig. 2B). We then examined whether there was a correlation between the levels of Gαolf and the intensity of AIMs, including all the lesioned animals that had received a chronic L-DOPA treatment. Although the experimental points appeared quite disperse, statistical analysis revealed a significant positive correlation between Gαolf immunofluorescence and the total AIMs score (Fig. 2C), the locomotor component of LID (LOC, Fig. 2D), and the composite score of axial-limb-orofacial (ALO) AIMs (Fig. 2E), which have been suggested to be more similar to the human LID than the locomotive AIMs (Lundblad et al., 2002). From these experiments, we concluded that the increase in Gαolf in the dorsolateral striatum of hemiparkinsonian mice was somehow associated with the severity of dyskinetic behavior. This result suggested a possible implication of Gαolf striatal up-regulation in the development of LID in the mouse model of hemiparkinsonism.
Effects of 6-OHDA lesions on Gαolf in Gnal +/- mice

Given the positive correlation between Gαolf increases and dyskinesia, we used a genetic model in which Gαolf levels are decreased to examine the consequences of this reduction on LID and signaling responses. Homozygous Gαolf knockout mice (Gnal -/-) have a severe phenotype due to the combination of olfactory and striatal deficits (Belluscio et al., 1998; Zhuang et al., 2000; Corvol et al., 2001; Herve et al., 2001). These mice usually die in the early post-natal period and could not be used in our study. In contrast, Gnal +/- mice which develop and breed normally, provide a very interesting model since they display a decrease of about 50% in Gαolf protein levels (Fig. 3A) and a 30-50% deficit in D1R- or A2AR-activated cAMP production in vitro when compared to wild type littermates (Corvol et al., 2001; Corvol et al., 2004). Since striatal Gαolf immunofluorescence was increased by about 50% in the lesioned striatum of dyskinetic mice whereas it was similar to the unlesioned tissue in weakly dyskinetic mice (see Fig. 2B), Gnal +/- mice in which Gαolf was reduced but not absent were a good model for testing the specific implication of Gαolf up-regulation.

Wild type (Gnal +/-) and heterozygous (Gnal +/) mice were lesioned with 6-OHDA and, after a 4-week recovery, treated with L-DOPA during 10 days (17 well lesioned mice in each group survived the whole experimental procedure, corresponding to about 60% of operated mice). They were killed 30 min after the last L-DOPA injection and proteins of interest were analyzed by immunofluorescence or immunoblotting. Immunoblotting showed that the two groups of mice displayed the same degree of TH reduction in their lesioned striatum indicating that 6-OHDA had a similar ability to destroy DA innervation in the two genotypes (Fig. 3B). Immunohistochemistry indicated that the TH-depleted area was similar in wild type and Gnal +/- mice (87 ± 2 % and 92 ± 2% of the total striatum area, respectively, Student t-test, t = 1.48, DF = 18, not significant). To compare the changes in Gαolf immunofluorescence in mutant and wild type mice, three sections from each animal were labeled and quantified using confocal imaging in 3 independent experiments and the results were averaged (Fig. 3C). As expected, the levels of Gαolf detected with this method were lower in unlesioned striatum of mutant mice compared to wild type (Fig. 3C). Interestingly, Gαolf immunofluorescence levels in Gnal +/- mice were increased in the lesioned striatum when compared to the contralateral intact striatum (Fig. 3C). The relative increase on the lesioned side compared to the contralateral side was similar in the two genotypes (Gnal +/- 132 ± 5 %, Gnal +/+, 145 ± 6 %, mean ± SEM, Student t-test t = 1.6, DF = 18, not significant). However, the levels of Gαolf immunofluorescence in the lesioned striatum of mutant mice remained below those in the intact striatum of wild type mice (-16%, Fig. 3C). Similar results were observed using the Odyssey-LI-COR system (data not shown). Altogether these results showed that the mechanisms of Gαolf increase following 6-OHDA lesion were unaltered in heterozygous mutant mice, but that the gene dosage effect maintained the levels of the protein below those in unlesioned wild type mice.

Behavioral responses of hemiparkinsonian Gnal +/- mice are similar to wild type

During L-DOPA treatment, the 6-OHDA-lesioned Gnal +/- and Gnal +/- mice were evaluated in behavioral tests. Mice were first subjected to the cylinder test before (baseline) and after the first administration of L-DOPA. Before L-DOPA, Gnal +/- and Gnal +/- mice displayed a marked reduction in the usage of the forelimb contralateral to the lesion (Fig. 4A), reflecting the hypokinetic effects of 6-OHDA-lesion. A similar recovery of the forelimb usage
was observed in both genotypes 1 h after L-DOPA treatment. Thus, the anti-parkinsonian effect of L-DOPA appeared to be preserved in Gnal +/− mice.

At the end of the 10-day L-DOPA treatment, the animals were scored for the various types of AIMs. No significant difference was observed between Gnal +/- and Gnal +/+ mice in axial, limb and orofacial AIMs induced by chronic L-DOPA treatment, and the ALO composite score was not significantly changed in the mutant mice (Fig. 4B). Similarly, no significant difference in locomotive score was observed between the two genotypes (Fig. 4C). The total AIMs score and the time course of L-DOPA effects were not significantly affected by Gnal haplodeficiency (Fig. 4D). We also compared LID in Gnal +/- and wild type mice after only 5 days of L-DOPA treatment (Fig. 4E) and we detected no significant difference between the two genotypes, suggesting the absence of alteration in the LID development in Gnal +/− mice. Thus our results clearly showed that the partial deficit in Gaolf in heterozygous mice did not change the incidence and intensity of LID following a 10-day treatment.

L-DOPA-induced cAMP-dependent phosphorylation is dramatically reduced in the lesioned striatum of Gnal +/− mice

We have previously reported a dissociated alteration of signaling pathways in Gnal +/- mice in response to psychostimulants in vivo, with markedly decreased cAMP-dependent phosphorylation and normal ERK signaling (Corvol et al., 2007). It was therefore particularly interesting to examine the signaling responses to L-DOPA in Gnal +/- mice. Since Gaolf is directly involved in the stimulation of cAMP/PKA signaling (Corvol et al., 2007), we compared the phosphorylation of several PKA substrates in chronically L-DOPA-treated Gnal +/- and wild type mice. In Gnal +/+ (n = 9) and Gnal +/- (n = 11) mice, we evaluated the number of neurons immunolabeled with an antibody recognizing the phosphorylated form of the PKA substrate consensus sequence, which has been validated for assessing the activity of cAMP/PKA pathway in neurons by immunofluorescence (Sindreu et al., 2007). A dramatic increase in the number of immunopositive neurons was observed after L-DOPA in the lesioned striatum of wild type mice showing a strong responsiveness of cAMP/PKA signaling (Fig. 5A and B). This increase was significantly lower in the lesioned striatum of Gnal +/- indicating a reduction of cAMP/PKA signaling in these mice. The persistence of some level of activation may correspond to a persistent partial PKA activation or to the reaction of the antibody with proteins phosphorylated by other kinases. Therefore, in a different set of lesioned animals, we examined by immunoblotting the phosphorylation of two well characterized PKA substrates, AMPA receptor subunit GluA1 at Ser845 (Snyder et al., 2000) and DARPP-32 at Thr34 (Hemmings et al., 1984). Phosphorylation of these sites has been shown to be correlated with the occurrence of LID (Santini et al., 2007). In the lesioned striatum of Gnal +/- mice, L-DOPA treatment induced an increased phosphorylation of GluA1 and DARPP-32 when compared with the unlesioned striatum (Fig. 5C-E), in agreement with previous results (Santini et al., 2007). In contrast, these responses were absent in Gnal +/- mice for Ser845-GluA1 (Fig. 5C, D) and markedly diminished for Thr34-DARPP-32 (Fig. 5C, E). A slight increase in baseline DARPP-32 phosphorylation was observed in Gnal +/- mice but it was not significant and of uncertain meaning given the variability in baseline DARPP-32 phosphorylation between animals. These results show that following a chronic treatment with L-DOPA, in vivo activation of the cAMP/PKA phosphorylation pathway is severely altered in Gnal +/- mice, in spite of the relative increase
in Goolf protein induced by the lesion as compared to the unlesioned striatum. This suggests a threshold effect of the amount of Goolf for cAMP/PKA pathway activation.

Hypersensitivity of L-DOPA-induced activation of the ERK pathway is preserved in the lesioned striatum of Gnal +/- mice

The second signaling pathway that has been reported to be strongly activated by L-DOPA in the lesioned striatum is the ERK cascade (Gerfen et al., 2002; Pavon et al., 2006; Santini et al., 2007; Westin et al., 2007). The activation of this pathway is D1R-dependent (Gerfen et al., 2002; Westin et al., 2007) and takes place in striatonigral MSNs which express this receptor (Santini et al., 2009b), although prolonged L-DOPA treatment was found to activate ERK pathway in the small population of cholinergic interneurons (Ding et al., 2010). Thirty minutes after injection of L-DOPA, following a 10-day treatment, we observed a dramatic increase in the number of neurons immunofluorescent for the di-phosphorylated active form of ERK1/2 (pERK) in the lesioned striatum as compared to the unlesioned side in Gnal ++/− mice (Fig. 6A). A similar response was observed in Gnal +/− mutant mice (Fig. 6A). We also analyzed the phosphorylation of ERK2 by immunoblotting and found no difference between the two genotypes (pERK2/total ERK2: ++/+, unlesioned, 100 ± 4, 6-OHDA, 142 ± 11; +/-, unlesioned, 102 ± 3, 6-OHDA 141 ± 6, means ± SEM. Two-way ANOVA, lesion effect F(1,57) = 42.1 p < 0.001, genotype effect F(1,57) = 0.01, not significant, interaction F(1,57) = 0.03, not significant). We then examined pAcH3 immunoreactivity that is known to be dependent on ERK activation and is strongly increased by L-DOPA in 6-OHDA-lesioned striatonigral MSNs (Santini et al., 2007; Santini et al., 2009b). A dramatic increase in the number of pAcH3-positive neurons was observed in the lesioned striatum of wild type mice (Fig. 6B). A virtually identical response was observed in Gnal +/− mice (Fig. 6B). Thus, there was a marked contrast between the identical pERK responses to L-DOPA in Gnal ++/− and +/− mice, and the clear reduction in the PKA pathway response in Gnal +/− mice.

Since chronic treatment by L-DOPA may change the response of ERK pathway (Santini et al., 2007; Cenci and Konradi, 2010; Ding et al., 2010), we wondered whether this treatment could blunt differences between Gnal ++/− and +/− mice. We therefore examined in a different batch of mice the responses to a single acute injection of L-DOPA. A very strong increase in pERK-positive (Fig. 6C) and pAcH3-positive (Fig. 6D) cells was observed in the lesioned striatum and this effect was identical in wild type and heterozygous Gnal mutant mice. These results clearly show that the decrease in Goolf did not alter ERK activation in response to L-DOPA.
DISCUSSION

The present study addressed the involvement of Gαolf up-regulation in LID in a mouse model of hemiparkinsonism. The use of mice partially deficient in Gαolf (Gnal +/-) allowed us to determine its contribution to LID and to intracellular signaling cascades presumably involved in LID development or expression.

**Gαolf up-regulation in dyskinetic mice**

We replicated in mice the up-regulation of Gαolf protein observed in the dorsal striatum of hemiparkinsonian rats and PD patients (Herve et al., 1993; Marcotte et al., 1994; Penit-Soria et al., 1997; Corvol et al., 2004; Rangel-Barajas et al., 2010). Previous data suggested that this up-regulation was post-transcriptional (Herve et al., 1993) and resulted from a lack of usage of the receptor (Herve et al., 2001). Although its precise mechanisms are still unknown, a parallel can be made with the highly related Gαs whose degradation increases following its activation in cultured cells (Levis and Bourne, 1992). In support of this hypothesis treatment of lesioned rats with L-DOPA or a D1R agonist normalized Gαolf levels (Corvol et al., 2004; Rangel-Barajas et al., 2010). However, Gαolf increase was observed in the putamen of PD patients who had received a prolonged L-DOPA treatment (Corvol et al., 2004). The present study shows a correlation between LID and maintenance of increased Gαolf levels. When the mice were stratified on AIMS intensity, animals with no or minimal dyskinesia had normal levels of Gαolf, whereas those with strong dyskinesia had the highest levels. If the degree of D1R usage was the only factor regulating Gαolf levels, their normalization should have been observed in all groups. Therefore our results, as those in patients (Corvol et al., 2004), suggest the existence of a dysregulation associated with the occurrence of dyskinesia and responsible for the “abnormal” persistence of high levels of Gαolf. It should be noted that no persistent increase in Gαolf was observed in dyskinetic L-DOPA-treated rats (Rangel-Barajas et al., 2010). We do not know whether this discrepancy results from differences between species or experimental protocols. Yet, the persistence of Gαolf increase in treated PD patients (see above) suggests that results in mice are relevant for the human disease.

**Gαolf up-regulation contributes to the L-DOPA activation of the D1R/cAMP/PKA pathway in the lesioned striatum**

The molecular mechanisms leading to the hypersensitivity of D1R-stimulated adenylyl-cyclase activity and signaling following DA lesion are unclear since the overall D1R concentration and its abundance at the plasma membrane are not increased (Savasta et al., 1988; Herve et al., 1993; Shinotoh et al., 1993; Berthet et al., 2009). Our results strongly support the role of Gαolf up-regulation in D1R-hypersignaling, since a marked decrease in the PKA stimulation was observed in Gnal +/- mutant mice. L-DOPA-induced phosphorylation was partially (general PKA-phosphosubstrates) or massively (phosphorylation of GluA1 or DARPP-32) decreased. Since in mutant mice the levels of Gαolf were below those in unlesioned wild type striatum, we can conclude that the increase in Gαolf is critical for D1R-dependent cAMP/PKA hypersensitivity. These results are in agreement with a previous study showing that the levels of Gαolf but not those of D1R are critical for the efficacy of the D1R/cAMP/PKA pathway (Corvol et al., 2007). They also account for the increased G-protein activation by D1R reported in MPTP-lesioned monkeys.
Additional factors may contribute to the cAMP/PKA hypersensitivity in DA-lesioned striatum, including increased levels of adenylyl-cyclase5/6 (Rangel-Barajas et al., 2010), or changes in phosphatases or phosphodiesterases levels and/or activities (Meurers et al., 2009).

**Gαolf/cAMP/PKA hypersensitivity does not account for the strong activation of ERK by L-DOPA in dyskinetic animals**

In animal models of PD, a strong ERK activation in response to L-DOPA is observed, correlated with the occurrence of LID (Gerfen et al., 2002; Pavon et al., 2006; Santini et al., 2007; Westin et al., 2007; Nicholas et al., 2008; Santini et al., 2009b; Fasano et al., 2010; Santini et al., 2010; Francardo et al., 2011). ERK activation occurs in D1R-expressing striatonigral neurons (Gerfen et al., 2002; Santini et al., 2009b), and following prolonged treatment in cholinergic interneurons (Ding et al., 2010). The ERK pathway may be important at several levels including regulation of transcription (Cenci et al., 1998; Aubert et al., 2007) and translation, through the mTORC1 complex (Santini et al., 2009a). The precise molecular mechanism of ERK-signaling hypersensitivity is not known. In the present study, despite a dramatic reduction in cAMP/PKA response, the L-DOPA-induced ERK activation was not affected by Gnal haplodeficiency. This result strongly suggests that the ERK hyper-responsiveness following DA degeneration does not result from the hypersensitivity of the cAMP/PKA pathway and that the two responses can be dissociated. These results are in line with our previous data on responses to amphetamine in Gnal +/- mice, showing no alteration in the ERK activation despite dramatic reduction in PKA signaling activity (Corvol et al., 2007). A cAMP-independent synergy of D1R and NMDAR for the ERK activation exists in neurons in culture (Pascoli et al., 2011). In vivo an amplification of this synergy by cAMP/PKA/DARPP-32 appears critical, possibly due to the high concentrations of regulatory proteins like DARPP-32 and striatal-enriched tyrosine phosphatase (STEP) (Valjent et al., 2005). The cause of the D1R-activated ERK pathway hypersensitivity remains to be identified in DA-lesioned mice, but could involve cAMP-independent mechanisms (Gerfen et al., 2008), including crosstalk between D1R and mGluR5 receptor (Rylander et al., 2009) or dysregulation of protein phosphatases (Meurers et al., 2009). It should be underlined however that a basal tonus of cAMP/PKA/DARPP-32 pathway appears required for ERK activation. For instance, the dramatic impairment of the cAMP pathway in Gnal +/- homozygous mutants prevented in vivo ERK activation by amphetamine (Corvol et al., 2007). In addition, in vivo complete inhibition of PKA (Lebel et al., 2010) or genetic deletion of an amplifier of PKA action, DARPP-32 (Santini et al., 2007), decreased the L-DOPA-induced ERK activation of in lesioned mice. Accordingly the existence of a basal level of phospho-DARPP-32 in Gnal +/- mice may play a permissive role for ERK activation.

**Hypersensitivity of the D1R/Gαolf/cAMP pathway is not responsible for dyskinetic behavior but have a permissive role**

A variety of dysregulations take place in DA-denervated striatum and are likely to concur to the occurrence of LID (Cenci, 2007; Jenner, 2008; Santini et al., 2008; Calabresi et al., 2010; Feyder et al., 2011; Gerfen and Surmeier, 2011; Murer and Moratalla, 2011). From a therapeutic stand-point it is important to dissect the mechanisms of induction and expression of LID, which can be targets for potential preventive or symptomatic treatments. Interfering with D1R (Westin et al., 2007; Darmopil et al., 2009), cAMP/PKA (Lebel et al., 2010),
DARPP-32 (Santini et al., 2007; Bateup et al., 2010), Ras-GRF1/ERK (Santini et al., 2007; Fasano et al., 2010), and mTOR (Santini et al., 2009a) decreases or prevents LID. Although these results suggest the involvement of all these signaling modules in the generation of LID, they do not prove that LID results from their up-regulation. They only show that their activity is necessary. Our results clarify this situation by showing that LID was still observed in \textit{Gnal} +/- mice in spite of a marked, but not complete, decrease in D1R/cAMP/PKA signaling. In addition, in \textit{Gnal} +/- mice the A2A receptor/cAMP/PKA signaling is known to be blunted (Corvol et al., 2001; Herve et al., 2001). A decrease in A2A signaling would be expected to reduce LID development (Xiao et al., 2006), but LID was not decreased in \textit{Gnal} +/- mice. Therefore we can rule out that increase of cAMP/PKA response is the main culprit for the occurrence of LID. As discussed above for ERK activation however, cAMP/PKA signaling appears necessary for LID. Indeed, the marked reduction of LID induced by pharmacological inhibition of PKA (Lebel et al., 2010) or by general or targeted DARPP-32 knockout (Santini et al., 2007; Bateup et al., 2010) shows the importance of this pathway. On the other hand, our results are compatible with a critical role of ERK in LID generation since its activation was similar in \textit{Gnal} +/- and wild type mice.

In conclusion, this study allows better understanding the importance of various aspects of D1R-signaling hypersensitivity in the generation of LID and of their relationships. Our results suggest several important conclusions: \textit{i}, \textit{G}αolf up-regulation is a conserved response to DA-lesion and its persistence during L-DOPA treatment correlates with LID; \textit{ii}, this up-regulation contributes to the hypersensitivity of the D1R/cAMP/PKA pathway, which was markedly decreased following reduction in \textit{G}αolf gene dosage; \textit{iii}, hypersensitivity of cAMP/PKA signaling does not explain ERK strong activation, which was unaltered in \textit{Gnal} mutant mice; \textit{iv}, although the cAMP/PKA pathway is necessary for LID occurrence, its hypersensitivity is unlikely to be its main cause. These results are compatible with a model in which a minimal level of cAMP/PKA pathway is necessary for LID generation, whereas increased ERK signaling may play a direct causal role. Identification of the mechanisms leading to hypersensitivity of the ERK pathway following lesion of DA neurons is therefore an important objective of future studies.
References


Figure Legends

Figure 1
Unilateral 6-OHDA lesion increases Gaolf levels in the dorsal striatum. Immunoreactivity was detected using Odyssey-LI-COR system (A-D) or confocal microscope (E, F). A, B, D: Serial coronal brain sections from a mouse that received a unilateral 6-OHDA injection into the right striatum. (A) Tyrosine hydroxylase immunofluorescence (TH IF). (B) Phospho-acetyl-histone H3 immunofluorescence (pAcH3 IF) showing the extent of signaling hypersensitivity in the dorsolateral part of the lesioned striatum. (C) Plot of Gaolf IF quantification with Odyssey-Li-Cor, as a function of the proportion of Gaolf protein in a dot blot assay. The samples were obtained by mixing various proportions of protein extracts from striatum (Str) expressing Gaolf and cortex (Cx) devoid of Gaolf. The immunofluorescence values were determined in parallel in two regions of the striatum, in two brain sections (i.e., non-homogenized striatal slices, right panel, Striatal regions). These values were within the linear range of the standard curve. (D) Gaolf immunofluorescence (Gaolf IF). Quantification of mean immunofluorescence intensity (IF) on the unlesioned (UL) and lesioned (6-OHDA) sides is shown in A, B, D in the dorsolateral part of the striatum (circled region of interest drawn in A). Data are means ± SEM (n = 11-13). Paired two-tailed Student’s t-test: TH: t = 4.57; pAcH3: t = 12.8; Gaolf: t = 4.00. Scale bar, 1 mm. (E and F) Single confocal sections in the dorsolateral striatum showing TH (E) and Gaolf immunofluorescence (F). Data are means ± SEM (n = 8-12). Paired two-tailed Student’s t-test: TH: t = 4.47, Gaolf: t = 3.19. Scale bar, 100 µm. **p < 0.01, ***p < 0.001. a.u. arbitrary units.

Figure 2
Striatal Gaolf levels increase correlates with dyskinetic behavior. (A) Quantification of Gaolf immunofluorescence (IF) in the dorsal striatum of sham-operated mice treated with L-DOPA for 10 days (Sham, n = 6), 6-OHDA-lesioned mice treated with a single injection of L-DOPA (Acute, n = 21) or daily for 10 days (Chronic, n = 44). Data are expressed as percentage of the unlesioned (UL) side in each group and are means ± SEM. One-way ANOVA F(2,68) = 4.16, *p < 0.05. Post hoc comparison (Bonferroni’s test): *p < 0.05 vs sham. (B) Comparison of Gaolf levels in the 25% 6-OHDA-lesioned chronically L-DOPA-treated mice in (A) that developed the weakest (total AIM score below 12.5, Low Dysk n = 11) and strongest (above 33.5, High Dysk, n = 11) dyskinesia. Data are means ± SEM. Unpaired two-tailed Student’s t-test: t = 2.89, **p < 0.01. (C-E) Correlation between Gaolf levels and (C) total AIMs score (r = 0.40, F(1,42) = 7.96, p < 0.01), (D) LOC (locomotor) AIMs (r = 0.41, F(1,42) = 8.56, p < 0.01), and (E) ALO (axial, limb, orofacial) AIMs (r = 0.34, F(1,42) = 5.38, p < 0.05).

Figure 3
Gaolf and tyrosine hydroxylase levels in 6-OHDA-lesioned Gnal+/− and control mice (A) Quantification of Gaolf levels in mice heterozygous for a null mutation of Gnal gene (+/−) and their wild type littermates (+/+ ) by immunoblotting. Data are expressed as percentage of the mean in wild type mice and are means ± SEM (n = 4-6). Unpaired two-tailed Student’s t-test: Gaolf t = 10.93, ***p < 0.0001. (B) Quantification of TH levels by immunoblotting. Data are expressed as percentage of the means of unlesioned (UL) side of Gnal +/+ mice and
are means ± SEM (n = 7-8). Two-way ANOVA: effect of lesion: F(1,24) = 357, p < 0.0001; effect of genotype: F(1,24) = 0.01, not significant (NS); interaction: F(1,24) = 0.90, NS. Post hoc comparison (Bonferroni’s test): *** p < 0.001, 6-OHDA vs. unlesioned (UL) for each genotype; (C) Quantification of Guolf mean immunofluorescence (IF) intensity in 6-OHDA-lesioned Gnal +/+ and wild type control (+/) mice chronically treated with L-DOPA. Guolf IF was measured in the dorsolateral striatum by confocal microscopy in 3 independent experiments for each mouse and normalized to the mean values in wild type mice for each experiment. Two-way ANOVA analysis effect of the lesion: F(1,18) = 133, p < 0.0001; effect of the genotype: F(1,18) = 13.97, p < 0.01; interaction: F(1,18) = 1.81, NS. Post hoc comparison (Bonferroni’s test): *** p < 0.001, 6-OHDA-lesioned vs. UL; ## p < 0.01 and ### p < 0.001, Gnal +/+ vs. Gnal +/-.

Figure 4

L-DOPA-induced dyskinesia is not altered in Gnal heterozygous mice. Heterozygous (+/-) Gnal mutant and wild type (+/+) mice were lesioned and treated with L-DOPA and benserazide during 10 days. (A) Forelimb use was determined using the cylinder test in the 6-OHDA lesioned mice before (-) and after administration of L-DOPA (+) on the first day of treatment. Data are means ± SEM (n = 7-8). Repeated-measure ANOVA (with the within-subjects factor of treatment and the between-subjects factor of genotype): effect of treatment: F(1,26) = 56, p < 0.0001; effect of genotype: F(1,26) = 0.49, NS; interaction between genotype and treatment: F(1,26) = 0.09, NS. Post hoc comparison (Bonferroni’s test): *** p < 0.001, before vs after L-DOPA. (B) Sum of axial, limb and orolinguinal (ALO) AIMs scored during 140-min period after L-DOPA. Comparison between +/- (n = 18) and +/- (n = 23) mice. Data are means ± SEM. Unpaired two-tailed Student’s t-test: t = 0.62, NS. (C) Sum of locomotive (LOC) AIMs scored during 140-min period after L-DOPA in the same animals. Data are means ± SEM. Unpaired two-tailed Student’s t-test: t = 0.61, NS. (D) Time course of total AIMs (sum of axial, limb, orofacial and locomotive AIMs) scored every 20 min over a period of 140 min after the last L-DOPA administration. Repeated-measure two-way ANOVA (with the within-subjects factor of time and the between-subjects factor of genotype): effect of genotype F(1,234) = 0.18, NS, effect of time F(6,234) = 71, p < 0.0001; interaction: F(6,234) = 0.43, NS. (E) Total AIMs (sum of axial, limb, orofacial and locomotive AIMs) scored during 140-min period after L-DOPA on days 5 of L-DOPA treatment. Data are means ± SEM. Student t-test, t = 0.181, DF = 25, NS.

Figure 5

L-DOPA-induced PKA-dependent phosphorylation is markedly impaired in Gnal heterozygous mice. (A) Single confocal section showing phospho-PKA substrate immunofluorescence in the unlesioned (UL) and 6-OHDA-lesioned dorsal striatum of Gnal +/- (upper panels) and Gnal +/+ (lower panels) mice, 30 min after the last injection of L-DOPA + benserazide. Scale bar, 100 µm. (B) Number of phospho-PKA substrate (pPKA substrate) positive cells in the dorsolateral striatum of wild type and heterozygous mice. Data are means ± SEM of positive cells in 375 x 375 µm confocal images (n = 8-11). Repeated-measure two-way ANOVA (with the within-subjects factor of lesion and the between-subjects factor of genotype): effect of lesion: F(1,234) = 0.18, NS, effect of time F(6,234) = 71, p < 0.0001; interaction: F(6,234) = 0.43, NS. (C) Total AIMs (sum of axial, limb, orofacial and locomotive AIMs) scored during 140-min period after L-DOPA on days 5 of L-DOPA treatment. Data are means ± SEM. Student t-test, t = 0.181, DF = 25, NS.
GluA1, phospho-Thr34-DARPP-32 (pThr34) and total DARPP-32 in the unlesioned (UL) and 6-OHDA-lesioned striatum of \textit{Gnal} +/+ and \textit{Gnal} +/- mice, 30 min after the last injection of L-DOPA + benserazide. (D) Quantification of pSer845 normalized to total GluA1, expressed as percentage of unlesioned (UL) striatum of \textit{Gnal}+/+ mice. Data are means ± SEM (n = 9-15). Two-way ANOVA, effect of the genotype; F_{(1,48)} = 6.02, p < 0.02; effect of the lesion: F_{(1,48)} = 10.8, p < 0.01; interaction: F_{(1,48)} = 6.8, p < 0.05. Post hoc comparison (Bonferroni’s test): *** p < 0.001, 6-OHDA vs. UL; ## p < 0.01 \textit{Gnal} +/- vs. \textit{Gnal} +/+. (E) Same representation for pThr34 normalized to total DARPP-32. Data are means ± SEM (n = 5-9). Two-way ANOVA, effect of the genotype; F_{(1,22)} = 1.13, NS; effect of the lesion: F_{(1,22)} = 15.5, p < 0.001; interaction: F_{(1,22)} = 10.1, p < 0.01. Post hoc comparison (Bonferroni’s test): *** p < 0.001, 6-OHDA vs. UL; # p < 0.05 \textit{Gnal} +/- vs. \textit{Gnal} +/-.

**Figure 6**

**Chronic or acute L-DOPA induces phosphorylation of ERK and histone H3 in the lesioned striatum of \textit{Gnal} heterozygous mice.** (A-B) Confocal sections through the dorsolateral striatum of unlesioned (UL) and 6-OHDA-lesioned dorsolateral striatum of \textit{Gnal} +/+ (upper panels) and \textit{Gnal} +/- (lower panels) mice, 30 min after the last injection of chronic L-DOPA and benserazide treatment showing phospho-ERK (pERK, A) and phospho-acetyl-histone H3 (pAcH3, B) immunofluorescence. Scale bar, 100 µm. Lower panels, number of pERK or pAcH3 positive cells in the dorsolateral striatum. Data are means ± SEM in 375 x 375 µm confocal images (n = 8-11). Repeated-measure two-way ANOVA (with the within-subjects factor of lesion and the between-subjects factor of genotype) (A) pERK: effect of the lesion F_{(1,17)} = 67, p < 0.0001 effect of the genotype: F_{(1,17)} = 0.89, NS; interaction F_{(1,17)} = 1.03, NS; (B) pAcH3: effect of lesion: F_{(1,17)} = 57, p < 0.0001; but no significant effect of genotype: F_{(1,17)} = 0.02, NS; and no significant interaction between genotype and lesion F_{(1,17)} = 0.01, NS. Post hoc comparison (Bonferroni’s test): *** p < 0.001, 6-OHDA vs. UL. (C, D) Same as in A, B but in different groups of mice which received a single injection of L-DOPA (n = 4-8 per group). (C) pERK, effect of lesion F_{(1,20)} = 893, p < 0.0001; effect of genotype: F_{(1,20)} = 0.79, NS; interaction: F_{(1,20)} = 0.79; NS. (D) pAcH3, effect of lesion: F_{(1,20)} = 1231, p < 0.0001; effect of genotype: F_{(1,20)} = 3.61, NS; interaction F_{(1,20)} = 3.61, NS. Post hoc comparison (Bonferroni’s test): *** p < 0.001, 6-OHDA vs. UL.
Fig. 1

A TH
Unlesioned 6-OHDA

B pAcH3
Unlesioned 6-OHDA

C Gaolf

D Gaolf

E TH
Unlesioned 6-OHDA

F Gaolf
Unlesioned 6-OHDA

Striatal regions
Fig. 2

A) Bar graph showing the percentage of Gaolf IF compared to the unlesioned side in Sham, Acute, and Chronic conditions. The percentage values are 100, 150, and 200, respectively, with significant differences indicated by *.

B) Bar graph showing the percentage of Gαolf%IF compared to the unlesioned side for Low Dysk and High Dysk conditions. The percentage values are 100 and 150, respectively, with significant differences indicated by **.

C) Scatter plot showing the relationship between Total AIMs score and Gaolf IF, percentage of unlesioned side, with a linear trend line.

D) Scatter plot showing the relationship between Total LOC score and Gaolf IF, percentage of unlesioned side, with a linear trend line.

E) Scatter plot showing the relationship between Total ALO score and Gaolf IF, percentage of unlesioned side, with a linear trend line.
Fig. 3

A

G\textalpha{olf} levels, % of wild type

\begin{tabular}{lcc}
Gnal & +/+ & +/- \\
\hline
0 & 100 & 100 \\
50 & 75 & 75 \\
100 & 50 & 50 \\
150 & 25 & 25 \\
\end{tabular}

B

TH levels, % of unlesioned side

\begin{tabular}{lcccc}
UL & 6-OHDA & \\
\hline
Gnal +/+ & Gnal +/- & \\
\hline
0 & 120 & 120 \\
50 & 80 & 80 \\
100 & 40 & 40 \\
150 & 0 & 0 \\
\end{tabular}

C

G\textalpha{olf} IF, % of +/+ unlesioned side

\begin{tabular}{lcc}
Gnal & +/+ & +/- \\
\hline
0 & 100 & 100 \\
50 & 75 & 75 \\
100 & 50 & 50 \\
150 & 25 & 25 \\
\end{tabular}

***

##

###
**Fig. 5**

A. **Phospho-PKA-substrate**

B. **pPKA-substrate positive cells**

C. **GluA1**

D. **pSer845/total GluA1**

E. **pThr34/total DARPP-32**
Fig. 6

A. pERK

B. pAcH3

C. Acute L-DOPA

D. Acute L-DOPA

Chronic L-DOPA

Unlesioned 6-OHDA

Gnal +/+ 6-OHDA

Gnal +/- 6-OHDA

Chronic L-DOPA

Unlesioned 6-OHDA

Gnal +/+ 6-OHDA

Gnal +/- 6-OHDA

Acute L-DOPA

Unlesioned 6-OHDA

Gnal +/+ 6-OHDA

Gnal +/- 6-OHDA

UL

6-OHDA

pERK-positive cells

pAcH3-positive cells

NS

***

UL

6-OHDA

pERK-positive cells

pAcH3-positive cells

NS

***

UL

6-OHDA

pERK-positive cells

pAcH3-positive cells

NS

***

UL

6-OHDA

pERK-positive cells

pAcH3-positive cells

NS

***

UL

6-OHDA