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Down-Regulation of GABA<sub>A</sub> Receptor via Promiscuity with the Vasoactive Peptide Urotensin II Receptor. Potential Involvement in Astrocyte Plasticity

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Abstract

GABA<sub>A</sub> receptor (GABA<sub>A</sub>R) expression level is inversely correlated with the proliferation rate of astrocytes after stroke or during malignancy of astrocytoma, leading to the hypothesis that GABA<sub>A</sub>R expression/activation may work as a cell proliferation repressor. A number of vasoactive peptides exhibit the potential to modulate astrocyte proliferation, and the question whether these mechanisms may imply alteration in GABA<sub>A</sub>R-mediated functions and/or plasma membrane densities is open. The peptide urotensin II (UII) activates a G protein-coupled receptor named UT, and mediates potent vasoconstriction or vasodilatation in mammalian vasculature. We have previously demonstrated that UII activates a PLC/PIPs/Ca<sup>2+</sup> transduction pathway, via both G<sub>a</sub> and G<sub>i/o</sub> proteins and stimulates astrocyte proliferation in culture. It was also shown that UT/G<sub>a</sub>/IP<sub>3</sub> coupling is regulated by the GABA<sub>A</sub>R in rat cultured astrocytes. Here we report that UT and GABA<sub>A</sub>R are co-expressed in cerebellar glial cells from rat brain slices, in human native astrocytes and in glioma cell line, and that UII inhibited the GABAergic activity in rat cultured astrocytes. In CHO cell line co-expressing human UT and combinations of GABA<sub>A</sub>R subunits, UII markedly depressed the GABA current (I<sub>GABA</sub>) in CHO and human astrocytes, and that UII partially reversed the GABA<sub>A</sub>R-mediated functional inhibition and endocytosis of GABA<sub>A</sub>R in CHO and human astrocytes, via its receptor C-terminus. This UII-induced disappearance of the repressor activity of GABA<sub>A</sub>R, may play a key role in the initiation of astrocyte proliferation.


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Introduction

Integrated brain function and dysfunction arise from the interactions between a network of multiple cell types including neurons, and the microvascular endothelial cells comprising the cerebral vasculature [1,2,3]. This micro-environment is a dynamic structure referred as neurovascular unit where polarized astrocytes have a pivotal role [4], rapidly transducing synaptic information [2,3,4,5]. In pathological conditions including stroke, the astroglial reactivity is characterized by proliferation, hypertrophy, process extension, increased synthesis of intermediate filament proteins and kinase activity, and expression of molecules involved in the regulation of cell proliferation and survival. These changes are associated with the development of glial scars and are a hallmark of the astrogliosis response to injury [6,7,8].

GABA<sub>A</sub> receptors (GABA<sub>A</sub>R) are believed to be pentameric heterooligomers mainly constructed from homologous subunit types α<sub>1–6</sub>, β<sub>1–3</sub>, γ<sub>1–3</sub>, δ and ε [9,10,11]. The GABA<sub>A</sub>R is expressed in neurons but also in glial cells in culture [12], brain slices [13], acutely isolated hippocampal slices [13], membrane fractions of the mature rodent brain [14] and also in vitro in healthy brain [15]. In pathological conditions, a significant decrease of benzodiazepine sites associated to the GABA<sub>A</sub>R has been demonstrated in patients with ischemic cerebrovascular [15,16,17], Parkinson [18] and Alzheimer [19,20] diseases. It was also observed a reduced chloride conductance [21], a decrease in receptor mediated inhibitory post-synaptic potentials [22] and a marked down-regulation of the GABA<sub>A</sub>R expression at the cell surface along with a fast time course [15,23,24]. In reactive and malignant astrocytes, mRNA levels of GABA<sub>A</sub>R have been shown to remain constant before diminution of functional GABA<sub>A</sub>R [15,25]. Thus, the disappearance of GABA<sub>A</sub>R expression is correlated with higher glial proliferation rate after stroke or during malignancy of astrocytoma [15,25,26], leading to the hypothesis that GABA<sub>A</sub>R expression/activation works as a repressor of cell proliferation. Investigations on alterations in GABA<sub>A</sub>R-mediated...
functions, receptor densities or modulation in astrocytes remain unchallenged. It has been demonstrated that simultaneous activation of different postsynaptic receptors induces cross-modulation of their activation properties and receptor membrane insertion/deletion. Thus, as many neurotransmitters and vasoactive peptides are released by endothelium and astrocytes, and their receptors are expressed by astrocytes, there is a potential for complex signaling within the neurovascular unit, involving receptor cross-talks.

Urotensin II (UII) and its paralog urotensin II-related peptide URP, are highly efficient vasoactive peptides, which share a fully conserved C-terminal cyclic CFWKYC core corresponding to the molecular pharmacophore [26,27,28]. The biological actions of UII and URP are mediated through activation of a G protein-coupled receptor named UT. It is now clearly established that activation of native UII receptors or UT-transfected cell lines is associated with an increase in polyphosphoinositol (PPIs) turnover promoting a cytosolic calcium concentration ([Ca\(^{2+}\)]\(_c\)) rise [29,30,31]. UII and UT are expressed in the mammalian cardiovascular system namely in the myocardium, vascular smooth muscle cells and endothelial cells [32,33,34,35], affecting cell proliferation [35,36] or neangiogenesis [37], stimulating collagen synthesis and cardiac hypertrophy [34].

In the brain, UII mRNA is particularly abundant in motoneurones of the medulla oblongata and spinal cord [38,39] while UT mRNA is widely expressed in various regions of the central nervous system including the olfactory system, hippocampus, amygdala, hypothalamus, or cerebellum [27]. However, immunohistochemical studies revealed that UT is expressed in astroglial processes in vivo [40] and in cultured rat cortical astrocytes [41]. In this cell type, we have previously demonstrated that UII activates a PLC/PPIs/Ca\(^{2+}\) transduction pathway, via both G\(_d\) and G\(_i/o\) proteins and stimulates cell proliferation [41,42]. Moreover, a functional interaction between GABA\(_{\alpha}R\) and UT suggested a cross-talk between these two receptors, involved in astrocyte activity [43]. In this study we demonstrate that activation of UT receptor induces a long-term inhibition of GABA\(_{\alpha}R\)-mediating chloride currents, a process potentially relevant for astrocyte proliferation.

## Results

GABA\(_{\alpha}R\) and UT functional coupling in rat cerebellar astrocytes

We and others have previously demonstrated that astrocytes express in vitro and in vivo UT mRNA and/or protein [38,40,41]. As shown in Figure 1Aa and 1b, UT is extensively expressed in astroglial processes, but also in the sparse mature granule cells present in co-cultures. In order to investigate a potential cross-talk between GABA\(_{\alpha}R\) and UT in both astrocytes and neurons, patch-clamp recording of the GABA\(_{\alpha}R\) agonist isoguvacine (Iso, \(10^{-4}\) M) responses was carried out on membrane potentials and currents of astrocytes in mono-culture and astrocytes or neurons in co-culture (Figure 1B and 1C). We established that flat and proliferating astrocytes in mono-culture (\(n = 31\)) (Figure 1B) were rarely responding (only 12% of cells are responding to Iso). In contrast, astrocytes cultured with cerebellar granule neurons present a slowly proliferating stellate astrocytic morphology (\(n = 78\)) (Figure 1B) and are more responding to the GABA\(_{\alpha}R\) agonist (48% of tested cells). In co-cultured astrocytes, local perfusion of rat UII (\(\mu UII, 10^{-7}\) M, 40 s) provoked a marked inhibition of the amplitude of the depolarization and chloride current (voltage clamp; \(-60\) mV) evoked by the GABA\(_{\alpha}R\) agonist isoguvacine by 24.2 ± 7.5% and 33 ± 6% (\(n = 5\)), respectively (Figure 1C). In contrast, \(\mu UII\) did not affect the Iso-evoked depolarization and current in cerebellar granule neurons (Figure 1C). In astrocytes, \(\mu UII\) induced a dose-dependent inhibition of the current with an EC\(_{50}\) value of 43.6 ± 23.7% M (Figure 1D). It can be noticed that in some cells, UII tested at 10\(^{-10}\) M, activated a small inward current. These data indicate that in astrocytes, unlike in neurons, \(\mu UII\) efficiently and markedly down regulated the GABAergic activity when UT and GABA\(_{\alpha}R\) are co-expressed.

In postnatal day 7 (P7) cerebellar slices, we investigated the topographic cellular and subcellular distribution of UT in the different layers. UT protein immunoreactivity was specifically distributed in particular zones of the cerebellar cortex, in the Purkinje cell layer (PCL), on fibers irradiating from the thin molecular cell layer (ML) toward the external granule cell layer (EGL), and on isolated cells residing in the internal granule cell layer (IGL) (Figure 2). In particular, UT receptor staining was present on NeuN-positive mature granule cell bodies (Figure 2A and 2A’) and also in Purkinje neuronal cell bodies and ramifications, as revealed by the strong immunofluorescence of UT in calbindin-positive Purkinje cell soma and dendrites (Figure 2B and 2B’). Doublecortin (DCX) is a distinctive marker of granule cells during the period of radial descent along the Bergmann glia into the cerebellar layers [44]. The DCX immunoreactivity appeared as a diffuse labeling in the IGL and densely in the ML, contiguously associated to UT-positive components but not co-localized (Figure 2C and 2C’). Moreover, UT was strongly co-localized with GFAP expressed by astrocytes of the white matter (not shown), and on glial cells (Figure 2D) including Bergmann cells [45]. At higher magnification, the double immunofluorescence analysis revealed that Bergmann cell bodies and fibers were surrounded by yellow co-localizing signals of UT varicosities and of GFAP-positive filaments (Figure 2D’). In addition, immunohistochemical analysis showed labeling of the \(\gamma_1\) subunit in Purkinje cells, and faint staining in fibers of the ML and in granule cell bodies of the IGL, co-localizing with UT (Figure 2E and 2E’). The \(\gamma_2\) subunit immunoreactivity was also mainly detected in Purkinje cells and fibers of the EGL (Figure 2F and 2F’). Thus, it appears that UT and GABA\(_{\alpha}R\) subunits are colocalized in cerebellar Purkinje and glial cell fibers in situ.

UT mediates inhibition of \(\gamma\)-composed GABA\(_{\alpha}R\) complexes

The molecular basis for the observed directional decrease in GABA\(_{\alpha}R\) function has been investigated in CHO co-expressing human UT and \(\beta\) and/or \(\gamma\) GABA\(_{\alpha}R\) subunits. GABA\(_{\alpha}R\) can be made from several different subunit families (\(\alpha_1-\alpha_6, \beta_1-\beta_3, \gamma_1-\gamma_9, \delta, \epsilon, \pi, \eta, \) and \(\theta\)), which come together in various combinations to form the pentameric receptor [46]. Most receptors are thought to contain \(\alpha, \beta\) and the third subunit type varying, being often the \(\gamma_2\). Expression of unitary subunits has produced conflicting results, but some subunits expressed alone [47,48] or as binary combinations, for instance \(\alpha_1\), \(\beta_3\) and \(\gamma_2\) appear to be able to produce GABA-gated ions channels [49,50,51]. In order to establish a direct functional link between GABA\(_{\alpha}R\) and UT, the effect of graded concentrations of \(\mu UII\) was studied on the current evoked by Iso on CHO-UT, expressing different subunit combinations of GABA\(_{\alpha}R\) (Figure 3A). It is observed that \(\mu UII\) induced inhibition of GABA\(_{\alpha}R\) complexes which systematically contained \(\gamma_1\) or \(\gamma_2\) subunit (Figure 3A and 3B), and the normalized data fit yielded various EC\(_{50}\) values and efficacies summarized in supplementary information (Table S1). In particular, \(\mu UII\) was less effective on the \(\alpha_2\), \(\beta_3\), and totally ineffective on the \(\alpha_2\), \(\beta_3\) and \(\gamma_2\) complexes (Figure 3A and 3B, Table S1).
Figure 1. UII-induced depression of GABAA\(_R\) in UT-expressing cerebellar astrocytes. (Aa, Ab) Double immunofluorescence labeling of UT (green) and the specific astrocyte marker GFAP (red, Aa), or the mature neuron marker NeuN (red, Ab) in astrocyte-neuron co-culture from P7 rat cerebellum. Astrocytes, recognized by strong GFAP staining show UT immunoreactivity (arrows), whereas few weaker UT-stained cells express NeuN (arrowheads), and were likely attributed to mature granule cells (arrowheads, Ab). Nuclei (blue) were counterstained with DAPI. Scale bars, 50 \(\mu\)m. (B) Phase contrast photomicrograph of astrocytes in mono-culture, or astrocytes and neurons in co-culture at 3 days in vitro. (C) Membrane depolarizations and currents evoked by the GABAA\(_R\) agonist isoguvacine (Iso, \(10^{-4}\) M, 2 s for membrane potential and 5 s for chloride current) in astrocytes and cerebellar granule neurons before, during rUII (\(10^{-7}\) M, 40 s) application and after 2-min washout. Right, normalized amplitudes deduced by the mean Iso-evoked depolarization or current obtained before rUII application. (D) Concentration-response relationship of Iso-evoked
As a control, the effect of the GABA\(_A\)R allosteric inverse modulator DMCM, \(\beta\)UII and other urotensinergic modulators, were tested on the \(\gamma_2\)\(\beta_3\)\(\alpha_2\) GABA\(_A\)R function in the absence of UT. Our data demonstrated that, as expected, DMCM induced inhibition of the current in most tested cells, and \(\beta\)UII and its paralog URP failed to affect the amplitude of the current from astrocytes yielding an EC\(_{50}\) value of 43.6\(\pm\)23.7 \(10^{-12}\) M. Data are mean \(\pm\) SEM of 4 to 6 cells. *, \(P<0.05\); ** \(P<0.01\) compared with the corresponding control Iso-evoked current. doi:10.1371/journal.pone.0036319.g001

Figure 2. Co-localization of UT with \(\gamma\) subunits in neuron and glial components in rat cerebellum. (A, A') Double-fluorescence staining for UT (green) and NeuN (red) showing the presence of UT in both mature (arrowhead, merge, A') and unidentified cells (arrows, merge, A') in the IGL. (B) Co-staining of UT and the marker of Purkinje cells, calbindin (red), in Purkinje cell soma and dendrites (arrowhead, B'). (C) Staining for UT and the marker of migrating neuroblasts doublecortin DCX (red) depicting a diffuse labeling in the ML. (C') UT immunopositive fibers contiguous to DCX-expressing migrating granule cells (merge, yellow, arrowhead). (D, D') Staining for UT and GFAP (red) in glial fibers (merge, yellow, arrowhead) of the ML. (E, F) Distribution of UT and the \(\gamma_1\) (E) and \(\gamma_2\) (F) GABA\(_A\)R subunits (red), in Purkinje cells (merge, arrowhead) and few extents of glia (merge, arrow) in the ML and IGL. Nuclei (blue) were counterstained with DAPI. Scale bars, 50 \(\mu\)m (A–F); 20 \(\mu\)m (A’–F’). EGL, external granule cell layer; IGL, internal granule cell layer; ML, molecular layer; PCL, Purkinje cell layer. (A’–F’) images of digitally zoomed regions corresponding to the white boxes in A–F. doi:10.1371/journal.pone.0036319.g002
Figure 3. Effect of hUII on different GABAAR subunit combinations. (A) Typical iso-evoked currents at the holding potential of ~60 mV, in the whole-cell configuration, on CHO stably expressing human UT (CHO-UT) and transiently transfected with cDNAs encoding α2β3γ2, α2β1γ1, α2β1γ1, α2β1γ1, β1γ2, β1γ2, or α2β1, subunits of the GABAAR. Iso (10⁻⁴ M) was repeatedly applied for 2 s at 2 min intervals and increasing concentrations of hUII (10⁻¹³ to 10⁻⁷ M) were bath perfused in the vicinity of cells. (B) Corresponding concentration-response curves for hUII on α2β1γ1 and α2β3γ2, α2β1γ1, and α2β1γ1, β1γ2 and β1γ2, α2β1, and α2β3, receptor subunits. Data are normalized to the control Iso response immediately prior to lower hUII concentration application. Data are mean ± SEM of 3 to 23 cells.

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Figure 4. Pharmacological characterization of the UT-mediated inhibition of the GABA_A R currents. (A) Whole-cell current response to Iso (10^{-2} M, 2 s) recorded in the absence or presence of the benzodiazepine site inverse agonist DMCM (10^{-5} M), hUII and URP (10^{-8} M, each), or UT antagonists [Orn^5]-URP and palosuran (10^{-6} M, each) in CHO expressing α_2, β_3 and γ_2 subunits. Below, summary of the various experimental conditions (n=3–18). (B) Comparison of the inhibitory effect of hUII and DMCM on CHO-UT-GABA_A R, as summarized in bar graphs (n=25). Bottom row, plot of the positive correlation (r^2 =0.8) of hUII-induced inhibition as function of the DMCM-evoked current decrease (n=28). (C) Comparison of the inhibitory effect of hUII and URP on CHO-UT-GABA_A R as summarized in bar graphs (n=12–54). (D, E) Effect of [Orn^5]-URP and palosuran in the absence or presence of hUII versus the effect of hUII alone. Right, summary of the various
(Figure 4A). The rapid development in recent years of several UT antagonists has led to the synthesis of [Orn$_5$]-URP which has been previously characterized in our laboratory [52,53] and palosuran, with one such high affinity toward human UT [54,55]. Thus, the specific rat UT antagonist [Orn$_5$]-URP (10$^{-6}$ M), and the primate UT specific antagonist palosuran (10$^{-8}$ M) did not modulate the Iso-evoked current.

To assess the role of $\gamma_2$ subunit into $\alpha_2$/$\beta$ binary complexes in the UII-induced GABA$_A$R regulation, we tested the effect of DMCM, as this allosteric modulator exhibits negative effect on GABA current in cells expressing $\gamma_2$ subunit [56,57] or null effect when $\gamma$ is not present [56]. We analyzed the effect of $\alpha$UII (10$^{-5}$ M) concomitantly to that of DMCM (10$^{-6}$ M) on CHO-UT co-expressing $\alpha_2$$\beta_3$$\gamma_2$ GABA$_A$R. The amplitude of the $\alpha$UII-induced inhibition was plotted as a function of the DMCM-induced inhibition of chloride currents recorded from twenty five CHO-UT-GABA$_A$R expressing cells. A linear regression analysis confirmed the significance of a correlation as represented on the scatter plot ($r^2 = 0.8$, $P < 0.001$) (Figure 4B), suggesting that $\gamma_2$ regulates the degree of UT-mediating inhibition of the GABAergic activity.

The effects of the different urotensinergic ligands were then tested on CHO-UT-GABA$_A$R composed of the $\gamma_2$ subunit. We found that $\alpha$UII and URP (10$^{-8}$ M, each) induced a marked current inhibition by 29.24±1.90% ($n = 54$) and 22.9±3.5% ($n = 12$), respectively, that persisted during washout for $\alpha$UII, but slightly recovered during washout for URP (Figure 4C). In order to examine whether UT antagonists might counteract the UII-induced decrease of the GABAergic activity, [Orn$_5$]-URP and palosuran were tested. [Orn$_5$]-URP (10$^{-6}$ M) significantly, but weakly inhibited the Iso-evoked current, blocked the effect of the acute application of $\alpha$UII, but failed to counteract the prolonged effect of the peptide on the same cell (Figure 4D). Surprisingly, palosuran mimicked the $\alpha$UII-induced inhibition of the Iso-evoked current, but abolished the irreversible action of $\alpha$UII during washout (Figure 4E). This indicates that [Orn$_5$]-URP or palosuran may keep their antagonist profile toward UT, but exhibit “agonistic activity” in regard to the GABA$_A$R function.

To test whether activation of the UT/[Ca$^{2+}$]$_i$, signaling pathway may be closely linked to the modulation of GABA$_A$R, changes in [Ca$^{2+}$]$_i$, evoked by $\alpha$UII, URP, [Orn$_5$]-URP and palosuran were measured by continuous Ca$^{2+}$ fluorescence imaging in CHO-UT. When applied to the bath solution, $\alpha$UII and URP evoked a significant and rapid amplitude of the baseline (Ca$^{2+}$), by 322% and 341%, respectively (Figure 5A and 5B). The effect of $\alpha$UII was irreversible, only partially recovering after 32-min washout (Figure 5A), as compared with URP whose effect totally recovered after 16-min washout (Figure 5B). It is also observed that [Orn$_5$]-URP (10$^{-6}$ M) behaved as a partial agonist (263% of [Ca$^{2+}$]), increase) but prevented the sustained effect of $\alpha$UII on [Ca$^{2+}$]. (Figure 5C). In contrast, palosuran (10$^{-8}$ M) failed to evoke a [Ca$^{2+}$], rise and completely blocked the $\alpha$UII-induced [Ca$^{2+}$], increase (Figure 5D).

Effect of UT on GABA$_A$R pharmacology and gating properties

According to the action of UT on GABA$_A$R, we asked the question whether the pharmacology and gating properties of GABA$_A$R were affected by UT. Thus, we found that the specific positive allosteric GABA$_A$R modulator pentobarbital (10$^{-5}$ M) directly activated a chloride current, and reversibly potentiated the Iso-evoked current by 196.28±12.31% (Figure S1A). In addition, SR95531 (10$^{-7}$ M) and picrotoxin (10$^{-7}$ M) induced attended current inhibition by 76.92±10.35%, and by 36.76±4.33%, respectively (Figure S1A). This indicates that pharmacological characteristics of the GABA$_A$R are not altered by the presence of UT. Next, to determine whether activation of UT might modify the conductance and selectivity of the GABA$_A$R-channel complex, the current-voltage (I-V) relationship was studied on CHO-UT co-expressing $\alpha_2$$\beta_3$$\gamma_2$ GABA$_A$R subunits. The voltage-dependence of the cell response to Iso (10$^{-3}$ M) was investigated in the absence or presence of $\alpha$UII and the amplitude of the current was measured at different holding potentials (Figure S1B). Local perfusion of $\alpha$UII (10$^{-8}$ M) decreased the slope of the I-V curve but did not significantly shift the chloride reversal potentials (EqCl$^-$, control, 4.84 mV; $\alpha$UII, 3.36 mV), closed to the theoretical EqCl$^-$ value calculated from the Nernst equation, based on the external and internal chloride concentrations used during recording (see Materials and Methods section). It is observed that $\alpha$UII-induced inhibition of the Iso-evoked current recorded at −60 mV (26.93±6.35%) and +60 mV (37.06±11.25%) was very similar and did not significantly depend on the holding potential (Figure S1B).

The subunit composition determines the GABA sensitivity and the pharmacological properties of the GABA$_A$R [9] as well as the time course of the GABA response referred as desensitization and deactivation of chloride current [58,59,60]. To clearly assess the mechanism of UT-mediating inhibitions of the GABA current, $\alpha$UII was applied on CHO-UT-GABA$_A$R on the fast component of current desensitization. As shown in Figure 6, the current evoked by Iso showed a slow decay during continuous agonist ejection as observed by an apparent desensitization of 59% in control, and 77% under $\alpha$UII perfusion. We then followed the time-dependent recovery from desensitization in the absence or presence of $\alpha$UII. Recovering of 50% receptors from this long exposure occurred with mean time constants of 16.2 s in control and 82.2 s in the presence of $\alpha$UII, demonstrating that the progressive recovery is delayed in the presence of the peptide (Figure 6). Thus, when coexpressed with UT, the pharmacological profile of GABA$_A$R is not altered, but UT rather affects macroscopic $\alpha_2$$\beta_3$$\gamma_2$R receptor current desensitization, and markedly slows the recovering process. Since recovery from desensitization does not involved membrane voltage [60], it is suggested that UT-mediating GABA$_A$R desensitization is mainly due to conformational changes of the ligand-bound receptor chloride channel, paralleled to a mechanism known to develop from the closed but fully bound conductive state of the receptor [61].

Mechanisms promoting fast short-term and long-term UT-induced GABA$_A$R current inhibition

In CHO-UT-GABA$_A$R, the Iso-activated currents were measured during a 28-min recording period from the initial application of the GABA$_A$R agonist. A 1-min application of $\alpha$UII (10$^{-6}$ M) provoked a fast and significant decrease of the current (23.44±2.47%, $n = 10$) followed by a progressive run-down, reaching 84.61±5.92% ($n = 4$) inhibition after 24-min washout (Figure 7A). The second large intracellular loop of several GABA$_A$R subunits possesses numerous consensus phosphorylation sites [62] and effective phosphorylation mechanisms have been shown to be involved in either short- or long-term regulation of
Figure 5. Role of specific UT ligands on cytosolic calcium in CHO-UT. (A, B) hUII (A) or URP (B) (10^{-8} M, each) provoked a robust increase of \([\text{Ca}^{2+}]_c\), which remained stable (A) or recovered to the basal line level (B) during washout. (C, D) Effect of the UT antagonists [Orn5]-URP (10^{-6} M, C) or palosuran (10^{-8} M, D), before and during hUII application. Right, bar graphs represent the percent increase of the [\text{Ca}^{2+}]_c during drug perfusion or during the washout period. Percent values were obtained by normalizing signals evoked during and after treatments to the value measured before
inhibitory synaptic transmission [63,64]. To further explore the mechanisms sustaining the hUII-induced current long-term depression, we hypothesized that various UT-couplings to G proteins, relay [Ca^{2+}], rise resulting in kinase or phosphatase activation, likely responsible for changes in GABA_{A}R subunits phosphorylation state. G-protein activation requires the exchange of bound GDP (resting state) with GTP (activated state). This is a common and necessary step of all G-protein mediated actions and is independent of G-protein type or the second messenger system involved. Therefore, blockade of this exchange will result in inability of the ligand-bound receptor to exert its action. Thus, G protein specific blockade with GDP_{S} (10^{-4} M) did not significantly prevent the fast and long-term inhibition of the current induced by hUII (Figure 7B). Intracellular dialysis with a cocktail of kinase and phosphatase inhibitors (KIC; phosphatase inhibitory cocktail, quercetin 10^{-5} M and staurosporine 10^{-5} M), failed to alter the fast hUII inhibitory effect but attenuated the rundown phenomenon (Figure 7C). It is also observed that hUII reduced the peak current amplitude after 5-s perfusion, but evoked a peak [Ca^{2+}]c increase only after a 10-s delay (Figure 7D). Consistent with this observation, intracellular BAPTA (10^{-4} M) dialysis reduced the long-term current inhibition by only 39.29 ± 10.16% (n = 9) (Figure 7E). Together, these observations tend to show that G proteins do not transduce UT-induced current inhibition and that calcium transient and phosphorylation mechanisms do not play a promoting role, but participate in the run-down of the GABA_{A}R current. To test a hypothetical role of UII in the dynamin-dependent GABA_{A}R endocytosis, the dynamin inhibitory peptide DIP, which competitively blocks binding of dynamin to amphiphysin [64], has been introduced in the intrapipette solution. As shown in Figure 7F, when cells were dialyzed with DIP (10^{-5} M), hUII retained its ability to induce a fast and highly reversible inhibition of the Iso-evoked current, but failed to reduce the current amplitude with time recording.

Inhibition of the GABAergic activity involves GABA_{A}R internalization and requires specific UT receptor domains in CHO and human astrocytes

To assess whether the C-terminus fragment of UT, corresponding to the cytoplasmic C-tail of the receptor contributes to the UII-induced run-down of the GABA_{A}R activity, we constructed four truncated mutants in which the last 19 (UTHA_{370}), 38 (UTHA_{351}), 57 (UTHA_{332}) or 70 (UTHA_{319}) residues were removed (Figure 8A). They are all present at the plasma membrane (Figure S2) and functionally expressed except the UTHA_{319} truncated form of UT (Figure S3). When UTHA_{370} is coexpressed with GABA_{A}R, the hUII-induced fast short-term inhibition was totally abolished, whereas the long-term inhibition was delayed, and significantly altered (Figure 8B). In the presence of shorter truncated forms, hUII totally failed to alter the GABA_{A}R activity, suggesting that the most distal C-terminus part of UT is involved in the functional cross-talk with GABA_{A}R (Figure 8B). Moreover, the peptidomi-

![Figure 6. UII-induced fast current inhibition and GABA_{A}R desensitization.](https://example.com/figure6.png)

(A, B) Examples of currents recorded from CHO-UT-GABA_{A}R during a long desensitizing pulse (25 s) of Iso (10^{-4} M), in the absence (black line) or presence (green line) of hUII (10^{-8} M, 1 min). (A) Exponential fit to the desensitizing current phases were shown overlaid on the currents. Bar graphs corresponding to the average desensitization constant parameter t in the absence (t_{Ctrl}) or presence (t_{hUII}) of hUII (n = 5). (B) Prolonged Iso (30 s) application eliciting current desensitization followed by a time course of the recovery from desensitization, in the absence (control) or presence of hUII. Graph represents the Iso-evoked current expressed as a fraction of the peak control current induced by the long Iso application to the current amplitude elicited by each short pulse, and plotted against interpulse intervals. Data are mean ± SEM from 3 to 8 cells. *, P < 0.05; ** P < 0.01; *** P < 0.001 compared with the corresponding control Iso-evoked current. doi:10.1371/journal.pone.0036319.g006

Figure 6. UII-induced fast current inhibition and GABA_{A}R desensitization. (A, B) Examples of currents recorded from CHO-UT-GABA_{A}R during a long desensitizing pulse (25 s) of Iso (10^{-4} M), in the absence (black line) or presence (green line) of hUII (10^{-8} M, 1 min). (A) Exponential fit to the desensitizing current phases were shown overlaid on the currents. Bar graphs corresponding to the average desensitization constant parameter t in the absence (t_{Ctrl}) or presence (t_{hUII}) of hUII (n = 5). (B) Prolonged Iso (30 s) application eliciting current desensitization followed by a time course of the recovery from desensitization, in the absence (control) or presence of hUII. Graph represents the Iso-evoked current expressed as a fraction of the peak control current induced by the long Iso application to the current amplitude elicited by each short pulse, and plotted against interpulse intervals. Data are mean ± SEM from 3 to 8 cells. *, P < 0.05; ** P < 0.01; *** P < 0.001 compared with the corresponding control Iso-evoked current. doi:10.1371/journal.pone.0036319.g006
Figure 7. Intracellular mechanisms of UT-triggering GABA_{A}R inhibition. (A) Traces of Iso (10^{-4} M, 2 s)-evoked current amplitude time-course on CHO-UT-GABA_{A}R, in control (above row) or during a 1-min application of hUII (10^{-8} M, bottom row). Corresponding average time course of the Iso-evoked current, in control or during and after hUII application. (B, C) Current traces before (1), during (2) a 1-min hUII application and after 20-min washout (3), in the absence or presence of GDPxS (B, 10^{-4} M, 15 min of dialysis) or the cocktail of kinase and phosphatase inhibitors (C, KIC, 

D}

E

F

G

hUII − BAPTA

hUII + BAPTA

hUII − DIP

hUII + DIP

Current (% of control)

Recording time (min)

Current (pA)

Ratio (G340/G380 nm)

GDPxS

KIC

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metric UT^{myc-319–389} (Figure 8A) completely inhibited the fast and long-term effects of UII on the GABA-evoked current (Figure 8C). Collectively, these results indicate that the C-terminus of UT relays the inhibition of the GABA_{AR} function, and may counteract a potential hUII-induced internalization process.

Our data thus suggest that UT activation likely regulates GABA_{AR} endocytosis. We first established the subcellular localization of both UT and GABA_{AR} in cultured CHO transiently transfected with cDNAs encoding recombinant human UT and the β_{2}γ_{2}-tagged (β_{2}γ_{2}HA) GABA_{AR} subunits and then, internalization of GABA_{AR} was followed by labeling the surface receptors with antibodies directed against the β_{2}γ_{2}HA co-expressed with β_{2}γ_{2} GABA_{AR} subunits and/or UT. In the absence of ligand, the immunoreactivity for β_{2}γ_{2} exhibited membrane localization (green) as enlightened by the intensity profiles (Figure 9Aa). In contrast, treatment with hUII (10^{-8} M), Iso (10^{-8} M) or the two agonists, drastically promoted GABA_{AR} endocytosis by 40.29±4.14%, 39.31±2.84% and 34.71±3.19%, respectively (Figure 9B), as seen by the increase of red punctuates in the cell soma (Figure 9Ab-8Ad). When GABA_{AR} was expressed alone, hUII failed to induce GABA_{AR} internalization whereas Iso or Iso combined with hUII remained able to provoke GABA_{AR} removal from the plasma membrane (Figure 9B).

In order to confirm the involvement of UII/UT in the internalization process of GABA_{AR} in CHO, we investigated the plasma membrane expression of the β_{2}γ_{2}HA GABA_{AR} subunit, as well as UT^{myc-319–389} after exposure to hUII by measuring the amount of surface immunolabelled receptors by ELISA. Typical bioluminescence and fluorescence values obtained from CHO expressing either UT^{myc-319–389} or UT^{myc-319–389}β_{2}γ_{2}HA, and UT^{myc-319–389}β_{2}γ_{2}HA or UT^{myc-319–389}β_{2}γ_{2}HA were, respectively, shown in Figure 10A. In CHO-UT^{myc-319–389}β_{2}γ_{2}HA, a 30 min treatment with hUII (10^{-8} M) led to approximately 40% loss of β_{2}γ_{2}HA subunit from the cell surface, without modifying cell membrane amounts of UT^{myc-319–389} (Figure 10B). In CHO-UT^{myc-319–389}β_{2}γ_{2}HA cotransfected with the cDNA encoding the UT^{myc-319–389}YFP peptidomimetric fragment, hUII incubation failed to remove the β_{2}γ_{2}HA subunit from the plasma membrane (Figure 10B), establishing that the C-terminus part of UT played a major role in the UII-induced GABA_{AR} internalization in a recombimant system. The physiological relevance of such mechanism was thus assessed in native human astrocytes and in the human glioma U87 cell line. Flow cytometry analysis showed that normal and tumoral glial cells in culture expressed β_{3} subunit (Figure 11A) and UT (Figure 11B), and that one population (around 8%) of astrocytes and U87 exhibited both receptors at the plasma membrane (Figure 11). hUII (10^{-8} M, 30 min) provoked more than 65% GABA_{AR} disappearance from the surface, whereas UT internalized in native astrocytes (Figure 11A) but was still present at the cell surface in U87 (Figure 11B). We then examined the cell surface stability of GABA_{AR} following UT activation in the absence or presence of the coexpressed UT^{myc-319–389}YFP on U87 glioma cell line. ELISA-based assay revealed that the UT^{myc-319–389} fragment reduced the β_{3}-associated GABA_{AR} subunit internalization from 20% to approximately 10% (Figure 11C), with an efficacy of transfection of around 32%, as controlled by flow cytometry (data not shown). As already shown in CHO, UT levels at the cell surface remained unchanged after exposition to hUII, in the absence or presence of the UT C-terminus (Figure 11C).

Discussion

The question of the regulation of the GABA_{AR} plasticity and cross-modulation in extrasynaptic glial cells by neurovascular factors, and its functional consequence were not explored. In rodent brain, mRNA encoding the vasoactive receptor UT are detected in the olfactory system, hippocampus, amygdala, tegmental nuclei, or cerebellum [27,65,66] and UII binding sites are restricted to few areas including the cerebellar cortex [39,67]. We and others have also shown that UT receptor expression may be specific to glial cells in vivo [40], and to astrocytes in culture [41], as well as to brain microvascular entities (unpublished data).

In the present study, to gain insight into the regulation of the GABAergic activity by the vasoactive peptide UII in astrocytes, the cellular and subcellular distribution and feature of the UT receptor, have been investigated in the cerebellum in situ, and in astrocyte-neuron co-cultures. We showed intense UT labeling in Purkinje cell bodies and ramifications as well as in glial Bergmann-GAPF positive long processes where it co-localizes with γ_{1} and, to a lesser extent, with γ_{2} subunits of the GABA_{AR}, and in astrocytes co-cultured with granule neurons. Together, this corroborates previous studies establishing expression of β_{3}, γ_{1} and γ_{2} subunits in the PCL [68,69], and of γ_{1} mRNA in Bergmann glia [70,71]. We confirmed the coexpression of GABA_{AR} subunits with UT in native human astrocytes and in the U87 glioma cell line. It is interesting to note that a majority of glial cells express at least the β_{3} GABA_{AR} subunit and also UT in the cytosolic compartments, but that only around 10% of cells corresponding to a common subpopulation, show these receptors at the plasma membrane. This is in a good agreement with the 12% responding rat cultured astrocytes to isoguvacine in mono-culture. In fact, the existence of two populations of astrocytes, exhibiting depolarized membrane potentials (around −30 mV) in a majority of cells and hyperpolarized membrane potentials (around −80 mV) in a minority (data not shown) was observed in our study, as already shown in cultured astrocytes [72], independent on patch-clamp recording conditions. Here, we determined that this hyperpolarized subpopulation represents astrocytes specifically responding to the GABA_{AR} activation. Since it was suggested that GABA acts as an anti-proliferating neurotransmitter in ventricular and subventricular zones [73] and in cortical progenitor cells [74] and that down regulation of functional GABA_{AR} is correlated with the proliferation rate of reactive or malignant astrocytes [13,25], we propose that rat and human cell subpopulation expressing GABA_{AR} likely correspond to quiescent astrocytes in culture.

These colocated features of UT and GABA_{AR} in astrocytes prompted us to investigate a potential functional cross regulation between the two receptors, likely involved in astrocyte plasticity. It has been shown that astrocytic GABA response is specific of early
We demonstrate that UII down regulates the Iso-evoked depolarization and chloride current amplitudes recorded from astrocytes co-cultured with granule neurons. This UII-induced GABAAR current inhibition is shown to be a very high affinity process, specific of astrocytes, which hardly recovered during washout. In CHO co-expressing human UT and αβ and/or γ GABAAR subunits, we found that UII was i) very potent on β1/γ1/α/γ2L GABAAR subunit complexes, ii) less potent on αβ1/γ1/LGABAAR complexes and iii) inactive on αβ binary complexes. These results

Figure 8. Receptor sequences involved in UT regulation of the GABAAR activity. (A) Schematic diagrams mixed with sequence alignments of the HA epitope-tagged human UT, C-terminus truncated UT^HA370, UT^HA351, UT^HA332, UT^HA319 mutants, and peptidomimetics corresponding to the entire C-terminus cytosolic fragment of UT (UT^c-myc319–389). (B and C) Traces of Iso (10^{-4} M, 2 s)-evoked current before (1), during (2) a 1-min UII (10^{-8} M) application and after 22-min washout (3). (B) Currents recorded from CHO coexpressing GABAAR and UT^HA (Control), UT^HA370, UT^HA351, UT^HA332 or UT^HA319. Corresponding average time course of the currents, in the absence or presence of UT truncated mutants. (C) Current traces recorded from CHO-UT-GABAAR, in the absence or presence of UT^c-myc319–389. Corresponding average time course of the Iso-evoked current, in the absence or presence of UT^c-myc319–389. In B, significance was only annotated above the time course graph during UII perfusion and after 18-min washout, for clarity. Data are mean ± SEM from 3 to 13 cells. ns, non significant; *, P<0.05; ** P<0.01 compared with the corresponding control Iso-evoked current.

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Figure 9. UT activation mediating GABA\(_{\alpha}R\) internalization. (Aa–Ad) CHO-UT transiently transfected with cDNA encoding \(\alpha_2\beta_3\gamma_2\) GABA\(_{\alpha}R\) subunits. Internalization was controlled through translocation of \(\beta_3\) subunit (red) in control (Aa) or after 60 min of hUII (10 \(^{-8}\) M, Ab), Iso (10 \(^{-4}\) M, Ac) or hUII-Iso (Ad) incubation. Fluorescence intensity plots of green and red fluorescences corresponding to the localization of GABA\(_{\alpha}R\) (\(\beta_3\)) at the plasma membrane and in the cytosol, respectively, across the regions delimited by the white line scans. A.u., arbitrary unit; scale bars, 25 \(\mu\)m. (B) Bar graphs of the fraction of fluorescence at the plasma membrane on CHOT-UT-GABA\(_{\alpha}R\) or CHO-GABA\(_{\alpha}R\) in the different conditions. Each bar corresponds to mean ± SEM percent obtained from 3 to 18 cells. ns, non significant; ***, \(P<0.001\) versus control in CHO-UT-GABA\(_{\alpha}R\); ####, \(P<0.001\) versus control in CHO-GABA\(_{\alpha}R\).

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thus establish that UII exhibits a very high affinity directional inhibition toward the GABA$_{A}$R specifically composed of the $\gamma_{2}$ subunit. This observation can be paralleled to the high affinity binding sites for UII determined on recombinant UT-expressing cells [76,77,78,79], and also on astrocytes [41]. The time course of the UT-evoked current inhibition can be distinguished by two phases, i.e. a short-term decrease detected immediately after and during UII administration, followed by a progressive run-down of the UII effect.

Figure 10. UII-induced GABA$_{A}$R loss from the plasma membrane through the C-terminus fragment of UT in CHO. The effect of hUII on the proportion of GABA$_{A}$R and UT at the cell surface of CHO was assessed by ELISA. (A) CHO transiently transfected with cDNA encoding UT$^{\text{c-myc}}$ and $\alpha_{2}\beta_{3}$ or $\alpha_{2}\beta_{3}\gamma_{2}^{\text{HA}}$ GABA$_{A}$R subunits (left), or UT$^{\text{c-myc}}$, and $\alpha_{2}\beta_{3}\gamma_{2}^{\text{HA}}$ GABA$_{A}$R subunits cotransfected with the cDNA encoding UT$_{319-389}$YFP (right). Background bioluminescence (left) and fluorescence (right) were measured after anti-HA antibody and colorimetric alkaline phosphatase substrate incubation, in the absence or presence of 30 min of hUII (10$^{-8}$ M, left), or directly on a fluorescent plate reader (right). (B) CHO transiently transfected with cDNA encoding UT$^{\text{c-myc}}$ and $\alpha_{2}\beta_{3}$ or $\alpha_{2}\beta_{3}\gamma_{2}^{\text{HA}}$ GABA$_{A}$R subunits (left), or cotransfected with the cDNA encoding UT$_{319-389}$YFP, and immunodetected with anti-HA (left) or anti-c-myc (right) antibodies. Percentage of cell surface $\gamma_{2}^{\text{HA}}$ GABA$_{A}$R subunit (left) or UT$^{\text{c-myc}}$ (right) are represented as the proportion of receptor at the plasma membrane (non permeabilized cells) to the total expressed receptor (permeabilized cells). One hundred percent correspond to values in the absence of 30 min treatment with hUII (10$^{-8}$ M, 37 °C). Each bar corresponds to mean ± SEM percent obtained from 5 to 7 independent experiments, in triplicates. ns, non significant; *, $P<0.05$; ***, $P<0.001$.

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Figure 11. UII-evoked GABAAR internalization in native human astrocytes and glioma. (A, B) Flow cytometric analysis of the $\beta_3$ GABAAR subunit and UT expression in native human astrocytes (A) and human U87 glioma cell line (B). Cells were stained with the anti-human $\beta_3$ subunit or anti-human UT in permeabilized or non-permeabilized conditions (membrane receptor). The black lines depict results from control staining with only secondary antibodies. The $\beta_3$ GABAAR subunit or UT cell surface expression was evaluated in the absence or presence of hUII (10$^{-8}$ M, 30 min) by flow cytometry. Data obtained in A and B illustrate two representative experiments showing $\beta_3$ (magenta line) and UT (yellow line) mean fluorescence in the cytosol and at the plasma membrane of a minority of non-permeabilized human astrocytes (A) or U87 (B) in culture. The exposure to hUII induced internalization of $\beta_3$ in both cell types and of UT only in U87 glioma. (C) U87 glioma cell line expressing UT and GABAAR composed of $\beta_3$ subunit, and transfected with the cDNA encoding UT$^{319-389}$YFP, and immunodetected with anti-$\beta_3$ (left) or anti-UT (right) antibodies. Percentage of
the current, leading to about 80% GABA\textsubscript{A}R current disappearance. However, URP which exhibits the same conserved biologically active cyclic sequence than UII, triggers a reversible inhibition. This is in accordance with the UII-induced long-lasting in the one hand, and the URP-evoked transient, on the other hand, increase in [Ca\textsuperscript{2+}]\textsubscript{c} in CHO-UT. The specific long-lasting phenomenon might be attributable to the slow dissociation rate of UII, as already described for rat and human UII on UT transfected cells, skeletal muscle myoblasts and astrocytes [41,54,79]. This could account for the sustained and washout-resistant contractile responses induced by UII on primate arteries, [Ca\textsuperscript{2+}]\textsubscript{c} increase in rat cortical astrocytes [42], and GABA\textsubscript{A}R current inhibition in our native and recombinant systems. We propose a mechanism whereby UII interacts reversibly with the classical binding site, but also with a secondary exosite in a wash resistant manner, resulting in persistent activation of UT and consequently, in a long-term inhibition of the GABA\textsubscript{A}R. Such process has already been demonstrated in vivo and cell culture, for exogenous agonists of M1 muscarinic and \(\beta_2\) adrenergic receptors [80,81]. Together, our previous work suggesting that GABA negatively controls UT-mediating signaling transduction in astrocytes [43], corroborated by the effect of benzodiazepines on UII-induced neurotransmitter release [82], supports at most the existence of a negative cross-talk between UT and the closely associated GABA\textsubscript{A}R, leading to a high affinity functional receptor complex in astrocytes. This functional complex may exhibit new pharmacological profile. Accordingly, we demonstrated that [Orn\textsuperscript{5}]-URP acts as a partial agonist and competitive antagonist on both GABA currents and [Ca\textsuperscript{2+}]\textsubscript{c} increase in CHO-UT. Surprisingly, palosuran as a specific primate UT antagonist [55], counteracts the UII-evoked [Ca\textsuperscript{2+}]\textsubscript{c}, increase but mimics the effect of UII by inhibiting the chloride current. Thus, palosuran behaving as an antagonist of the UT-mediating [Ca\textsuperscript{2+}]\textsubscript{c} transduction signaling can be considered as a partial UT “agonist” toward the GABA\textsubscript{A}R effector pathway, then suggesting a different UT pharmacology when co-expressed with GABA\textsubscript{A}R.

Here we found that initiation of UT and GABA\textsubscript{A}R functional interaction is independent on G protein, calcium and phosphorylation mechanisms, but that UII-induced current run-down partially requires calcium and kinase/phosphatase activities. In this context, the Ca\textsuperscript{2+}/calmodulin requirement for membrane fusion in endocytic pathways [83], as well as the clathrin-mediated internalization depending on kinase and phosphatase activities[84,85,86], support here a possible role of UT in a delayed calcium/kinase dependent GABA\textsubscript{A}R endocytosis. GABA\textsubscript{A}R internalization is primarily thought to occur via a clathrin- and dynamin-dependent mechanism [87]. Here DIP, known to block endocytosis by disrupting the interaction between dynamin and amphiphysin, did not interfere with the UII-induced fast short-term but totally abolished the long-term inhibition of the current, supporting a role of UT in the dynamin-dependent GABA\textsubscript{A}R internalization. We then show that the GABA\textsubscript{A}R agonist triggered removal of \(\beta_3\) or \(\gamma_2\) GABA\textsubscript{A}R subunit from CHO plasma membrane, or from native human astrocyte and glioma cell surface. Together, the interesting point resides in the ability of the UII/UT system in the promotion of the marked GABA\textsubscript{A}R internalization in the absence of co-activation of GABA\textsubscript{A}R. In addition, 30 min incubation with UII failed to internalize UT in CHO and U87, but led to UT loss from human astrocyte surface, a discrepancy unexplained but needing further investigations. Thus, a constitutive tight promiscuity between UT and GABA\textsubscript{A}R might be responsible for the high affinity effect of UII on GABA\textsubscript{A}R disappearance from the plasma membrane.

Here we produced and expressed truncated UT receptors in order to identify which specific receptor determinants are involved in the GABA\textsubscript{A}R modulation. The deletion of up to 57 residues of the UT C-terminus, did not affect the expression and ability of truncated mutants to stimulate the [Ca\textsuperscript{2+}]\textsubscript{c}, as already observed for rat UT truncated mutants [88]. We demonstrate that the entire UT C-terminus totally abolished the UII-mediating current inhibition and that the most distal part likely relays the fast and long-term inhibitions of the GABA\textsubscript{A}R function. In rat, the last 19 residues of the C-terminus contain motifs that are not crucial for UT internalization [88]. In fact, the serine cluster localized upstream rather displays consensus motifs for PKC and casein kinase I important for rat UT internalization. In humans, the distal UT C-terminus (351–389) exhibits serine residues and a combination of two polyproline motifs (Figure 8A), the last one possibly interacts with SH3 domain proteins [89]. Our present data in CHO co-expressing human UT and \(\alpha_2\beta_3\) and \(\gamma_2\) GABA\textsubscript{A}R and in U87 transfected with the cDNA encoding the UT\textsubscript{319–389} peptidomimetic, indicate that the UII-induced internalization of GABA\textsubscript{A}R formed from \(\gamma_2\) or \(\beta_3\) subunit, requires at least in part, the C-terminus fragment of UT. Therefore, the question of whether UT and GABA\textsubscript{A}R physically associate directly or whether SH3 proteins relayed GABA\textsubscript{A}R down regulation, has to be elucidated. Together, the functional cross-modulation between UT and GABA\textsubscript{A}R is mediated through the most distal part of the UT C-terminus, which would directly interact with \(\gamma\) subunits, or recruit intermediate proteins implicated in GABA\textsubscript{A}R inhibitory transactivation.

Therefore, our observations suggest a model in which UT and GABA\textsubscript{A}R are closely associated to depress the GABAergic activity (Figure 12). When UT and \(\alpha_2\beta_3\) GABA\textsubscript{A}R subunits are co-expressed, as in native and tumoral glial cells, U87 efficiently activates its receptor, leading to a fast short-term decrease of the chloride current, independently of G proteins, calcium, phosphorylation and endocytosis processes, and involving the last 19 amino acids of the UT C-terminus. During washout, a long-term inhibition referred as run-down, develops via a dynamin-dependent internalization requiring the 351–370 sequence of UT, and calcium- and phosphorylation-dependent endocytic mechanisms. This directional cross-talk between UT and the GABA\textsubscript{A}R leads to the extinction of the GABA\textsubscript{A}R expression at the plasma membrane that would play a key role in the induction of cell proliferation (Figure 12).

Materials and Methods

Animals

Wistar rats (Depré, Saint-Doulchard, France) were kept in a temperature-controlled room (21 ± 1°C), under an established photoperiod (lights on 07.00–19.00 h) with free access to food and tap water. The work described in this article was carried out in accordance with the Directive 2010/63/EU of the European parliament and of the council of 22\textsuperscript{nd} September 2010 on the
Figure 12. Schematic model depicting the mechanism of UT-mediated GABA\(\alpha\)R down-regulation. UTI efficiently activates the G protein-coupled receptor UT, leading to a fast short-term decrease of the chloride current not sustained by G proteins, calcium, phosphorylation and endocytosis processes. This rapid effect involves the distal 19 C-terminal amino acids of UT and the presence of \(\gamma\) subunits within of the GABA\(\alpha\)R complex (1). During the washout period, a long-term inhibition develops via a dynamin-, calcium- and phosphorylation-dependent endocytic mechanisms, requiring at least in part the 351–370 sequence of UT and GABA\(\alpha\)R \(\gamma\) subunits (2). It is hypothesized that the directional cross-talk between UT and GABA\(\alpha\)R, and the extinction of the latter at the plasma membrane, may relay transition from quiescent to proliferant astrocytes.

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For astrocyte-neuron co-culture, granule cell cultures were prepared from cerebella of P7 Wistar rats as described previously [91]. Isolated cells were plated on 14-mm culture dishes coated with poly-L-lysine (5 mM) at a density of \(1.5 \times 10^6\) cells/dish and incubated at 37°C in a humidified atmosphere (5% CO\(_2\)) for 1 to 10 days before use. Culture medium consisted of DMEM/Ham’s F12 (75%/25%) supplemented with 10% FBS, 2 mM glutamine, 5 \(\mu\)g/ml insulin, 25 mM KCl and 1% of antibiotic-antimycotic solution. Co-cultures are obtained by seeding granule cells (\(1.5 \times 10^6\) cells/ml) on cerebellar astrocytes plated in 24-well plates after 12 hours. Co-cultures are maintained in the specific medium for neuron and incubated at 37°C in a humidified atmosphere (5% CO\(_2\)) for several days.

Human native astrocytes and glioma cell line

The human cell line from glioblastoma U87 was obtained from the American Type Culture Collection (LGC Standards, Mol-sheim, France). U87 cells were maintained in DMEM containing 10% FBS and 1% sodium pyruvate. NHA-Astrocytes (Lonza, Walkersville, MD, USA) were cultivated in DMEM culture medium supplemented with 2 mM glutamine, 5 mM HEPES, 1% non essential amino acids, 1% sodium pyruvate, B27, 25 ng/ml EGF, 1% of the antibiotic-antimycotic solution and 10% FBS. All cells were incubated at 37°C in a humidified atmosphere containing 5% CO2. Culture media were replaced every three days.

CHO recombinant cell line and plasmid transfection

CHO-CHO-K1 cell lines were obtained from American Type Culture Collection (Manassas, VA, USA). The human UT stable CHO (CHO-UT) was generously provided by Dr Christophe Dubessy (Inserm, Rouen University, France) et generated by CHO electroporation with 20 \(\mu\)g pIRES-neo2-UT DNA and 500 \(\mu\)g
sterile sonicated salmon sperm DNA using the EasyJet One electroporation system (Equibio, Angleur, Belgium), followed by repetitive rounds of limiting dilution of cells in G-418 (400 µg/ml) for selection. Normal or stable CHO was grown in Ham-F12 medium supplemented with 10% FBS, 1% antibiotic-antimycotic solution and 2 mM glutamine, at 37°C in a humidified incubator with an atmosphere of 5% CO2. The stable CHO-UT medium was supplemented with the antibiotic G-418 (40 µg/ml).

For transfection, cells were trypsinized (0.05%), triturated in Ham-F12 containing 10% FBS media, pelleted by centrifugation, resuspended in 100 µl solution V for nucleofection by an Amaza Nucleofector Device (Koln, Germany; set to program U016). Experiments were performed on normal CHO or on the stable CHO-UT cells and transiently transfected with combinations of cDNA encoding α2β2γ2, α2β3γ1, α2β2γ2, α2β2γ1, α2β2γ5 (4 µg of cDNA total/transfection) GABAαR subunits. Cells were seeded on 14-mm poly-L-lysine-coated glass bottom insert dishes at 5 x 10^5 cells in a volume of 0.5 ml/dish. Cells were incubated overnight at 37°C in a humidified incubator (5% CO2) during 16 h before electrophysiological or immunocytochemical experiments.

Recombinant receptors

For UT epitope-tagged with HA (UTHA), or c-myc (UT-cmyc), human UT receptor cDNA inserted into pcDNA3.1 (Ressource Center, MI, USA) was amplified and PCR products were subcloned using the EcoRI and Xhol sites of pCMV-HA or pCMV-c-myc. Mutant UT receptor cDNAs were constructed by oligonucleotide-directed mutagenesis (Expand High Fidelity PCR System; Roche) using the human UT receptor cDNA inserted into pcDNA3.1 as a template. Two sets of forward and reverse oligonucleotides were used (Table S2) to introduce stop codons in frame of Leu370, Phe351, Gly332 and Thr319 (to generate pcDNA3.1 as a template. Two sets of forward and reverse System; Roche) using the human UT receptor cDNA inserted into oligonucleotide-directed mutagenesis (Expand High Fidelity PCR pCMV-c-myc. Mutant UT receptor cDNAs were constructed by enzymes.

16 h before electrophysiological or immunocytochemical experiments.

The 5' and 3' oligonucleotides incorporated Sall and NotI or EcoRI and BamHI sites, respectively, to facilitate subcloning into pCMV-c-myc or pEYFP-N1 (BD Biosciences Clontech, Mississauga, ON, Canada), and incorporated initiation and stop codons where appropriate (Table S2).

In order to generate mini-peptides corresponding to the C-terminus of UT, cDNA encoding the UT-319-319 or UT-319-389 YFP fragment was amplified by PCR. The 5' and 3' oligonucleotides incorporated Sall and NotI or EcoRI and BamHI sites, respectively, to facilitate subcloning into pCMV-c-myc or pEYFP-N1 (BD Biosciences Clontech, Mississauga, ON, Canada), and incorporated initiation and stop codons where appropriate (Table S2).

Immunocytochemistry on co-culture

Astrocyte-neuron co-cultured on glass coverslips were washed three times in PBS, fixed in 4% paraformaldehyde at 4°C for 20 min, and washed three times in PBS. Cells were permeabilized in PBS containing 0.1% Triton X-100 (10 min) and pre-incubated with normal goat and/or normal donkey antiseraum (1:50, Santa-Cruz, Tebu bio, Le Perray en Yvelines, France) for 1 h. Then, cells were incubated at 20°C for 1 h with a goat anti-UT (1:200), a mouse anti-NeuN (1:200, Santa-Cruz) or a rabbit anti-GFAP (1:1000, Dako, Trappes, France). Specificity of the UT immuno-

labelling on astrocytes has already been demonstrated [40]. After several rinses in PBS, cells were incubated at 20°C for 2 h with Alexa 488-conjugated donkey anti-goat and anti-Alexa 594-conjugated donkey anti-rabbit IgGs diluted 1:300 (Invitrogen, Boulouge Billancourt, France).

Receptor cell surface internalization

For double-immunofluorescence, non-permeabilized living CHO cells expressing UT and/or α2β2γ2, GABAαR receptors were washed two times in PBS, and incubated with monoclonal mouse anti-c-myc and rabbit polyclonal anti-UT-HA (1:200, Santa-cruz) for 1 h on ice in DMEM. Excess antibody was removed and cells were incubated with the different receptor agonists for 30 min at RT. After several rinses in PBS, cells were fixed in 4% paraformaldehyde at 4°C for 20 min, washed three times in PBS and then incubated at RT for 2 h with appropriate secondary antibodies, i.e. Alexa-488- and 594-conjugated donkey anti-mouse IgGs diluted 1:300 (Invitrogen). After washing, coverslips were mounted in Eukitt (VWR International, Strasbourg, France).

All preparations were examined using a confocal laser-scanning microscope (Leica, Heidelberg, Germany) equipped with a Diaplax optical system, a UV laser (excitation wavelength 405 nm) and argon/krypton ion (excitation wavelengths 488/594 nm) laser.

Immunohistochemistry on cerebellar sections

Following decapitation, the cerebellum of 7-day-old (P7) Wistar rats was extracted and immersed in iced PBS. Cerebella of P7 rats were sectioned transversely into 180 µm-thick slices on a vibrating blade microtome (VT1000S, Leica Instruments). The slices were subsequently washed in PBS (pH 7.4) and postfixed in a 4% PFA solution for 20 min. Thereafter, free-floating sections were rinsed and non-specific binding was blocked by 10% normal donkey serum, 0.1% BSA and 0.3% Triton X-100 in PBS for 1 h. The tissue sections were incubated overnight at 4°C with a goat antiserum directed against rat UT (1:200, Santa Cruz), a mouse anti-calbindin (1:400, Sigma-Aldrich, Saint-Quentin Fallavier, France), a mouse anti-GFAP (1:1000, Dako), a mouse anti-NeuN raised in mouse (1:200, Santa-Cruz), an anti-doublecortin (DCX) raised in goat (1:400, Santa-Cruz), or anti-γ- and anti-γ2 GABAαR subunits (1:200, generous gift from Dr Sieghart, Brain Research Institute, Vienna, Austria). The sections were rinsed three times with PBS and incubated for 2 h at RT with Alexa 488- or 594-conjugated donkey anti-rabbit, donkey anti-goat or donkey anti-mouse (1:200, Invitrogen). After washing, slices were incubated with 4,6-diamidino-2-phenylindole (DAPI, 1:10000, Sigma-Aldrich) for 5 min to label nuclei. Finally, the sections were rinsed in PBS, and mounted with mowiol. To study the specificity of UT and other marker immunoreactivities, the following controls were performed (1) substitution of each antiserum by PBS, and evaluation of the level of fluorescence given by each type of antibody, (2) systematic mono-immunolabeling of each protein marker. The preparations were examined on a Leica SP2 upright confocal laser scanning microscope (DM RXA-UV) equipped with Acousto-Optical Beam Splitter (AOBS) system. For confocal images, Alexa-488 and Alexa-594 were excited respectively at 488 and 594 nm.

Electrophysiology

The conventional whole-cell configuration of the patch-clamp technique was used to study the GABA-gated currents in astrocytes and granule neurons, UT stably transfected CHO, CHO-UT, and CHO transiently transfected with diverse variants of UT and GABAAR subunits. After 24-h transfection, cell culture coverslips were placed in a small chamber (1.5 ml) on a stage of a...
right microscope DMLFSA (Leica, Heidelberg, Germany) and superﬁced continuously with the following bath solution containing (in mM): NaCl, 140; KCl, 2.5; HEPES, 5; CaCl2, 2; MgCl2, 1; glucose, 10 (pH 7.4 adjusted with NaOH). The patch pipettes were fabricated from 1.5 mm (outer diameter) soft glass tubes on a two-step vertical pipette puller (List-Medical, L/M-3P-A, Darmstadt, Germany). Patch electrodes had a ﬁnal resistance of 1–6 MΩ when ﬁlled with an internal pipette solution containing (in mM): KCl, 130; MgCl2, 2; CaCl2, 0.5; EGTA, 5; HEPES, 10; ATP, 1; GTP, 0.1 (pH 7.4 adjusted with KOH). ATP and GTP were added to the internal solution used to ﬁll electrode just before recording. All recordings were obtained at RT with cells voltage-clamped at −70 mV. The GABAα receptor agonist isoguvacine was prepared in the extracellular solution and was applied to cells by pneumatic pressure ejection. To prevent desensitization, isoguvacine was more often ejected during 5 s at 2-min intervals. ATP, GTP, or guanosine 5′-O-(2-thiodiphosphate) (GDPβS) were administered through the patch pipette solution. We investigated the effect of competitive inhibition of GDP-GTP exchange by including GDPβS in the pipette solution (in addition to the normal amount of GTP). For GDPβS and the KIC, immediately following patch rupture, GABA current recordings were performed and the experiment was commenced after an equilibration period of 15 min.

All peptide ligands, inhibitors or allosteric modulators of the GABAα receptor function were applied via gravity through a plastic tubing positioned in the vicinity of the cell body in order to maintain a sustained perfusion. In these conditions, drugs could reach the cell of interest after 30 s of perfusion. Isoguvacine was applied focally by pneumatic pressure ejection from a micropipette.

All current signals were ampliﬁed from an Axopatch 200A Ampliﬁer (Axon Instruments, Union City, CA, USA) and ﬁltered at 2 kHz (3 dB, four-pole, low-pass Bessel ﬁlter). Data acquisition and analysis were performed through a digitized 1200 interface using the pClamp 8 suite programs (Axon Instruments, Union City, CA, USA) and/or the Origin 4.1 analysis software (Microcal Software, Northampton, MA, USA).

The decrease of the chloride current (normalized current, I) was deﬁned as [I-Iso+I-UII/Iso]-1 where I-Iso+I-UII is the current response in the presence of various concentrations of UII and I-Iso is the control GABAαR current. Concentration response curves were generated and the data were ﬁtted by a non-linear regression analysis using Microcal Origin Software. Dose-response curves were ﬁtted using a nonlinear square-ﬁtting program to the equation: F(x) = Bmax/[1 + (EC50/X)^n], where x is the drug concentration, EC50 is the concentration of drug eliciting a half-maximal response and n is the Hill coefﬁcient.

**Cell calcium imaging**

For cell calcium imaging, Fura-2 AM (5 mM, Molecular Probe; Fisher, Cergy-Pontoise, France) was dissolved in 20% pluronic F-127 (w/v, DMEM) and then added to culture medium at ﬁnal concentrations of 5 μM and 0.02%, with 2.5 mM probenecid (Sigma-Aldrich) respectively. Cells were incubated in the dye solution for 1 h in an humididiﬁed atmosphere (37 °C, 5% CO2) and then rinsed in the standard extracellular solution used for patch-clamp experiments. For simultaneous measurements of intracellular calcium and chloride-evoked currents in CHO, patch-clamp electrodes were ﬁlled with an internal solution containing: KCl, 140; MgCl2, 4; Fura-2-pentoK, 0.25; HEPES, 10; ATP, 1; GTP, 0.1 (pH 7.4 adjusted with KOH). Fluorescence images were acquired with the right microscope DMLFSA (Leica) equipped with a digital CCD camera Coolmap HQ (Photometrics, Roper scientific, Evry, France). A high-speed scanning polychromatic light source was used for alternate excitations at wavelengths of 340 and 380 nm. The ﬂuorescence intensities at both wavelengths (340 and 390) were measured every 500 ms. Image acquisition and analysis were obtained with a MetaFluor/Metamorph Imaging System (Roper scientiﬁc). The ratio between the two images was proportional to the [Ca2+]i in the cell under study. Before an experiment, the bath ground level of ﬂuorescence (attributable to autofluorescence and camera noise) was determined and systematically subtracted.

**Cell surface expression of receptors by ELISA**

Receptor surface expression was assessed by ELISA 24 h posttransfection of CHO with cDNA encoding UT1HA or UT2HA truncated mutants (supplementary Fig. S2) or encoding UT1-UII and UT2-UII. UT1HA, UT1-UII or UT2HA, UT2-UII subunits, or U97 glial cell line before and after bUII treatment, after quantiﬁcation of HA, c-myc, β3 subunit or UT immunoreactivity. Cells were plated at 50 000 cells/well in 96-well plates coated with 0.1 mg/ml poly-L-ornithine (Sigma-Aldrich). After transfection of CHO cells with human UT1-UII and UT2-UII cDNA (Supplementary Table S2), cells were serum starved for 2 h before exposure to bUII 10−10 M during 30 min at 37°C. Cells were ﬁxed with 4% PFA in PBS for 5 min at RT, washed in PBS, and then permeabilized in 0.05% triton X100 (only for permeabilized cells) and non speciﬁc binding were blocked with PBS containing 1% FBS for 30 min at RT. The ﬁrst rat anti-HA monoclonal antibody (0.5 μg/ml, 3F10 clone, Roche, Meylan, France) and/or the anti-UT antibody (1 μg/ml, Santa Cruz), mouse anti-c-myc monoclonal antibody (1.33 μg/ml, 9E10 clone, Roche), rabbit anti-UT antibody (1 μg/ml, TEBU, Santa Cruz) or rabbit anti-β3 antibody (1:200, Abcam, Paris, France), were added for 1 h at 30°C at RT. Incubation with goat anti-rat (Thermo scientiﬁc, Fisher, Brevières, France), goat anti-mouse (Santa Cruz) or goat anti-rabbit (Tebu, Santa Cruz) conjugated alkaline phosphatase diluted at 1:1000 in PBS/FBS was carried out for 30 min at RT. The cells were washed four times with PBS, a colorimetric alkaline phosphatase substrate was added (SuperSignal ELISA, Thermo scientiﬁc, Fisher) and the resulting color reaction was measured using a Viktor multilabel plate reader (PerkinElmer, Courtaboeuf, France). Background absorbance from samples transfected with non-tagged receptors or from cells without primary antibodies were systematically measured. Results are expressed as the percentage of membrane receptor corresponding to the proportion of receptor at the plasma membrane (non permeabilized cells) to the total receptor (permeabilized cells), and normalized to the values obtained in the absence of UII. All experiments were done at least three times in triplicates.

**Receptor expression by Flow cytometry**

Human astrocytes and glioma U87 grown in 75-cm2 ﬂasks until conﬂuence were washed in PBS, detached and spun down at 4°C. The cell pellet was washed, re-suspended in PBS containing 1% BSA and incubated with 10 μg/ml non-immune rabbit IgGs for 30 min. For total receptor detection, cells were permeabilized in PBS containing 1% BSA and 0.1% saponin for 30 min. Then, cells were incubated with antibodies directed against rabbit anti-β3 subunit (1:100, Abcam) or anti-UT (1:100, Santa Cruz), diluted in PBS containing 1% BSA and 0.1% saponin at RT (22°C) for 30 min. Non permeabilized cells were directly incubated with rabbit IgGs (10 μg/ml, 30 min), and exposed to the first antibodies for 30 min at 4°C. In negative controls, the primary antibodies were omitted. Cells were incubated with an FITC- or PerCP-conjugated goat anti-rabbit secondary antibody (Santa Cruz) diluted 1:200 and measured in the FL1 (530 nm) or FL3 (670 nm) channel. Cells were analyzed on a FACSScalibur ﬂow cytometer (BD Biosciences) operated with the CellQuestTM software.
Samples in which the primary antibodies were omitted were used as control to normalize mean fluorescence levels. Dead cells and debris were excluded from the analysis by gating living astrocytes or gliona cells from size/structure density plots. Data were displayed on a logarithmic scale in arbitrary units corresponding to the mean fluorescence intensity. Each histogram plot was recorded from at least 10,000 gated events.

**Calcium mobilization assay**

CHO cells stably transiently transfected with the human UT<sup>HA</sup> or their truncated forms were plated at a density of 4×10<sup>4</sup> cells/well in flat clear bottom black 96-well plates. After 24 h in culture, cells were incubated at 37°C with 20 µl of 2 µM Fluo-4 AM dye containing 20% pluronic acid for 40 min in a 5% CO<sub>2</sub> atmosphere. Cells were washed twice with modified HBSS, and the effects of graded concentrations of UII on [Ca<sup>2+</sup>]<sub>i</sub>, were measured with a fluorometric imaging plate reader FlexStation II (Molecular Devices, Sunnyvale, CA) during 150 s with an excitation wavelength of 480 nm and an emission wavelength of 525 nm. After 18 s recording in basal conditions, 50 µl of graded concentration of UII (four-fold final concentration) was added to the incubation medium with a built-in eight-channel pipettor to assess activity. After subtraction of mean fluorescence background, baseline was normalized to 100%. Fluorescence peak values were determined for each concentration of UII, and potency (IC<sub>50</sub>) and efficacy (Emax) were calculated with the Prism 4.0 software using a logistic equation. Results were expressed as mean ± SEM.

**Chemicals**

Rat UII (hUII, rQHGTAPECFWKYCI), human UII (hUII, ETPDCFWKYCV), URP and [Orn<sup>5</sup>]URP were synthesized by the solid phase methodology on a Pioneer PerSeptive Biosystem peptide synthesizer (Applera France, Courtaboeuf, France) using the standard manufacturer’s procedures as previously described [32]. All peptides were purified on a 2.2×25-cm Vydac C<sub>18</sub> column (Alltech, Templemars, France) (>98% pure) and characterized by MALDI-TOF MS on a Voyager DE-PRO mass spectrometer (Applera France). B27, DMEM, glutamine, HEPES, non essential amino acids, sodium pyruvate and the antibiotic-antimycotic solution were purchased from InVitrogen (Fischer, Illkirch, France). EGF was obtained from Abcys (Les Ulis, France). ATP, GTP, guanosine 5’-[β;γ]triphosphate (GDPβS), the GABA<sub>3</sub> receptor agonist isoguvacine, pentobarbital, poly-D-ornithine, methyl 6,7-dimethoxy-4-ethyl-β-carboline-3-carboxylate (DMCM), a phosphatase inhibitor cocktail 2 (sodium vanadate, sodium molybdate, sodium tartrate and imidazole), quercetin, staurosporine, picrotoxin, SR95531, Ham-F12, insulin, D(+)-glucose, Tri-reagent, probenicid, and bovine serum albumin (BSA) were obtained from Sigma. FBS was from Eurobio or Lonza (France). Phloronic acid and Fura-2 pentapotassium (Fura-2 AM pentOK) salt and Fura-2 acetoxymethyl ester (Fura-2 AM) were from Molecular Probes (Leiden, Netherlands). The dynamin inhibitory peptide (DIP) was obtained from Tocris Bioscience (Ellisbury, MI, USA).

**Statistics**

All data are presented as mean ± SEM. Statistical comparisons were assessed with One-way ANOVA followed by Mann and Whitney, Newman-Keuls or Friedman post hoc tests, as relevant. P<0.05 was taken as significance.

**Supporting Information**

**Figure S1 Pharmacological and gating properties of hUII-induced regulation of GABA<sub>AR</sub>.** (A) Iso-evoked current in the absence or presence of pentobarbital (10<sup>−5</sup> M), SR95531 (10<sup>−5</sup> M) and picrotoxin (10<sup>−4</sup> M, 2 s) in CHO-GABA<sub>AR</sub>. Right, summary of the effects of modulators on the GABAergic activity. (B) Current-Voltage (I–V) relationship of the Iso-evoked current, in the absence or presence of hUII (10<sup>−8</sup> M). Data are mean ± SEM from 5 to 9 cells. *, P<0.05; **, P<0.01; ***, P<0.001 compared with the control Iso-evoked current. NS, non significant. (PPT)

**Figure S2 Expression of the UT C-terminus truncated mutants.** (A) Confocal microscope images of CHO expressing UT<sup>H<sub>4</sub>A</sup>, UT<sub>319</sub> HA, UT<sub>322</sub> HA, UT<sub>351</sub> HA, UT<sub>570</sub> HA (green). (B) Expression of the different UT<sup>HA</sup> mutants expressed as receptors in whole cells (permeabilized) or only at the cell plasma membrane (non-permeabilized) using anti-HA antibody. Data are mean ± SEM from a representative experiment in triplicate. *, P<0.05; **, P<0.01; ***, P<0.001 compared to control. Mock, empty pCMV-HA vector. (PPT)

**Figure S3 Functional expression of the UT C-terminus truncated mutants.** (A) Dose-response curves of the mean of maximum amplitude of [Ca<sup>2+</sup>]<sub>i</sub>, transients induced by hUII in CHO expressing UT<sup>H<sub>4</sub>A</sup>, UT<sub>319</sub> HA, UT<sub>322</sub> HA, UT<sub>351</sub> HA, UT<sub>570</sub> HA. The results are expressed as percentages of the corresponding control values in the absence of hUII. (B) Corresponding table summarizing EC<sub>50</sub> values and percentage of efficacy of the effect of hUII on each UT construction. Data are mean ± SEM from 3 independent experiments in duplicate. The Pearson coefficient r<sup>2</sup> close to 1 is used for significance. (PPT)

**Table S1 EC<sub>50</sub> and maximum inhibitory effects of hUII on different GABA<sub>AR</sub> subunit combinations.** Data are mean ± SEM from 3 to 23 independent experiments. ND, not determined; r<sup>2</sup>, Pearson coefficient. (PPT)

**Table S2 Primer sequences and restriction enzymes used for the different UT and GABA<sub>AR</sub> subunit constructions.** (PPT)

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**Author Contributions**

Conceived and designed the experiments: HC PG MCT VC. Performed the experiments: LD TL CL. MTS JL FM. Analyzed the data: HC LD TL. Contributed reagents/materials/analysis tools: JL FP. Wrote the paper: HC LD.