1	Estrogen receptor alpha and beta differentially mediate C5aR agonist evoked
2	Ca <sup>2+</sup> -influx in neurons through L-type voltage-gated Ca <sup>2+</sup> channels
3	Imre Farkas <sup>a</sup> , Miklós Sárvári <sup>a</sup> , Máté Aller <sup>b</sup> , Noriko Okada <sup>c</sup> , Hidechika Okada <sup>d</sup> , István
4	Likó <sup>e</sup> , Zsolt Liposits <sup>a,f</sup>
5	<sup>a</sup> Laboratory of Endocrine Neurobiology and <sup>b</sup> Laboratory of Cellular Pharmacology,
6	Institute of Experimental Medicine, Hungarian Academy of Sciences, Budapest,
7	Hungary, <sup>c</sup> Department of Immunology, Nagoya City University, Nagoya, Japan,
8	<sup>d</sup> Choju Medical Institute, Toyohashi, Japan, <sup>e</sup> Gedeon Richter Plc., Budapest,
9	Hungary, <sup>f</sup> Department of Neuroscience, Faculty of Information Technology, Pázmány
10	Péter Catholic University, Budapest, Hungary
11	
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14	Corresponding author and to whom reprint requests should be addressed:
15	Imre Farkas, PhD
16	Laboratory of Endocrine Neurobiology, Institute of Experimental Medicine,
17	Hungarian Academy of Sciences, Szigony. u. 43., H-1083 Budapest, Hungary
18	Tel.: 36-1-210-9400; Fax: 36-1-210-9944
19	Email: farkas@koki.hu

## 21 Abstract

22 Complement C5a is associated primarily with inflammation. The widespread 23 expression of its receptors, C5aR and C5L2 in neuronal cells, however, suggests 24 additional regulatory roles for C5a in the CNS. C5aR agonist (PL37-MAP) evokes  $Ca^{2+}$ -influx in GT1-7 neuronal cell line and the  $Ca^{2+}$ -influx is regulated by estradiol. 25 In the present study, we examined further the mechanism of  $Ca^{2+}$ -influx and the 26 27 contribution of the two estrogen receptor (ER) isotypes, ER $\alpha$  and ER $\beta$ , to estrogenic modulation of intracellular Ca<sup>2+</sup>-content. GT1-7 neurons were treated with isotype 28 selective ER agonists for 24h then C5aR agonist evoked Ca<sup>2+</sup>-responses were 29 measured by  $Ca^{2+}$ -imaging. Transcriptional changes were followed by real-time PCR. 30 31 We found that not only estradiol (100pM), but the ERa selective agonist PPT (100pM) enhanced the PL37-MAP-evoked Ca<sup>2+</sup>-influx (E2: 215%, PPT: 175%, 32 33 compared to the PL37-MAP-evoked  $Ca^{2+}$ -influx). In contrast, the ER $\beta$  selective agonist DPN (100pM) significantly reduced the Ca<sup>2+</sup>-influx (32%). Attenuated Ca<sup>2+</sup>-34 response (25%) was observed in Ca-free environment and depletion of the Ca<sup>2+</sup>-pool 35 by CPA eliminated the remaining elevation in the  $Ca^{2+}$ -content, demonstrating that 36 the majority of Ca<sup>2+</sup> originated from the extracellular compartment. L-type voltage-37 gated Ca<sup>2+</sup>-channel (L-VGCC) blocker nifedipine abolished the Ca<sup>2+</sup>-influx, while R-38 type Ca<sup>2+</sup>-channel blocker SNX-482 had no effect, exemplifying the predominant role 39 40 of L-VGCC in this process. Acute pre-treatments (8min) with ER agonists did not 41 affect the evoked Ca<sup>2+</sup>-influx, revealing that the observed effects of estrogens were 42 genomic. Therefore, we checked estrogenic regulation of C5a receptors and L-VGCC 43 subunits. ER agonists increased C5aR mRNA expression, whereas they differentially 44 regulated C5L2. Estradiol decreased transcription of Ca<sub>v</sub>1.3 L-VGCC subunit. Based 45 on these results we propose that estradiol may differentially modulate C5a-induced

- 46  $Ca^{2+}$ -influx via L-VGCCs in neurons depending on the expression of the two ER
- 47 isotypes.
- 48
- 49 Keywords: GT1-7 neuron, complement C5a receptor, estrogen receptor alpha,
- 50 estrogen receptor beta; voltage-gated calcium channel
- 51
- 52 Abbreviations:
- 53 C complement system
- 54 C5aR "classical" complement 5a receptor (G-protein coupled)
- 55 C5L2 "second" complement 5a receptor (non-G-protein coupled)
- 56 CPA cyclopiazonic acid
- 57 DPN Diarylpropionitrile
- 58  $E2 17\beta$ -estradiol
- 59 ER $\alpha$  estrogen receptor alpha
- 60  $ER\beta$  estrogen receptor beta
- 61 GnRH gonadotropin-releasing hormone
- 62 HBSS Hanks' Balanced Salt Solution
- 63 L-VGCC L-type voltage-gated calcium channel
- 64 PPT -4,4',4''-(4-Propyl-[1H]-pyrazole-1,3,5-triyl)trisphenol
- 65 PVN paraventricular nucleus
- 66 SON supraoptic nucleus

#### 1. Introduction

70 The complement (C) system is an ancient immune pathway comprised of 71 numerous elements activated in a cascade leading to the elimination of pathogens 72 (Speth et al., 2008). During C activation, the C5a anaphylatoxin, a 74 amino-acid long 73 fragment of the fifth component of C is released when C5 is cleaved by the C5 74 convertase. C5a binds to two receptors, the "classical" G-protein coupled C5aR 75 (CD88) and the non-G-protein coupled receptor C5L2 (GPR77). Both receptors are 76 expressed on immune and non-immune cell types. C5a binding to C5aR leads to 77 various events such as increased intracellular calcium level and activation of 78 intracellular signaling cascades resulting in functional responses e.g. recruiting and 79 activation of inflammatory cells, degranulation, delayed or enhanced apoptosis, 80 phagocytosis, histamine release, and chemotaxis (Fujita et al., 2004; Guo and Ward, 81 2005). C5L2 may function as a decoy receptor regulating the inflammatory response 82 resulted from the C5a/C5aR binding (Bamberg et al., 2010; Woodruff et al., 2011).

83 Expression of C5aR has been demonstrated in astrocytes, microglia and 84 neurons of the central nervous system (CNS) (Woodruff et al., 2010). The cellular 85 expression pattern of C5L2 is similar to that seen for C5aR. Pyramidal neurons in the 86 hippocampus and the cortex, Purkinje cells of the cerebellum and neuroblastoma cells 87 express C5aR (Farkas et al., 1999; Farkas et al., 1998a; Farkas et al., 1998b). Function 88 of the C5aR in neurons remains elusive. A C5aR-related apoptotic pathway and the 89 role of this receptor in neurodegenerative diseases such as Alzheimer's disease have 90 been suggested (Farkas et al., 2003; Fonseca et al., 2009; Fonseca et al., 2011). In 91 contrast, the neuroprotective role of C5a has also been demonstrated (Woodruff et al., 92 2010).

93	C5aR has recently been identified in hypothalamic neurons, including
94	gonadotropin-releasing hormone (GnRH)-producing cells, immortalised GnRH-
95	producing GT1-7 neurons and neurons of the paraventricular (PVN) and supraoptic
96	(SON) nuclei (Farkas et al., 2008). GT1-7 neurons establish neuronal network with
97	co-ordinated activity and produce GnRH in a pulsatile fashion (Liposits et al., 1991;
98	Wetsel et al., 1992). Pulsatility and volume of the secretion is in strong correlation
99	with synchronised firing (Moenter et al., 2003; Thiery and Pelletier, 1981; Wilson et
100	al., 1984). Various factors released during inflammation can play role in the function
101	of these cells (Karsch et al., 2002). Cannabinoids also affect GnRH neurons by
102	utilising the retrograde endocannabinoid signaling mechanism (Farkas et al., 2010).

103 The estrogen receptor alpha and beta (ER $\alpha$  and ER $\beta$ ) are expressed in 104 numerous hypothalamic neurons, such as GnRH cells and the neurons of PVN and 105 SON (Hrabovszky et al., 2004; Shughrue et al., 1997; Shughrue and Merchenthaler, 106 2001), GT1-7 cells express both ER subtypes (Roy et al., 1999). 17 $\beta$ -estradiol (E2) 107 can modulate the electric function of GT1-7 cells and exert both negative and positive 108 feedback on the firing (Christian et al., 2005; Farkas et al., 2007).

109 Our previous experiments have shown that administration of a C5aR agonist results in robust calcium ( $Ca^{2+}$ ) influx in GnRH neurons. In addition, E2 pre-treatment 110 elevates this Ca<sup>2+</sup>-response suggesting that the signal transduction pathways related to 111 112 the C5aRs and the ERs, respectively, can modulate each other (Farkas et al., 2008). Change in the intracellular Ca<sup>2+</sup>-milieu can heavily affect firing properties of the 113 114 neurons. Firing can be fine-tuned for example by the opening and closing of the Ltype voltage-gated Ca<sup>2+</sup>-channels (L-VGCCs). Ca<sub>v</sub>1.2 and Ca<sub>v</sub>1.3 subunits of the L-115 116 VGCCs are strongly involved in spontaneous firing and pacemaking (Zuccotti et al., 117 2011). In the present study, therefore, we investigated further, how C5aR and ER

118 subtypes interact using the immortalised GnRH-producing GT1-7 neurons as a 119 neuronal model with  $Ca^{2+}$ -imaging and quantitative real-time PCR methods. Genomic 120 and non-genomic actions of ER subtypes on the C5aR-mediated  $Ca^{2+}$ -influx were 121 examined. Potential sources of the increased  $Ca^{2+}$ -content were also studied, including 122 the role of various VGCCs and the intracellular  $Ca^{2+}$ -pool.

#### 124 **2.** Materials and methods

### 125 2.1 Cell culture

126 GnRH-producing immortalised GT1-7 neurons were cultured in Dulbecco 127 Modified Eagle Medium (DMEM) containing high-glucose and supplemented with 128 10% fetal calf serum (FCS) and 5% horse serum (HS). Prior to ER agonist treatment 129 the culturing medium was replaced with a steroid/thyroid- and phenol red-free one 130 and cells were cultured in this medium for 24hrs. Subsequently, the cells were treated 131 with 17 $\beta$ -estradiol (E2, SIGMA), the highly potent ER $\beta$  receptor agonist DPN 132 (Diarylpropionitrile, Tocris) and the ERa receptor agonist PPT (4,4',4"-(4-Propyl-133 [1H]-pyrazole-1,3,5-trivl)trisphenol, Tocris) at various concentrations (100pM -134 20nM) for 24h and then used for calcium imaging and RT-PCR experiments. In the 135 experiments examining effect of acute treatment, the E2, DPN, and PPT were added 136 to the cells 8min before starting the calcium imaging recording.

137

## 138 2.2 Calcium imaging

139 Cultured GT1-7 cells were loaded with the calcium-sensitive fluorescent dye Fura-2 AM (1µM; Molecular Probes, Eugene, Oregon) in loading buffer Hanks' 140 141 Balanced Salt Solution (HBSS) containing 0.1% DMSO and Pluronic-F127 (1µM, 142 Molecular Probes) for 1.5h at room temperature (RT). After washing with HBSS, the 143 experiments were carried out at RT. The antisense homology box peptide fragment of 144 the C5a (RAARISLGPRCIKAFTE) was synthesised in multiple antigenic peptide 145 form (termed PL37-MAP, 2.5µM). Sequence of the PL37 is a fragment sequence of 146 the C5a, representing a "strong" antisense homology box region in the C5a (Baranyi 147 et al., 1995). Our previous works applying both C5a and PL37, respectively, 148 demonstrated that PL37 is a potent agonist of the C5aR preserving the biological

149 activity of the C5a and triggering responses similar to that of the C5a (Baranyi et al., 1996; Fujita et al., 2004). The peptide was pipetted directly onto the cells in HBSS 150 151 after a 1min baseline recording and then the diluted peptide remained in the HBSS 152 during recording. In the case of E2, DPN, PPT pre-treatment, the cells were pre-153 treated with them as described in the "Cell culture" section and all of the rinsing and extracellular solutions contained the same concentration of E2, DPN and PPT. After 154 155 the 1min baseline recording the PL37-MAP peptide was introduced into the bath fluid 156 containing E2, DPN, and PPT and then the diluted peptide remained in the HBSS-E2 or HBSS-DPN or HBSS-PPT mixture during recording. 157

158 When PL37-MAP was applied in  $Ca^{2+}$ -free extracellular solution (phosphate 159 buffered salt solution=PBS, pH 7.4), the HBSS was changed to PBS just before 160 starting the recording, except when the cells were treated with cyclopiazonic acid 161 (CPA, 10µM, Tocris). CPA is a specific blocker of the Ca<sup>2+</sup>-ATP-ase of the 162 intracellular Ca<sup>2+</sup>-store endoplasmic reticulum and depletes these Ca<sup>2+</sup>-stores. CPA 163 was applied to the GT1-7 neurons in PBS 30min before starting the measurements.

164 The VGCC blockers nifedipine (10 $\mu$ M, SIGMA) and SNX-482 (100nM) were 165 added to the HBSS just before starting the Ca<sup>2+</sup>-imaging measurement and remained 166 in the HBSS during recording.

167 The experiments were carried out with an ARGUS HiSCA Ca<sup>2+</sup>-imaging 168 system (Hamamatsu Photonics, Hamamatsu, Japan) or with an Olympus BX50WI 169 microscope equipped with a Polychrome II monochromator (TILL Photonics), a 170 cooled CCD camera (Photometrics Quantix, Tucson, AZ, USA), and controlled by the 171 Axon Imaging Workbench 6.0 software (Axon Instruments, Union City, CA, USA). 172 The ratio of the fluorescent signals obtained at excitation wavelengths of 340 and 380 173 nm was used to determine changes in the intracellular Ca<sup>2+</sup>-concentration. The surface density of the cultured cells was 500,000-750,000 cells/cm<sup>2</sup>, the magnification of the objective lens used was 40x, and the area of the calcium imaging acquisition was 0.038 mm<sup>2</sup>. Before starting the calcium imaging recordings, an area was chosen where at least 7 individual cells without overlaps could be clearly observed and measured (7-15 neurons depending on the surface density of the cells in the area measured).

180

### 181 2.3 Total RNA isolation from GT1-7 cells

Total RNA was isolated from GT1-7 cells using the RNeasy Mini Kit
(QIAGEN, Hilden, Germany). RNA analytics included capillary electrophoresis using
Agilent 2100 Bioanalyzer (Santa Clara, CA). All RNA samples displayed RNA
integrity numbers (RIN) above 8.5.

186

### 187 2.4 Quantitative real-time PCR

188 Inventoried TaqMan assays were selected to study in depth the regulation of 189 genes of our interest by quantitative real-time PCR. Each assay consisted of a FAM 190 dye-labeled TaqMan MGB probe and two PCR primers. Every assay was optimised 191 by the manufacturer to run under universal thermal cycling conditions with a final 192 reaction concentration of 250nM for the probe and 900nM for each primer. Reverse 193 transcription and real-time PCR were run as described earlier (Sarvari et al., 2010). 194 RealTime StatMiner (Integromics, Granada, Spain) software and relative 195 quantification against calibrator samples ( $\Delta\Delta$ Ct) were used for analysis of Applied 196 Biosystems TaqMan gene expression assays. Two house-keeping genes (Gapdh, Hprt) 197 were applied as internal controls. The geometric mean of Ct values of Gapdh and 198 Hprt1 was used for subsequent  $\Delta$ Ct calculation (Vandesompele et al., 2002). Relative

199	quantity (RQ) represents the change in the expression of a given gene in response to a
200	treatment compared to basal (control) expression of the given gene. We considered
201	changes with RQ>1.5 as up-regulation or RQ<0.67 as down-regulation (21).
202	
203	2.5 Statistical analysis
204	Ca <sup>2+</sup> -imaging recordings using the fluorescence ratio obtained at 340 and 380
205	nm wavelengths were baseline corrected, then the area-under-curve data of the
206	records representing the net $Ca^{2+}$ -influx were analyzed. Group data of the cells (n $\geq$ 7)
207	were expressed as mean ± standard error (SEM). Statistical significance was analyzed
208	using ANOVA followed by Newman-Keuls (NK) test (GraphPad Software Inc.,
209	USA), and considered at p<0.05.

#### **3. Results**

212 3.1 Estrogens differentially modulate the  $Ca^{2+}$ -influx evoked by the C5aR agonist

The C5aR agonist peptide PL37-MAP (2.5µM) triggered robust Ca<sup>2+</sup>-influx in 213 GT1-7 neurons (Fig. 1a). Onset of the elevation of intracellular Ca<sup>2+</sup>-concentration 214 215 was within 1-1.5min after application of the peptide. When the neurons were pretreated with E2 (100pM, 24h), the evoked Ca<sup>2+</sup>- influx started earlier (in less than 30 216 217 seconds) and the amplitude of the records was higher than that of the recordings 218 evoked by the PL37-MAP alone (Fig. 1b). Pre-treatment with the selective ERa agonist PPT (100pM, 24h) elevated the PL37-MAP-triggered Ca<sup>2+</sup>-influx. The 219 elevation was similar to the increased  $Ca^{2+}$ -influx measured with E2 (Fig. 1c). In 220 221 contrast, when the cells were pre-treated with the selective ER<sub>β</sub> agonist DPN (100pM, 24h), the elevation of the  $Ca^{2+}$ -content was much lower and started in 1.5-2min after 222 223 introducing the PL37-MAP peptide into the bath fluid (Fig. 1d).

224 Histogram of the normalised area-under-curve data representing the net changes in the intracellular free Ca<sup>2+</sup> concentration showed significant increase when 225 PL37-MAP was applied to GT1-7 neurons in the presence of either E2 (100pM, 24h) 226 or PPT (100pM, 24h), revealing facilitation of the PL37-MAP-evoked Ca<sup>2+</sup>-influx by 227 228 both E2 and PPT. On the other hand, DPN (100pM, 24h) significantly decreased (Fig. 1e) the Ca<sup>2+</sup>-influx evoked by PL37-MAP (E2: 226.6  $\pm$  27.52%; PPT: 159.1  $\pm$ 229 16.74%; DPN 18.6  $\pm$  10.47% of the Ca<sup>2+</sup>-influx evoked by PL37-MAP alone; 230 231 p<0.001). Application of higher concentrations of ER agonists resulted in similar 232 effect except in the case of PPT (Fig. 1e). E2 (20nM, 24h) increased the PL37-MAPevoked Ca<sup>2+</sup>-influx whereas DPN (20nM, 24h) attenuated it significantly (E2: 169.7  $\pm$ 233 16.67%; DPN 10.9  $\pm$  3.65% of the Ca<sup>2+</sup>-influx evoked by PL37-MAP alone; 234 235 p<0.001).

# 237 3.2 The effects of ER agonists on the $Ca^{2+}$ -influx are genomic

In order to determine whether the observed effect of ER agonists on the PL37-238 MAP-evoked Ca<sup>2+</sup>-influx was genomic, PL37-MAP was applied after an acute (8min) 239 application of E2, PPT, and DPN. Ca<sup>2+</sup>-imaging experiments demonstrated that short 240 241 administration of ER agonists failed to influence the change in the intracellular free 242 Ca<sup>2+</sup>-concentration triggered by PL37-MAP (Fig. 1f). Examination of the area-undercurve data showed no significant differences (E2:  $115.1 \pm 13.78\%$ ; PPT:  $132.2 \pm$ 243 13.97%; DPN 146.9  $\pm$  45.22% of the Ca<sup>2+</sup>-influx evoked by PL37-MAP alone). The 244 245 results revealed that genomic effects of the E2, DPN, and PPT were necessary to modulate the Ca<sup>2+</sup>-response. 246

# 247 3.3 The major $Ca^{2+}$ -source is extracellular

Potential sources of the increase in the intracellular Ca<sup>2+</sup>-content were also 248 investigated with Ca<sup>2+</sup>-imaging on the GT1-7 neurons (Fig. 2a-g). Application of 249 PL37-MAP in Ca<sup>2+</sup>-free extracellular fluid (PBS) resulted in a significantly lower 250 Ca<sup>2+</sup>-response than in HBSS, which was independent from the ER agonist pre-251 252 treatment (p<0.001). Administration of PL37-MAP resulted in a significantly attenuated elevation in  $Ca^{2+}$ -concentration (25.3 ± 4.91% of the  $Ca^{2+}$ -increase evoked 253 by PL37-MAP in Ca<sup>2+</sup>-containing extracellular fluid). Nevertheless, the effect of the 254 PL37-MAP was not entirely eliminated in PBS, suggesting, that intracellular Ca<sup>2+</sup>-255 256 sources were also activated during the process. Therefore, the effect of PL37-MAP 257 was also examined when CPA (10µM, 30min) was present in the PBS. Depletion of the  $Ca^{2+}$ -store in the endoplasmic reticulum by CPA resulted in ablation of the 258 increase in the intracellular free Ca<sup>2+</sup>-content (3.2  $\pm$  0.66% of the Ca<sup>2+</sup>-increase 259 evoked by PL37-MAP in Ca<sup>2+</sup>-containing extracellular fluid). Application of PL37-260

MAP in PBS in the presence of E2, DPN, or PPT (20nM, 24h) demonstrated significant decrease of the Ca<sup>2+</sup>-response (E2: 16.9  $\pm$  1.63%; PPT: 22.9  $\pm$  5.82%; DPN: 27.5  $\pm$  5.45% of the Ca<sup>2+</sup>-increase evoked by PL37-MAP in Ca<sup>2+</sup>-containing extracellular fluid), however, these data did not differ from the Ca<sup>2+</sup>-increase evoked by PL37-MAP in Ca<sup>2+</sup>-free solution.

266 3.4 L-type but not the R-type voltage-gated  $Ca^{2+}$ -channels are involved in the PL37-

267 *MAP-evoked*  $Ca^{2+}$ *-influx* 

In order to investigate which Ca<sup>2+</sup>-channel was involved in the Ca<sup>2+</sup>-influx 268 triggered by C5aR activation, PL37-MAP was applied to GT1-7 neurons in the 269 presence of blockers of various VGCCs. The L- and the R-type Ca<sup>2+</sup>-channels have 270 271 been reported as the most abundant ones in the GT1-7 cells (Watanabe et al., 2004), therefore these two channels were examined. Application of nifedipine (10 µM), the 272 273 inhibitor of the L-VGCC, resulted in a significantly reduced Ca<sup>2+</sup>-influx evoked by 274 PL37-MAP (Fig. 3a-f). Pre-treatment the cells with E2, PPT or DPN (20nM, 24h) did 275 not modify effect of nifedipine (nifedipine alone:  $39.7 \pm 3.29\%$ ; E2:  $35.8 \pm 6.94\%$ ; PPT:  $32.4 \pm 4.66\%$ ; DPN:  $34.2 \pm 6.48\%$  of the data measured with PL37-MAP alone; 276 p<0.001). 277

In contrast to nifedipine, SNX-482 (100nM), the inhibitor of the R-type  $Ca^{2+}$ -278 channels, had no effect on the changes in the intracellular Ca<sup>2+</sup>-concentration evoked 279 by PL37-MAP (Fig. 4a-b, f). In the presence of SNX-482, the triggered Ca<sup>2+</sup>-influx 280 did not differ from the one measured with PL37-MAP alone (SNX:  $81.6 \pm 13.95$  %). 281 In addition, block of the R-type Ca<sup>2+</sup>-channels did not influence effect of the pre-282 treatment with E2, PPT or DPN (20 nM, 24 h) (Figs. 4c-f). The Ca<sup>2+</sup>-influx increased 283 upon E2 whereas decreased upon DPN pre-treatment significantly (E2: 139.5 ± 284 14.63%, PPT: 86.6 ± 5.33%, DPN: 43.3 ± 6.15%, p<0.001). 285

- 287 3.5 Estrogenic modulation of the transcription of the C5a receptors and the  $Ca_v 1.3$ 288 subunit of the L-type  $Ca^{2+}$ -channel in GT1-7 cells
- We examined the effects of E2 and isotype selective ER agonists on the transcription of genes encoding C5a receptors C5aR and C5L2, and L-VGCC subunits Ca<sub>v</sub>1.2, and Ca<sub>v</sub>1.3, by real-time PCR. We demonstrated estrogenic regulation of the C5a receptor genes (Table 1). C5ar1 (the classical C5aR) was up-regulated by the three ER agonists. C5L2 was regulated differentially, E2 increased while PPT decreased its transcription. Cacnald (gene for Ca<sub>v</sub>1.3) was regulated only by E2 whereas Cacnalc (Ca<sub>v</sub>1.2) showed no estrogenic regulation.

**4. Discussion** 

In the present study, we examined further estrogenic modulation of the C5aR agonist-evoked Ca<sup>2+</sup>-response using the GnRH-producing GT1-7 cell line as a neuronal model and applying isotype selective ER agonists. We demonstrated that i) ER $\alpha$  and ER $\beta$  agonists differentially modulated the C5aR agonist-evoked Ca<sup>2+</sup>-influx, ii) estrogenic modulation was dependent on genomic effects, iii) Ca<sup>2+</sup>-influx was mediated primarily through L-VGCC, iv) estrogens up-regulated C5aR mRNA expression while differentially regulated C5L2.

4.1 Estrogens differentially modulate the PL37-MAP-evoked Ca<sup>2+</sup>-influx

Our present results showed that the C5aR agonist-evoked Ca<sup>2+</sup>-influx was 306 307 differentially mediated by various ER agonists in GT1-7 neurons. Expression of 308 C5aR, and ER $\alpha$  and ER $\beta$  has long been reported in various types of neurons (Farkas 309 et al., 2003; Farkas et al., 2008; Hrabovszky et al., 2004; Hrabovszky et al., 2000; 310 Hrabovszky et al., 2001; Shughrue et al., 1997; Shughrue and Merchenthaler, 2001; Stahel et al., 1997a; Stahel et al., 1997b; Wilson et al., 2002; Woodruff et al., 2010). 311 Differential modulation by ER $\alpha$  and ER $\beta$  could be important, because Ca<sup>2+</sup>-influx 312 313 evoked by the activation of C5aR can differentially affect functions of a neuronal cell, 314 such as firing pattern, shape of after-hyperpolarisation and depolarising after-315 potentials, neurotransmitter release, plasticity, gene transcription, and vulnerability 316 (Berridge, 1998; Zuccotti et al., 2011).

In the present experiments DPN decreased the  $Ca^{2+}$ -influx evoked by PL37-MAP. Nevertheless, the amplitude of the inward ion current in the ER $\beta$ -expressing GnRH neurons from slices obtained from E2 substituted mice was higher than those from ovariectomized mice (Farkas et al., 2008). The reasons of the discrepancy may lie in the differences between the two models.

322 4.2 Estrogenic modulation of the evoked  $Ca^{2+}$ -influx is dependent on genomic effects

323 Estrogenic modulation of the  $Ca^{2+}$ -signal evoked by the activated C5aR was genomic rather than rapid in our experiments. Numerous estradiol-regulated genes 324 325 have already been identified in GT1-7 neurons by expression profiling (Varju et al., 326 2009). Majority of the responding genes were up-regulated in these cells, including 327 potassium channel subunits and transporters, transcription factors, molecules related 328 to cell death, immune response, neurotransmitter, hormone and neuropeptide 329 receptors, regulators of G-protein signaling. Those results support our present data 330 showing up-regulation of C5aR.

In our present experiments, we found no acute effect of E2 on the  $Ca^{2+}$ -influx 331 332 evoked in the GT1-7 cells. In another model, published from another laboratory (Sun 333 et al., 2010), both genomic and rapid changes resulted from the E2 administration 334 were reported by potentiating the Ca<sup>2+</sup>-current in GnRH neurons in the acute brain slice in 5min. This report described, however, that percentage of the responding cells 335 336 depended upon the concentration of E2 and only doses of E2 much higher than used 337 in our experiments could evoke response with high rate of success. In, addition, the 338 GnRH neurons presented an "all or none" ability to respond to acutely administered 339 E2 (Sun et al., 2010). The observed discrepancies may reflect differences in the basic 340 physiology and regulation of GnRH neurons integrated within the preoptic brain slice 341 preparation versus the immortalised GT1-7 neurons cultured in vitro.

342 4.3 L-type but not R-type Ca<sup>2+</sup>-channels are involved in the PL37-MAP-evoked Ca<sup>2+</sup>343 influx

In GT1-7 cells, the two major VGCCs are the L- and R-type channels (Watanabe et al., 2004). Both of them play a critical role in the regulation of  $Ca^{2+}$ dependent GnRH-release. In addition, R-type  $Ca^{2+}$ -channels have been reported to be 347 involved in the release of neurotransmitters in calyx-type synapses of the medial 348 nucleus of the trapezoid body, oxytocin neurons, and adrenal chromaffin cells 349 (Albillos et al., 2000; Wang et al., 1999; Wu et al., 1998; Wu et al., 1999). R-type channels are responsible for the dendritic Ca<sup>2+</sup>-influx induced by action potentials in 350 351 CA1 pyramidal neurons of the hippocampus (Magee and Johnston, 1995; Sabatini and Svoboda, 2000). L-VGCCs contribute to Ca<sup>2+</sup>-dependent gene transcription and can 352 353 modulate firing properties of neurons (Gomez-Ospina et al., 2006; Zuccotti et al., 2011). Our results have now revealed that the  $Ca^{2+}$ -ion current resulted from the 354 355 activation of the C5aR in GT1-7 neurons passed through the L-VGCCs but not via Rtype channels. In addition, differential modulation of the  $Ca^{2+}$ -influx by ER agonists 356 357 affected the function of the L-type channel. These data suggest that physiological 358 functions of the L-VGCC such as regulation of the GnRH release, parameters of the 359 firing, and various gene transcriptional events are affected by C5aR activation. 360 Modulation of the L-VGCC by E2 and DPN has recently been reported in GnRH 361 neurons of the acute brain slice of the mice demonstrating changes in the ion current 362 via these channels under physiological conditions (Sun et al., 2010). Similarly, modulation of the T-type voltage-gated Ca<sup>2+</sup>-channels by estradiol was also 363 364 demonstrated (Bosch et al., 2009; Qiu et al., 2006; Zhang et al., 2009).

It is an intriguing question, how the activation of C5aR can regulate opening of a VGCC. One possibility is a change in the threshold level by phosphorylation of the L-VGCC, occurring as a downstream event of the C5aR activation. G-proteincoupled receptor activation can result in activation of diverse pathways involving enzymes such as protein kinase A or protein kinase C yielding phosphorylation of the L-VGCC, in particular its  $Ca_v 1.2$  or  $Ca_v 1.3$  subunits (Dai et al., 2009; Dolphin, 2009). This phosphorylation could eventually modify various electric parameters of the 372 neurons such as open probability of the channel or the threshold level. Nevertheless, 373 the existence of a C5aR-related phosphorylation of L-VGCC requires further 374 examination. Other G-protein mediated mechanisms can also be involved such as 375 direct G-protein related modulation of the L-VGCCs (Currie, 2010; Tedford and 376 Zamponi, 2006). In this paradigm, the L-VGCC molecule possesses residues interacting directly with the  $\beta\gamma$  subunits of the G-protein. The interaction is 377 membrane-delimited, i.e. involves a second messenger molecule that remains 378 379 associated with the plasma membrane, rather than diffusing to the channel via a 380 cytoplasmic pathway (Hille, 1994).

Our Ca<sup>2+</sup>-imaging measurements showed that in addition to the extracellular 381 sources, intracellular Ca2+-stores were also involved in the elevation of the Ca2+-382 content evoked by the activation of the C5aR. The intracellular Ca<sup>2+</sup>-stores could be 383 triggered to release Ca<sup>2+</sup> directly via the C5aR-related signal transduction pathway 384 385 (Nishiura et al., 2010). Another possible pathway for this action is that C5aRactivation opens the L-VGCCs first and then the L-VGCCs activate the intracellular 386 Ca<sup>2+</sup>-stores via a putative coupling (Kim et al., 2007; Kolarow et al., 2007), however, 387 388 these opportunities require further elaboration.

389

# 390 4.4 C5aR agonist and estrogens modulate function of L-VGCC crucial in firing

The characteristic firing pattern is a crucial feature of the hormone secreting neurons. Pulsatile release of the GnRH, for example, is indispensable for the proper function of the reproductive system (Moenter et al., 2003). In addition, the synchronous firing of the GnRH-producing neurons correlates with this pulsatility (Moenter et al., 2003). The Ca<sup>2+</sup>-channels mediates how the neurons fire, therefore, effects disturbing the intracellular Ca<sup>2+</sup>-milieu could have an effect on the firing 397 properties of the GnRH cells and consequently, the pulsatile secretion of GnRH. The 398 L-VGCC is considered as one of the key mediators of the firing pattern (Zuccotti et 399 al., 2011). Its Ca<sub>v</sub>1.2 and Ca<sub>v</sub>1.3 subunits expressed in neurons and endocrine cells, 400 such as pancreatic beta, adrenal chromaffin cells (Catterall et al., 2005) and contribute 401 to the spontaneous firing and pacemaking of the neurons (Zuccotti et al., 2011). 402 However, literature data show activity-dependent differences between them. Cav1.3 is 403 more effective at low levels of activity such as during interburst intervals, and Cav1.2 404 is more efficient at high levels of activity such as during interspike intervals in the 405 bursts (Zhang et al., 2006). It has recently been reported that E2 decreases mRNA 406 expression of the  $Ca_v 1.2$ , but has no effect on the  $Ca_v 1.3$  subunit in the hippocampus 407 of aged female rats (Brewer et al., 2009). Our real-time PCR measurements revealed 408 that the Ca<sub>v</sub>1.3 subunit was down-regulated by E2 in GT1-7 cells. This discrepancy 409 might originate from the cell-type differences in the two experimental models.

410 The point of convergence of the signals coming from C5aR and ERs, was not the expression of the  $Ca^{2+}$ -channel, because the  $Ca_v 1.2$  and  $Ca_v 1.3$  subunits were not 411 regulated by DPN and PPT. Therefore, our present data suggest that any differential 412 413 transcriptional regulation of elements of C5a/C5aR signaling which might be involved in the explored differential effects of the ER agonists on the evoked Ca<sup>2+</sup>-influx 414 415 should be upstream of L-VGCC. Possible candidates could be the regulator molecules 416 of G-protein signaling (RGS2, RGS9 and RGS 10) which were earlier shown to be 417 regulated by E2 in the GT1-7 cells (Varju et al., 2009).

418 *4.5 C5aR is up-regulated by ER agonists* 

In our studies, the expression of classical C5aR was up-regulated by E2 and the used isotype selective ER agonists, suggesting that neurons could respond more effectively to the inflammatory mediator C5a in the presence of estrogens. In contrast,

422 the expression of C5L2 was differentially regulated by ER agonists, displaying the 423 up-regulation of this receptor by E2 and the attenuation of its expression by PPT. As 424 the decoy receptor modulates the performance of the classical C5aR, the elucidation 425 of this inverse regulatory trend warrants further investigation.

426 The results of the present in vitro study raise the questions of how activation of C5aR could occur by its ligand, the C5a in hormone secreting neurons in vivo. The 427 428 hypophysiotrophic axonal projections of numerous hormone secreting neurons 429 terminate outside the blood brain barrier, suggesting that these neurons are capable of 430 monitoring C5a released either in the hypothalamus or the blood. Since several rodent 431 hypothalamic neurons have been shown to express functional C5a receptors (Farkas et 432 al., 2008), it is reasonable to assume that the inflammatory mediator C5a can alter the 433 physiological properties and cellular functions of these neurons. In these mechanisms, 434 the Ca<sup>2+</sup>-influx occurring via L-VGCCs might have a pivotal role.

435

### 5. Conclusions

Summing up, this study provided evidence that C5aR-mediated Ca<sup>2+</sup>-signaling 438 439 can be differentially modulated via ER $\alpha$  and ER $\beta$ . In addition, estrogens potentiate the 440 sensitivity of GT1-7 neurons for C5a by up-regulation of C5aR through ERa and ER $\beta$ . C5aR activation leads to Ca<sup>2+</sup>-influx through L-VGCCs. Although the 441 442 transcription of the Ca<sub>v</sub>1.3 L-VGCC subunit is regulated by E2, the isotype specific ligands had no effect, indicating that these subunits are not the primary targets of ERa 443 444 and ERβ agonist actions upon C5a/C5aR signaling. The significance of the present 445 findings relates to the better understanding of the differential impact of estrogens on 446 the C5a-evoked response of neurons which express ER $\alpha$  and/or ER $\beta$ .

447

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456

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646 Figure legends

Figure 1. Ca<sup>2+</sup>-influx evoked by the C5aR agonist PL37-MAP (PL37) in the 647 presence of E2, DPN, or PPT in the GT1-7 neurons. a) The PL37-MAP induced 648 robust Ca<sup>2+</sup>-influx in the cells. b) Pre-treatment of the GT1-7 cells with E2 (100pM, 649 24h) potentiated the Ca<sup>2+</sup>-influx significantly. Onset of the response started earlier and 650 651 the amplitude of it was higher than without pre-treatment. c) Pre-treatment with the 652 PPT (100pM, 24h) resulted in an elevated Ca<sup>2+</sup>-influx. d) DPN pre-treatment (100pM, 24h), however, attenuated the  $Ca^{2+}$ -influx evoked by the PL37-MAP, significantly. e) 653 The histogram shows the area-under-curve values, representing the net  $Ca^{2+}$ -influx in 654 the neurons, in the percentage of the  $Ca^{2+}$ -influx measured with the PL37-MAP alone 655 656 (PL37+100 pM E2: 226,6±27,52%; PL37+100 pM PPT: 159,1±16,74%; PL37+100 pM DPN: 18,6±10,47%; PL37+20 nM E2: 169,7±16,67%; PL37+20 nM PPT: 657 658 80,2±9,73%; PL37+20 nM DPN: 10,9±3,65%) f) Acute (8min) pre-treatments with the E2, DPN, or PPT showed no significant effect on the PL37-MAP-evoked Ca<sup>2+</sup>-659 influx, demonstrating that the observed effect of E2, DPN, or PPT was genomic 660 (PL37+E2: 115,1±13,78%; PL37+PPT: 132,1±13,97%; PL37+DPN: 146,9±45,22%). 661 \*=p<0.05, \*\*=p<0.01. Arrow shows the onset of the administration of the PL37-662 663 MAP.

664

Figure 2.  $Ca^{2+}$ -influx evoked by the PL37-MAP (PL37) in the presence of CPA, E2, DPN, or PPT in the GT1-7 neurons in  $Ca^{2+}$ -free extracellular solution. a-f) Calcium imaging recordings show that extracellular calcium sources play important role in the  $Ca^{2+}$ -influx. g) The histogram reveals that the  $Ca^{2+}$ -free extracellular environment decreased the  $Ca^{2+}$ -response significantly, demonstrating that majority of the elevation in the  $Ca^{2+}$ -concentration was from the extracellular source. This change 671 was independent from the pre-treatment with E2, DPN or PPT. CPA, a depletor of the 672  $Ca^{2+}$ -pool in the endoplasmic reticulum, however, eliminated the remaining  $Ca^{2+}$ -673 response in the neurons, showing that the endoplasmic reticulum was the intracellular 674 source of the remaining  $Ca^{2+}$ -response (PL37 in  $Ca^{2+}$ -free: 25.3±4.91%; PL37+CPA 675 in  $Ca^{2+}$ -free: 3.2±0.66%; PL37+E2 in  $Ca^{2+}$ -free: 16.9±1.63%; PL37+PPT in  $Ca^{2+}$ -676 free: 22.9±5.82%; PL37+DPN in  $Ca^{2+}$ -free: 27.5±5.45%). \*=p<0.05, \*\*=p<0.01. 677 Arrow shows the onset of the administration of the PL37-MAP.

678

Figure 3. Ca<sup>2+</sup>-influx evoked by the PL37-MAP in the presence of nifedipine 679 680 (blocker of L-type Ca<sup>2+</sup>-channels), E2, DPN, or PPT in the GT1-7 neurons. a-b) 681 Calcium imaging recordings show that comparing to the control, nifedipine (Nif) eliminated the Ca<sup>2+</sup>-influx significantly. c-e) This decrease was independent from the 682 683 presence of the E2, DPN, or PPT, demonstrating role of the L-type Ca<sup>2+</sup>-channels. f) The histogram of the area-under-curve data shows the values expressed in the 684 percentage of the Ca<sup>2+</sup>-influx measured with the PL37-MAP alone (PL37+Nif: 685 35.8±6.94%; 686 39.7±3.29%; PL37+E2+Nif: PL37+PPT+Nif: 32.4±4.66%; PL37+DPN+Nif: 32.4±4.66%). \*=p<0.05. Arrow shows the onset of the 687 688 administration of the PL37-MAP.

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Figure 4. Ca<sup>2+</sup>-influx evoked by the PL37-MAP in the presence of SNX-482 (blocker of R-type Ca<sup>2+</sup>-channels), E2, DPN, or PPT in the GT1-7 neurons. a-e) The ratiometric graphs revealed that SNX-482 (SNX) showed no effect on the Ca<sup>2+</sup>- influx evoked by the PL37-MAP, and it exerted no influence on the effect of the ER agonists, demonstrating that the R-type Ca<sup>2+</sup>-channel plays no role in these processes.
f) The histogram of the area-under-curve data shows the values expressed in the

696	percentage of the Ca <sup>2+</sup> -influx measured with the PL37-MAP alone (PL37+SNX:
697	81.6±13.95%; PL37+E2+SNX: 139.5±14.63%; PL37+PPT+SNX: 86.6±5.33%;
698	PL37+DPN+SNX: 43.3±6.15%). *=p<0.05. Arrow shows the onset of the
699	administration of the PL37-MAP.
700	

701 **Table legend** 

702 Table 1. Transcriptional modulation of L-type Ca<sup>2+</sup>-channel subunits and C5a

703 **receptors.** Transcription of L-type  $Ca^{2+}$ -channel  $Ca_v 1.2$  (Cacna1c) and  $Ca_v 1.3$ 704 (Cacna1d) subunits, and C5a receptors CD88 (C5ar1) and C5L2 (Gpr77) was 705 followed by real-time PCR. Table shows the arithmetic mean and standard deviation 706 of relative quantities (RQ) from two independent experiments. Arrows show direction

- 707 of regulation of the respective transcript where change is significant. The lack of
- rotation evidence for regulation is denoted with "-".
- 709
- 710











Fig. 1.



712

Ratio











719	
720	

Gene symbol and physiologic name	E2		РРТ		DPN	
Cacna1c (L-type Ca <sup>2+</sup> - channel Ca <sub>v</sub> 1.2 subunit)	0.867±0.164		0.905±0.073	-	0.748±0.052	-
Cacna1d (L-type Ca <sup>2+</sup> - channel Ca <sub>v</sub> 1.3 subunit)	0.413±0.326	ţ	0.960±0.023	-	0.795±0.117	-
C5ar1 (C5aR)	2.476±0.715	1	2.039±0.241	1	1.938±0.743	1
Gpr77 (C5L2)	1.634±0.336	1	0.562±0.101	ţ	0.955±0.577	-

Table 1.