

1 **LOW DEGREE OF OVERLAP BETWEEN KISSPEPTIN, NEUROKININ B AND DYNORPHIN**
2 **IMMUNOREACTIVITIES IN THE INFUNDIBULAR NUCLEUS OF YOUNG MALE HUMAN**
3 **SUBJECTS CHALLENGES THE ‘KNDY NEURON’ CONCEPT**

4
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8 Abbreviated title: KNDy neuron concept challenged

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43 **Abstract**

44 Previous immunohistochemical and *in situ* hybridization studies of sheep, goats and rodents
45 indicated that kisspeptin (KP), neurokinin B (NKB) and dynorphin A (DYN) are extensively
46 colocalized in the hypothalamic arcuate nucleus (ARC), thus providing a basis for the ‘KNDy
47 neuron’ concept; in both sexes, KNDy neuropeptides have been implicated in the generation of
48 gonadotropin releasing hormone (GnRH) neurosecretory pulses and in the negative feedback effects
49 of sexual steroids to the reproductive axis.

50 To test the validity and limitations of the KNDy neuron concept in the human, we carried out the
51 comparative immunohistochemical analysis of the three neuropeptides in the infundibular nucleus
52 (Inf=ARC) and stalk (InfS) of young male human individuals (<37 years).

53 Results of quantitative immunohistochemical experiments established that the regional densities of
54 NKB immunoreactive (IR) perikarya and fibers, and the incidence of afferent contacts they formed
55 onto GnRH neurons, were about 5 times as high as those of the KP-IR elements. Dual-
56 immunofluorescent studies confirmed that considerable subsets of the NKB-IR and KP-IR cell
57 bodies and fibers are separate and only about 33% of NKB-IR perikarya and 75% of KP-IR
58 perikarya were dual-labeled. Furthermore, very few DYN-IR cell bodies could be visualized in the
59 Inf. DYN-IR fibers were also rare, and with few exceptions, distinct from the KP-IR fibers. The
60 abundance and colocalization patterns of the three immunoreactivities showed similar trends in the
61 InfS around portal blood vessels.

62 Together, these results indicate that most NKB neurons in the Inf do not synthesize detectable
63 amounts of KP and DYN in young male human individuals. These data call for a critical use of the
64 ‘KNDy neuron’ terminology while referring to the putative pulse generator system of the
65 mediobasal hypothalamus. We conclude that the functional importance of these three neuropeptides
66 in reproductive regulation considerably varies among species, between sexes and at different ages.

67 **Introduction**

68 Information accumulated from immunohistochemical and *in situ* hybridization studies has recently
69 formed the basis for the kisspeptin/neurokinin B/dynorphin A ('KNDy') neuron concept and terminology
70 (1-4). As shown first for the sheep (5), many of these neurons with cell bodies located in the
71 hypothalamic arcuate nucleus (ARC; called infundibular nucleus in humans; Inf) co-synthesize kisspeptin
72 (KP), neurokinin B (NKB) and dynorphin A (DYN). They have been implicated in negative sex steroid
73 feedback to gonadotropin-releasing hormone (GnRH) neurons (5-7) and proposed to also serve as
74 pacemakers for the GnRH neurosecretory pulses (3, 4, 8, 9). Recent models of the GnRH pulse generator
75 (3, 4, 8) suggest that KNDy neurons communicate with one another via NKB and its receptor, NK3, and
76 possibly, also DYN and its receptor, KOR. In ovariectomized goats, central NKB increases and DYN
77 decreases the frequencies of multiunit activity volleys and LH secretory pulses (8). Pulse generator cells,
78 in turn, appear to communicate with GnRH neurons primarily via KP/KISS1R signaling. GnRH neurons
79 express KISS1R (10-12) and the majority of GnRH neurosecretory pulses show temporal association
80 with KP pulses in the median eminence of monkeys (13).

81 The general consensus that KP, NKB, DYN, NK3 and KOR are expressed by the same neurons relies
82 on combined neuroanatomical data from sheep (1, 5), rats (14), mice (3, 4, 15), goats (8), monkeys (16)
83 and humans (17, 18). However, closer analysis of these reports, in retrospect, reveals that neuropeptide
84 and receptor colocalizations are often only partial and also variable in the different studies, species, sexes
85 and age groups. Notably, in our recent immunohistochemical study of aged human individuals, we have
86 detected robust sex differences in the abundance of KP-IR (17, 18) and NKB-IR (18) neuronal elements
87 in the Inf. These studies have also revealed that the incidences of NKB-IR cell bodies, fibers and
88 appositions onto GnRH neurons exceed several-fold those of KP-IR elements, with particularly robust
89 differences in males (18). These results suggested that NKB-IR neurons and their fibers are partly
90 distinct from the KP-IR elements in these human models, thus challenging the universal validity of the

91 KNDy neuron concept. Moreover, in triple-immunofluorescent studies of aged human males, GnRH
92 neurons tended to receive primarily single-labeled afferent inputs from these peptidergic systems, with
93 KP/NKB double-labeled axons representing only 10.2% of all KP-IR afferents and NKB/KP double-
94 labeled axons about 8.8% of all NKB-IR afferents (18). From the above findings and preliminary
95 observations indicating that KP immunolabeling is even weaker in young than in aged men, we predicted
96 that the degree of overlap between the KNDy neuropeptides is much lower in young male humans than
97 suggested earlier for female sheep (1, 5), goats (8) or mice (3).

98 In the present study we investigated the universal validity of the KNDy neuron concept via the
99 parallel immunohistochemical analysis of NKB-, KP- and DYN immunoreactivities in the Inf and the
100 infundibular stalk (InfS) of young men. Specifically, i) we compared the immunoreactive perikaryon and
101 fiber densities in the Inf and the InfS, ii) we addressed the colocalization of KP with NKB and KP with
102 DYN in perikarya and fibers, and finally, iii) we compared quantitatively the incidences of NKB-IR vs.
103 KP-IR afferent contacts onto GnRH-IR neurons.

104

105 **Materials and methods**

106 *Human subjects*

107 Human hypothalamic samples were obtained from autopsies at the Forensic Medicine Department of
108 the University of Debrecen with permission from the Regional Committee of Science and Research
109 Ethics of the University of Debrecen (DEOEC RKEB/IKEB: 3183-2010). Selection criteria included
110 sudden causes of death, lack of history of neurological and endocrine disorders and *post mortem* delay
111 below 36h. Tissue specimens from six young male individuals (age 21-37 years) were used.

112 *Section preparation*

113 Following dissection, the hypothalamic tissue blocks were rinsed with running tap water and then,
114 immersion-fixed with 4% formaldehyde in 0.1M phosphate buffer saline (PBS; pH 7.4) for 7-14 days.
115 After fixation, the blocks were trimmed in a way to include the optic chiasma rostrally, the mammillary

116 bodies caudally and the anterior commissure dorsally (17-19). Bilateral sagittal cuts were placed 2cm
117 lateral from the midline. The blocks were bisected into right and left halves and then, infiltrated with
118 20% sucrose for 5 days at 4°C. The right hemihypothalami were sectioned coronally at 30µm with a
119 Leica SM 2000R freezing microtome (Leica Microsystems). All experiments were performed on every
120 24th hemihypothalamic section from each subject.

121 *Pretreatments*

122 The tissues were permeabilized and endogenous peroxidase activity reduced using a mixture of 0.2%
123 Triton X-100 and 0.5% H₂O₂ in PBS for 30 min. Antigen epitopes were unmasked by incubating sections
124 in 0.1M citrate buffer (pH 6.0) at 80 °C for 30 min (18). Dual-immunofluorescent experiments also used
125 a Sudan black pretreatment against autofluorescence (17, 18).

126 *Immunohistochemical detection of KP*

127 To detect KP immunoreactivity, sections were incubated in a sheep polyclonal antiserum against
128 human kisspeptin-54 (GQ2; 1:200,000). This antiserum recognizes human KP-54, KP-14 and KP-10,
129 exhibits less than 0.01% cross-reactivity *in vitro* with other related human RF amide peptides (20), and
130 was used successfully in previous immunohistochemical experiments on primate hypothalami (16-18,
131 21). Incubation in the primary antibodies for 48 h at 4C was followed by biotinylated secondary
132 antibodies (Jackson ImmunoResearch Laboratories, West Grove, PA, USA; 1:500) and the ABC Elite
133 reagent (Vector, Burlingame, CA; 1:1000) for 60 min each. The peroxidase reaction was visualized with
134 nickel-intensified diaminobenzidine chromogen (19) and then, post-intensified with silver-gold (22).

135 *Immunohistochemical detection of NKB and DYN*

136 Another two parallel series of sections were used to visualize NKB- and DYN, respectively. NKB
137 neurons were detected with rabbit polyclonal antibodies against the C-terminal 28 amino acids of human
138 NKB (IS-682, P. Ciofi; 1:100,000) (17), whereas DYN neurons were labeled with rabbit polyclonal
139 antibodies against amino acids 1-17 of porcine (and human) DYN (T-4268; Peninsula Laboratories; San
140 Carlos, CA; 1:100,000). The primary antibodies were reacted with biotinylated antirabbit IgG (Jackson

141 ImmunoResearch Laboratories; 1:500; 1h) and then, ABC Elite reagent (1:1000; 1h). The peroxidase
142 signal was developed with silver-gold-intensified nickel-diaminobenzidine. For control purposes,
143 mapping studies of DYN were replicated using a previously characterized second antiserum (IS-35;
144 1:200,000) against a different prodynorphin-derived peptide, dynorphin B (23).

145 *Dual-immunoperoxidase detection of KP and GnRH or NKB and GnRH*

146 Two section series were processed for the immunohistochemical detection of KP and NKB,
147 respectively, as above. Then, they were processed to detect GnRH using a guinea pig primary antiserum
148 (#1018; 1:5000) (18) which was reacted with biotinylated antiguinea pig IgG (Jackson ImmunoResearch
149 Laboratories; 1:500; 1h) and ABC Elite solution (Vector; 1:1000; 1h). The peroxidase signal was
150 visualized with brown diaminobenzidine.

151 *Fluorescent immunohistochemistry*

152 Other series of sections were processed for the dual-immunofluorescent studies of the colocalization
153 between NKB and KP or DYN and KP. Incubation in a cocktail of primary antibodies (rabbit anti-NKB,
154 1:1000 and sheep anti-KP, 1:1000; or rabbit anti-DYN, 1:1000 and sheep anti-KP, 1:1000; 48h; 4°C) was
155 followed by a cocktail of fluorochrom-conjugated secondary antibodies (Jackson ImmunoResearch; anti-
156 rabbit-FITC, 1:250; anti-sheep-Cy3, 1:1000) for 5h at room temperature.

157 To maximize sensitivity, other dual-immunofluorescent studies used tyramide signal amplification. In
158 these experiments KP was detected first using sequential incubations in KP antibodies (1:30,000; 48h;
159 4°C), biotinylated antgoat IgG (Jackson ImmunoResearch Laboratories; 1:500; 1h), the ABC Elite
160 reagent (Vector; 1:1000; 1h), biotin tyramide working solution (1:1000, in 0.05M Tris-HCl buffer, pH
161 7.6, containing 0.003% H₂O₂; 30 min) (24) and finally, avidin-Cy-3 (Jackson ImmunoResearch; 1:1000;
162 1h). Then, the sections were treated for 30 min with 0.5% H₂O₂ and 0.1% sodium azide in PBS, to
163 inactivate horseradish peroxidase. To detect NKB or DYN, the rabbit primary antibodies were used at
164 1:50,000 (48h; 4°C) and reacted with antirabbit-peroxidase (Jackson ImmunoResearch; 1:500; 1h). Then,
165 FITC-tyramide (24) (diluted 1:500 with 0.05M Tris-HCl buffer, pH 7.6, containing 0.003% H₂O₂; 30

166 min) was deposited on the peroxidase sites. Control experiments included the omission of the NKB and
167 DYN primary antibodies. Lack of FITC labeling in these control sections indicated that no FITC-
168 tyramide deposition is caused by residual peroxidase activity on KP-IR sites.

169 *Section mounting and coverslipping*

170 Following section mounting, immuno-peroxidase labeled sections were coverslipped with DPX (Fluka
171 Chemie; Buchs, Switzerland) and immunofluorescent specimens with Mowiol.

172 *Digital photography*

173 The light and fluorescent microscopic images were scanned with an AxioCam MRc 5 digital
174 camera mounted on a Zeiss AxioImager M1 microscope using the AxioVision 4.6 software (Carl
175 Zeiss, Göttingen, Germany). The light and fluorescent microscopic images were scanned with an
176 AxioCam MRc 5 digital camera mounted on a Zeiss AxioImager M1 microscope using the
177 AxioVision 4.6 software (Carl Zeiss, Göttingen, Germany). Confocal images were prepared with
178 an inverted Nikon Eclipse Ti-E microscope equipped with an A1R confocal system (Nikon,
179 Austria). The digital images were processed with the Adobe Photoshop CS software (Adobe Systems,
180 San José, CA, USA) at a 300 dpi resolution. Quantitative data were expressed as mean \pm SEM and
181 statistical comparisons used one-way ANOVA.

182 **Experiment 1. Studies of the incidences and overlaps of IR cell bodies in the Inf**

183 The number of immunoreactive cell bodies was counted at 100X magnification within a 0.25 mm²
184 counting area with the aid of a 5X5 ocular grid, as described previously (17, 18). Each subject was
185 characterized by the maximal number of immunoreactive perikarya in this counting area (determined
186 from 2-6 sections.)

187 The overlaps between NKB- and KP immunoreactivities and between DYN- and KP
188 immunoreactivities were first assessed in confocal images of dual-immunofluorescent specimens. The
189 percentages of single-labeled and double-labeled NKB-IR and KP-IR perikarya were also determined

190 quantitatively from the specimens in which the tyramide signal amplification was used, using 1-3
191 representative confocal images per subject.

192 **Experiment 2. Studies of the regional abundance of IR fibers in the Inf**

193 The regional density of immunoreactive fibers was determined as described recently (18). First,
194 digital images were taken from the bulk of kisspeptin-IR and NKB-IR neurons in the Inf. The files were
195 opened with the Adobe Photoshop CS software. The immunolabeled cell bodies were erased (“eraser
196 tool”) from the photomicrographs. The remaining images were compiled into TIF files and opened with
197 the Image J software (public domain at <http://rsbweb.nih.gov/ij/download.html>). The regional fiber
198 density in each photograph was defined as the area occupied by immunoreactive fibers/total area. For
199 each subject, the mean fiber density was derived from 1-3 digital images. The overlap between NKB-IR
200 and KP-IR axons or DYN-IR and KP-IR axons was also studied qualitatively in confocal images of dual-
201 immunofluorescent specimens.

202 **Experiment 3. Studies of immunoreactive fibers in the InfS**

203 Projections of NKB-, KP-, DYN- and GnRH-IR axons around the portal blood vessels of the InfS
204 were analyzed in this experiment. Based on previous immunohistochemical results in the median
205 eminence of different species (16, 23), we assumed that fibers containing KNDy peptides around the
206 portal vasculature arise mostly from the ARC/Inf. First, sections labeled with peroxidase-based
207 immunohistochemistry were used to study the relationship of fibers with the superficial and deep
208 capillary plexuses of the human postinfundibular eminence (25). Then, the extents of overlap between
209 NKB and KP immunoreactivities and between DYN and KP immunoreactivities were assessed from
210 dual-immunofluorescent specimens.

211 **Experiment 4. Studies of the incidences of KP-IR and NKB-IR appositions onto GnRH-IR neurons** 212 **of the Inf**

213 Dual-immunoperoxidase labeled sections were selected (1-2 from each individual) to determine the
214 number of axonal contacts along the outlines of GnRH-IR cell bodies and dendrites. Counting of the

215 appositions was carried out using a 63X oil-immersion objective and contacts defined using stringent
216 criteria (18, 26, 27). For each subject, the mean number of contacts per GnRH soma and 100 μ m GnRH
217 dendrite were calculated (18).

218

219 **Results**

220 The comparative analysis of NKB, KP and DYN immunoreactivities in
221 immunoperoxidase-labeled sections of the Inf (Fig. 1) and the InfS (Fig. 2) revealed
222 strikingly different labeling intensities for KNDy neuropeptides. In general, NKB-IR
223 elements showed much higher abundance than KP-IR elements. DYN immunoreactivity,
224 both in perikarya and fibers, was relatively sparse and weak.

225 *Experiment 1. Incidence of IR perikarya in the Inf*

226 In peroxidase-based immunohistochemistry, many NKB-IR perikarya were identified in the
227 Inf (Figs. 1A, B). KP-IR cell bodies occurred in much lower numbers in neighboring sections
228 (Figs. 1C, D). Quantitative analysis showed that the density of KP neurons was about 5 times
229 lower than that of NKB-IR perikarya (Graph 1; $P=0.01$ by ANOVA). DYN-IR perikarya were
230 either entirely absent in some subjects (Figs. 1E, F) or extremely rare in the Inf of others,
231 preventing quantitative studies. In contrast, the supraoptic nucleus contained many labeled
232 perikarya (Fig. 1G), making it unlikely that the low DYN signal in the Inf reflects technical
233 limitations.

234 A surprising segregation of NKB-IR and KP-IR perikarya was revealed in dual-
235 immunofluorescent specimens (Figs. 3A, B). Tyramide signal amplification was crucial for
236 sensitive detection of NKB/KP dual-labeled cell bodies which represented only $32.9\pm 4.7\%$ of the

237 NKB-IR and $75.2 \pm 6.6\%$ of the KP-IR perikarya (Figs. 3A, B). Tyramide signal amplification was
238 capable of visualizing only a few DYN-IR perikarya (not shown).

239 ***Experiment 2. Abundance of IR fibers in the Inf***

240 The incidence of immunolabeled fibers in the Inf followed a similar trend as labeled
241 perikarya. The most frequently encountered phenotype was, again, IR for NKB. These axons
242 established many appositions (Fig. 1B) to NKB-IR cell bodies and their dendritic processes.
243 Quantitative analysis of the area covered by immunohistochemical signal established that the
244 mean incidence of NKB-IR fibers was about 5 times as high as that of KP-IR fibers (Graph
245 2; $P=0.0001$ by ANOVA). DYN-IR fibers were also detectable in the Inf, although less
246 frequently than either NKB-IR or KP-IR axons (Figs. 1E, F).

247 In immunofluorescent specimens, many NKB-IR fibers without KP immunolabeling as
248 well as KP-IR fibers without NKB labeling could be seen in the Inf, in addition to dual-
249 labeled axons (Figs. 3A, B). DYN-IR fibers showed a high intensity of labeling only if the
250 tyramide signal amplification approach was also used. Most of them were distinct from KP-
251 IR axons, although dual-labeled KP/DYN-IR fibers occasionally occurred (Figs. 3F, G).
252 Similarly, the majority of KP-IR fibers were also devoid of dynorphin B immunoreactivity in
253 the Inf and the InfS (Figs. 3I, J), whereas this second antiserum also performed well in
254 regions rich in DYN fibers, including the ventromedial nucleus (Fig. 3K).

255 ***Experiment 3. Abundance of immunoreactive fibers in the InfS***

256 The InfS was associated with the superficial and the deep capillary plexuses of the
257 postinfundibular eminence (25). Both were abundantly innervated by GnRH-IR axons
258 (brown color in Figs. 2A-F), suggesting they contribute to the GnRH supply of

259 adenohipophysial gonadotropes. The relative abundance of the different types of labeled
260 fibers around the two capillary plexuses matched what was seen in the Inf. Accordingly,
261 portal blood vessels were surrounded by dense networks of NKB-IR fibers (Figs. 2A-C) and
262 innervated only moderately by KP-IR fibers (Figs. 2D-F). Very few DYN-IR fibers occurred
263 in the proximity of the portal capillaries (Figs. 2G, I, J). This low level of DYN signal did
264 not reflect a technical limitation, considering that the magnocellular neurosecretory tract was
265 immunolabeled heavily in the same sections (Fig. 2H).

266 The analysis of immunofluorescent specimens confirmed that NKB dominates over KP
267 around the portal vasculature and NKB-IR fibers often lack KP labeling (Figs. 3C-E). Similarly
268 to the Inf, the InfS contained both single-labeled and double-labeled KP-IR fibers (Figs. 3C-E).
269 In sections dual-labeled for KP and DYN, labeled fibers were mostly distinct, although rare
270 colocalization cases were also detectable (Fig. 3H).

271 *Experiment 4. Frequency of kisspeptin-IR and NKB-IR appositions onto GnRH-IR neurons*

272 Sections double-labeled with the silver-gold-intensified nickel-diaminobenzidine and
273 diaminobenzidine chromogens were used to obtain quantitative estimates about NKB-IR and KP-IR
274 inputs to GnRH-IR neurons. Microscopic analysis confirmed that NKB-IR (18) and KP-IR (17, 18) axons
275 provide axo-somatic and axo-dendritic inputs to GnRH neurons in the Inf (Fig. 4). Quantitative analysis
276 (Graph 3) established that GnRH-IR perikarya and dendrites, respectively, received 6 and 5 times heavier
277 NKB-IR (Fig. 4A) than KP-IR (Fig. 4B) innervation (GnRH perikarya: $P=0.004$; GnRH dendrites:
278 $P=0.005$, by ANOVA).

279 **Discussion**

280 Immunohistochemical results of this study provide evidence for limitations of the 'KNDy
281 neuron' terminology and concept. Specifically, our observations indicate that in young male humans

282 the majority of NKB-IR neurons in the Inf, their processes and contacts onto GnRH neurons do not
283 contain detectable amounts of KP immunoreactivity. Furthermore, KP-IR neuronal elements without
284 NKB labeling also occur frequently, in addition to NKB/KP dual-phenotype structures. Finally, we
285 observed that most KP-IR neurons and fibers are devoid of DYN immunoreactivity in this human
286 model.

287 **Species differences in the colocalization of KNDy peptides**

288 We propose that the different colocalization patterns in the present human study and in previous
289 animal experiments partly reflect species differences in reproductive mechanisms (28). These may
290 include the absence of DYN signal from most KP neurons and fibers in humans, which is in contrast
291 with the extensive coexpression in the rodent (3, 4, 14), sheep (5) and goat (8) ARC. Another putative
292 species difference is the large excess of NKB-IR over KP-IR perikarya in the present human study, as
293 opposed to the two-fold excess in the ARC of male mice (4).

294 **Sex-dependent variations in the absolute and relative abundances of KP and NKB and their** 295 **colocalization pattern**

296 In a previous study of KP-IR and NKB-IR neurons we identified a series of sex-dependent
297 morphological differences between aged human males and females (18). In particular, KP
298 immunoreactivity was highly sexually dimorphic; the number of KP-IR perikarya, the density of KP-IR
299 fibers and the incidence of KP-IR afferents on GnRH neurons were much higher in aged women
300 compared with men (18). It is worth of note that the NKB/KP neuron density ratio also differed, being
301 2.18 in aged males (>50 years) and 1.55 in postmenopausal females (>55 years) (18). In further support
302 of the idea that the KP/NKB colocalization pattern is sexually dimorphic, the ratio of dual-labeled NKB-
303 IR as well as KP-IR afferent contacts onto GnRH-IR neurons was significantly higher in postmenopausal
304 women (\approx 25-30%) than in aged men (\approx 8-10%) (18). We proposed that sex differences are either due to

305 organizational effects of sex steroids during critical period(s) of sexual differentiation or alternatively, to
306 the loss of negative sex steroid feedback in postmenopausal women, unlike in aged men who maintain
307 relative high testosterone levels. In our previous dual-immunofluorescent study we demonstrated a high
308 degree of overlap between the KP and NKB systems of postmenopausal women; above 80% of KP-IR
309 perikarya and NKB-IR perikarya contained also the other neuropeptide (17). The degree of this
310 colocalization is likely to be sexually dimorphic and much lower in aged males whose Inf contained over
311 twice as many NKB-IR as KP-IR perikarya (18).

312 It is worthy of note that the sexual dimorphism of KP and NKB neurons in the ARC/Inf region is not
313 unique to humans. The ARC of the sheep contains higher NKB (29) and KP (1) cell numbers in females
314 than in males. In rats, sex differences were reported in the projection fields of NKB-IR axons within the
315 infundibular area (23).

316 **Ageing-dependent variations in the absolute and relative abundances of KP and NKB**

317 **The human NKB and KP systems, at least in the male, also exhibit robust ageing-dependent changes.**
318 **Our preliminary data indicate that the enhancements of the absolute NKB and KP cell numbers coincide**
319 **with a significantly enhanced percentage of KP-expressing NKB neurons in aged men (Molnár et al., in**
320 **preparation).**

321 **Role of NKB in the regulation of the human reproductive axis**

322 The tachykinin peptide NKB plays a crucial role in human reproduction and inactivating mutations of
323 the NKB and NK3 encoding genes cause normosmic hypogonadotropic hypogonadism (30, 31).
324 Although the first reports did not indicate fertility deficits in the NK3 mutant mice (32), later analysis
325 focusing on the reproductive phenotype noticed subfertility (33), suggesting functional similarities in
326 NKB/NK3 signaling between the human and mouse species.

327 Out of the three KNDy peptides, NKB provided the heaviest immunohistochemical signal in the Inf
328 and the InfS of young men (present report), aged men (18) and aged women (18). In immunoperoxidase-
329 based studies, the dominance of NKB over KP was highest in young men (present study) where the
330 density of NKB-IR perikarya and fibers were about 5 times as high as those of the KP-IR elements
331 and NKB-IR axons established about 6-times as many axo-somatic and 5-times as many axo-dendritic
332 contacts onto GnRH neurons as did KP-IR axons.

333 Conflicting results of previous experiments suggest that the net effect of NKB on LH secretion
334 depends on animal species and endocrine paradigms. Intraperitoneal or intracerebroventricular NKB
335 administration to male mice had no effect on serum LH (34), whereas the intracerebroventricular
336 injection of the selective NK3 agonist senktide reduced LH secretion in ovariectomized rats treated with
337 a low dose of estradiol (35). Reduced LH secretion in response to senktide was also observed in
338 ovariectomized and in ovariectomized and estradiol treated rats (36) and in ovariectomized mice (3),
339 whereas another study on rats found stimulatory effect on LH secretion in the presence of physiological
340 levels of estradiol (37). Senktide also stimulated LH secretion in castrated male monkeys (16), and in the
341 follicular, but not in the luteal, phase in ewes (38), whereas reduced net LH secretion was observed in
342 ovariectomized goats (8).

343 Multiunit activity recorded in the ARC is considered to be an electrophysiological correlate of the
344 GnRH pulse generator activity. The coordinated bursts of neuronal firing occur in synchrony with the LH
345 secretory pulses in various animal species (8, 39), including primates (40). Senktide dose-dependently
346 suppressed the frequency of pulsatile LH secretion and inhibited hypothalamic multiunit activity volleys
347 in ovariectomized rats, independently of gonadal steroid levels (36). In contrast, a robust increase in the
348 frequency of multiunit activity volleys was observed in ovariectomized goats (8).

349 NKB mainly acts upstream from GnRH neurons. For example, the stimulatory effect of intravenous
350 NKB and senktide on LH secretion could be abolished by the GnRH receptor antagonist acyline (16).

351 One major target site for NKB actions appears to be on NKB (KNDy) neurons of the ARC/Inf.
352 Accordingly, i) we found numerous NKB-IR afferent contacts on human NKB neurons, ii) similar
353 contacts were reported previously in rats (14) and sheep (5, 41), iii) NK3 receptors are present on these
354 cells in rodents and sheep (3, 14, 42) and iv) KP (KNDy) neurons in the ARC of male mice respond with
355 c-Fos expression (4) and depolarization (4) to senktide.

356 There is little consensus regarding the possibility that NKB also influences GnRH neurons directly.
357 In sheep, GnRH neurons do not express NK3 immunoreactivity (42). In mice, while single-cell
358 microarray and RT-PCR studies provided proof for NK3 mRNA expression in GnRH neurons (43), *in*
359 *situ* hybridization studies were unable to confirm this finding (4). Also, senktide did not activate mouse
360 KNDy neurons *in vitro* (4). In rats, while immunohistochemical studies found evidence for NK3
361 immunoreactivity in only 16% of GnRH-IR cell bodies (44), the receptor was more abundant on GnRH-
362 IR axon terminals in the median eminence (44) where frequent appositions between GnRH-IR and NKB-
363 IR axons occurred (23, 44). Although NKB in itself did not alter GnRH release from hypothalamic
364 explants of male rats, it abrogated the KP-induced release of GnRH, suggesting a complex mode of
365 action which is likely parallel with, and not upstream from, the KP action (45). Recent functional
366 evidence from KISS1R-KO mice indicates that intact KP/KISS1R signaling is required for the
367 suppression of LH secretion by senktide. This finding provides support for the concept that the dominant
368 action of NKB is upstream from KP neurons, instead of being exerted directly on GnRH cells (45).

369 Results of the present and a previous (18) human study indicate that NKB-IR axons abundantly
370 innervate human GnRH neurons and the incidence of these contacts is several times as high as those of
371 KP-IR axons. It will require clarification whether this anatomical pathway uses NKB/NK3 signaling.
372 Alternatively, neurotransmitter (s) other than NKB may act in this communication, which is not likely to
373 be KP or DYN, in view of their relative paucity in young male individuals. In addition to innervating
374 GnRH-IR cell bodies and dendrites, NKB-IR axons also represented the most abundant KNDy peptide

375 around the portal capillary plexuses of the human postinfundibular eminence (25). This hypothalamic site
376 that lies outside the blood-brain barrier may represent an important site of interaction between NKB-IR
377 and GnRH-IR axons. It also remains possible that NKB is released into the hypophysial portal circulation
378 to influence adeno-hypophysial functions. *In vitro* evidence from rats, indeed, indicates that NKB
379 can induce prolactin secretion from perfused pituitary cells (46).

380 **Role of KP in the regulation of the human reproductive axis**

381 In humans, KP/KISSR1 signaling plays a pivotal role in reproduction. Loss of function mutations of
382 the genes encoding KISSR1 (47-49) and KP (50) result in hypogonadotropic hypogonadism. In recent
383 models of the GnRH pulse generator, KP was proposed to provide the main output signal of the pulse
384 generator neuronal network toward GnRH neurons, whereas NKB and DYN seem to primarily account
385 for the intranuclear communication of KNDy neurons via acting on NK3 and KOR, respectively (3, 8). It
386 is generally believed that independently from the species, KP acts directly on GnRH neurons that express
387 KISS1R mRNA (10-12) and in mice, respond with depolarization to KP (11, 51, 52).

388 As in laboratory and domestic animals, KP increases LH secretion in men (20, 53, 54) and women
389 (55, 56), most potently during the preovulatory phase of the menstrual cycle in the latter. It is interesting
390 to note that the continuous intravenous infusion of KP enhanced the LH pulse frequency in men (53),
391 indicating that KP not only acts on GnRH neurons, but also upstream from the pulse generator network.
392 This finding suggests a species difference from the mouse in which KNDy neurons do not appear to
393 synthesize KISS1R (57) and express only NK3 (3) and KOR (3) mRNAs.

394 In humans, axo-somatic, axo-dendritic and axo-axonal contacts may serve as communication
395 pathways between KP and GnRH neurons (17). In previous studies we showed that the KP system of
396 aged human individuals exhibits a robust sexual dimorphism (18) with postmenopausal women having
397 several times higher densities of KP-IR cell bodies and fibers in the Inf and higher incidences of KP-IR

398 afferent contacts onto GnRH neurons, than aged men above 50 years. Preliminary data that KP
399 immunoreactivity is even much lower in the hypothalamus of young men (Molnár et al., in preparation)
400 led us to carry out the present study to challenge the validity of the KNDy neuron terminology and
401 concept for the young male human model. Our present immunohistochemical data indicate that the
402 density of KP-IR perikarya and fibers in the Inf were about 5 times lower than those of the
403 corresponding NKB-IR elements and the number of KP-IR appositions to GnRH-IR cell bodies and
404 dendrites only reached about one-fifth and one-sixth, respectively, of those established by NKB-IR
405 axons. The functional consequences of the surprisingly low level of KP immunoreactivity in young
406 men (in somata as well as fibers) requires clarification. In view that KP is thought to represent the
407 main neurotransmitter output of the pulse generator system (3, 8), its low level in the mediobasal
408 hypothalamus of young men is compatible with only a moderate stimulation of GnRH/LH secretion.
409 Much higher raw numbers of the KP-IR cell bodies, fibers and contacts onto GnRH neurons in aged
410 male human subjects (18) and their highest incidence in postmenopausal women (18) are in
411 accordance with the idea that KP immunoreactivity and serum LH levels are linked. A future
412 challenge will be to correlate the immunohistochemical images of KP and the other KNDy peptides
413 with the GnRH neurosecretory output at the different human age and sex groups.

414 **Absence of DYN immunoreactivity from most KP neurons and their fiber projections**

415 The opioid peptide DYN is rather ubiquitous and may have multiple sites of action upon the
416 reproductive axis. These sites are likely upstream from the GnRH neuron that does not appear to express
417 KOR in rats (58). DYN is critically involved in progesterone negative feedback to GnRH neurons in
418 ewes; the majority of DYN cells in the ARC of ovariectomized ewes contain progesterone receptor (59)
419 and progesterone treatment increases preprodynorphin mRNA expression in the ARC and DYN levels in
420 the cerebrospinal fluid (60).

421 DYN is an important regulator of the pulse generator system. In sheep, KOR antagonists stimulate
422 the episodic secretion of LH during the luteal phase (61). In ovariectomized goats, central administration
423 of DYN decreases and KOR antagonist increases the frequencies of the multiunit activity volleys and of
424 the LH secretory pulses (8). Opioid peptides also regulate negatively the pulsatile release of prolactin and
425 LH in humans; this inhibitory tone can be suspended by the blockade of opioid receptors with naloxone
426 (62, 63).

427 The concept and terminology of the 'KNDy neurons' rely on the similar results of colocalization
428 experiments from several animal species. DYN has been detected in NKB (and/or KP) neurons in the
429 ARC of sheep (5, 41), mice (3, 4), rats (14, 23) and goats (8). Moreover, the DYN receptor KOR is
430 present in subsets of KNDy neurons in the ARC of mice (3, 4). Our present immunohistochemical study
431 to address the presence of DYN immunoreactivity in human KP neurons was also informed by previous
432 reports in which preprodynorphin mRNA expression was detected in the human Inf (64) and the monkey
433 ARC (65).

434 The somewhat unexpected absence of DYN immunoreactivity in most KP-IR somata and fibers of
435 young male humans questions the universal importance of DYN peptides within NKB and KP neurons of
436 the Inf and reveals an important difference from the rodent, sheep and goat species (1-5, 8, 23). It is
437 worthy to note that species also vary considerably regarding the sex steroid regulation of DYN in the
438 ARC/Inf. Preprodynorphin expressing neurons showed reduced numbers in postmenopausal women (64)
439 and in ovariectomized ewes (60), whereas there was no change in mRNA expression in postmenopausal
440 monkeys (65), whereas preprodynorphin mRNA was increased in the absence of sex steroids in mice (3).

441 The absence of DYN immunoreactivity from most KP-IR neurons and their fibers we report in this
442 study is unlikely to be entirely caused by the limited sensitivity of the applied immunohistochemical
443 method because i) DYN-IR cell bodies (e.g. magnocellular perikarya in the supraoptic nucleus) and fibers
444 (e.g. a dense fiber plexus in the ventromedial nucleus) were readily detectable elsewhere in the

445 hypothalamus, ii) substantial colocalization with KP was also undetectable using the highly sensitive
446 tyramide signal amplification method to visualize DYN or iii) using an antiserum against a different
447 prodynorphin cleavage product, dynorphin B.

448 **Summary of neuroanatomical findings**

449 i) The regional density of NKB-IR cell bodies, fibers and contacts onto GnRH neurons exceed about
450 five-fold those of KP-IR neuronal elements in the Inf.

451 ii) In addition to NKB-IR cell bodies and processes (in both the Inf and the InfS) that are devoid of
452 KP labeling, KP-IR elements lacking NKB immunoreactivity are also highly abundant, as established in
453 dual-immunofluorescent studies. In this study only $32.9\pm 4.7\%$ of the NKB-IR and $75.2\pm 6.6\%$ of the
454 KP-IR perikarya were dual-labeled.

455 iii) DYN-IR cell bodies and fibers occur much less frequently than either NKB-IR or KP-IR
456 elements; KP-IR axons in the Inf and the InfS contain DYN immunoreactivity only occasionally.

457 In conclusion, the immunohistochemical observations we made on hypothalamic tissue samples of
458 young male human subjects question the universal validity of the KNDy neuron concept and terminology
459 and suggest that the abundance of these peptides and their overlap are species-, sex- and age-dependent.

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470 **Legends**

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472 **Figure 1. Relative abundances of NKB-, KP- and DYN-IR cell bodies and fibers in the Inf of young**473 **male humans.** The silver-gold intensified nickel-diaminobenzidine chromogen was used to visualize474 NKB (**A, B**), KP (**C, D**) and DYN (**E-G**) immunoreactivities in adjacent sections of the Inf from a 31-475 year-old male subject. NKB-IR perikarya as well as nerve fibers (**A, B**) occur in much higher numbers476 than do KP-IR elements (**C, D**). For results of the quantitative analyses which reveal about five-fold

477 differences for both perikaryon and fiber densities, see Graphs 1 and 2, respectively. Arrowheads point to

478 NKB-IR cell bodies in **B** and a KP-IR cell body in **D**. Note that NKB-IR neurons receive numerous479 afferent contacts (arrows in **B**) from NKB-IR varicose axons; analogous juxtapositions in other species

480 were proposed to underlie the main peptidergic signaling mechanism among the putative pulse-

481 generating KNDy neurons. Few if any DYN-IR cell bodies are detectable in the Inf (none visible in this

482 specific case) and only scattered DYN-IR fibers occur (**E, F**). This low level of the DYN signal does not

483 appear to reflect a technical limitation of the immunohistochemical approach, given that IR perikarya and

484 fibers are abundant elsewhere in the hypothalamus, including the supraoptic nucleus (SO; **G**). Scale485 bar=100 μ m for **A, C, E** and 9 μ m elsewhere.

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489 **Figure 2. Results of immuno-peroxidase studies illustrating the differential innervation of the**
490 **portal capillary plexus by NKB-, KP- and DYN-IR fibers in a 31 year old men. A-I:** The black
491 silver-gold intensified nickel-diaminobenzidine chromogen was used to detect NKB (**A-C**), KP (**D-F**)
492 and DYN (**G-J**) immunoreactivities in adjacent sections of the InfS. Note that GnRH has also been
493 visualized in **A-F** with brown diaminobenzidine. The postinfundibular eminence with its deep (**B, E, I**)
494 and superficial (**C, F, J**) plexuses of portal blood vessels (BV) is surrounded by GnRH-IR
495 hypophysiotropic axons (brown color in **A-F**). Out of the axons immunoreactive for the three KNDy
496 peptides (black color), those with NKB immunoreactivity represent the most frequently encountered
497 phenotype (**A**) and densely innervate both the deep (**B**) and the superficial (**C**) portal capillaries; this
498 innervation raises the possibility of NKB release into the hypophysial portal circulation. The KP-IR
499 innervation of the portal BVs is of much lower density (**D-F**). Although DYN-IR axons are readily
500 detectable in the InfS (**G**) and contribute to the magnocellular axon tract (**H**), they only occur rarely
501 around the portal BVs (**H-I**). Scale bar=100 μ m for **A, D, G** and 10 μ m elsewhere.

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504 **Figure 3. Results of immunofluorescent studies revealing a significant degree of mismatch between**
505 **KP and NKB immunoreactivities (A-E) and KP and DYN (or dynorphin B) immunoreactivities (F-**
506 **K) in the Inf (A, B, F, G) and the InfS (C-E, H).** The dual-immunofluorescent visualization of NKB
507 (green) and KP (red) immunoreactivities in the Inf (**A, B**) not only confirms the dominance of NKB-IR
508 (green) over KP-IR (red) cell bodies (arrowheads) and axons (arrows) in the Inf, but also reveals a
509 considerable degree of segregation between the two different perikaryon (arrowheads) and fiber (arrows)
510 populations. Yellow double-arrows and arrowheads point to dual-labeled fibers and cell bodies,
511 respectively. Single-labeled NKB-IR axons are also typical around the portal blood vessels (BV) of the
512 postinfundibular eminence in the InfS (**C-E**), in addition to single-labeled KP-IR fibers (red) and
513 NKB/KP-dual-labeled (yellow) axons. The dual-immunofluorescent visualization of KP and DYN (**F-H**)
514 illustrates the absence of DYN immunoreactivity from KP-IR cell bodies in the Inf (red arrowheads in **F**).
515 With the exceptions of a few scattered dual-labeled fibers (yellow double-arrows) in the Inf (**F, G**) and
516 the InfS (**H**), the majority of KP-IR (red) and DYN-IR (green) axons are separate. **Negative**
517 **colocalization results are reproducible in the Inf (I) and the InfS (J) using an antiserum against a different**
518 **prodynorphin cleavage product, dynorphin B. Note the high density of dynorphin B-IR fibers in the**
519 **neighboring ventromedial nucleus (VMH) in K. High-power micrographs (A, B, D-K) represent single**
520 **optical slices (0.7 μ m). Scale bar=100 μ m for C, 5 μ m for D and 10 μ m for A, B, E-K.**

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525 **Figure 4. Different incidences of NKB-IR and KP-IR afferent contacts onto GnRH-IR neurons.**

526 NKB-IR axons (black color in **A**) show a much higher abundance in the Inf and establish considerably
527 more axo-somatic and axo-dendritic juxtapositions (arrows) onto GnRH-IR neurons (brown color) than
528 do KP-IR axons (black color in **B**). For quantification of these results, see Graph 3. Scale bar=20µm.

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Graph 1. Regional abundance of NKB-IR and KP-IR neuronal perikarya in the Inf of young male humans. The maximal number of immunoreactive cell bodies per 0.25mm² counting frame (1-6 per subject) was determined with the aid of an ocular frame and used as the index of the density of labeled perikarya. Results show that in young male human individuals, the mean incidence of NKB-IR cell bodies is about 5-times as high as the incidence of KP-IR cell bodies. *P<0.05 by ANOVA.

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541 **Graph 2. Density of NKB-IR and KP-IR fibers in the Inf of young men.** The area covered by
542 immunoreactive fibers (divided by the total area analyzed) was determined with the ImageJ software in
543 digital photographs of the Inf and used as a fiber density measure. Areas that were occupied by labeled
544 cell bodies and their thick proximal dendrites were deleted from the photographs with the eraser tool of
545 the Adobe Photoshop software and thus, excluded from the analysis. The density of NKB-IR fibers
546 (expressed in arbitrary units) is 5.4 times as high as the density of KP-IR fibers. *P<0.0005 by ANOVA.

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550 **Graph 3. Incidences of NKB-IR and KP-IR contacts onto GnRH-IR neurons in the Inf in young**
551 **men**

552 High-power light microscopic analysis of sections, dual-immunolabeled with the combined use of silver-
553 gold-intensified nickel-diaminobenzidine and diaminobenzidine chromogens, was carried out to
554 determine the relative incidences of NKB-IR and KP-IR neuronal appositions to the somata (left
555 columns) and the dendrites (right columns) of GnRH-IR neurons. The counts were obtained from all
556 GnRH-IR cell bodies and dendrites identified in 1-3 infundibular sections of each individual. The number
557 of NKB-IR contacts is about 6-times as high on the somatic and 5-times as high on the dendritic
558 compartment of GnRH neurons as those established by KP-IR axons. *P<0.01 by ANOVA.

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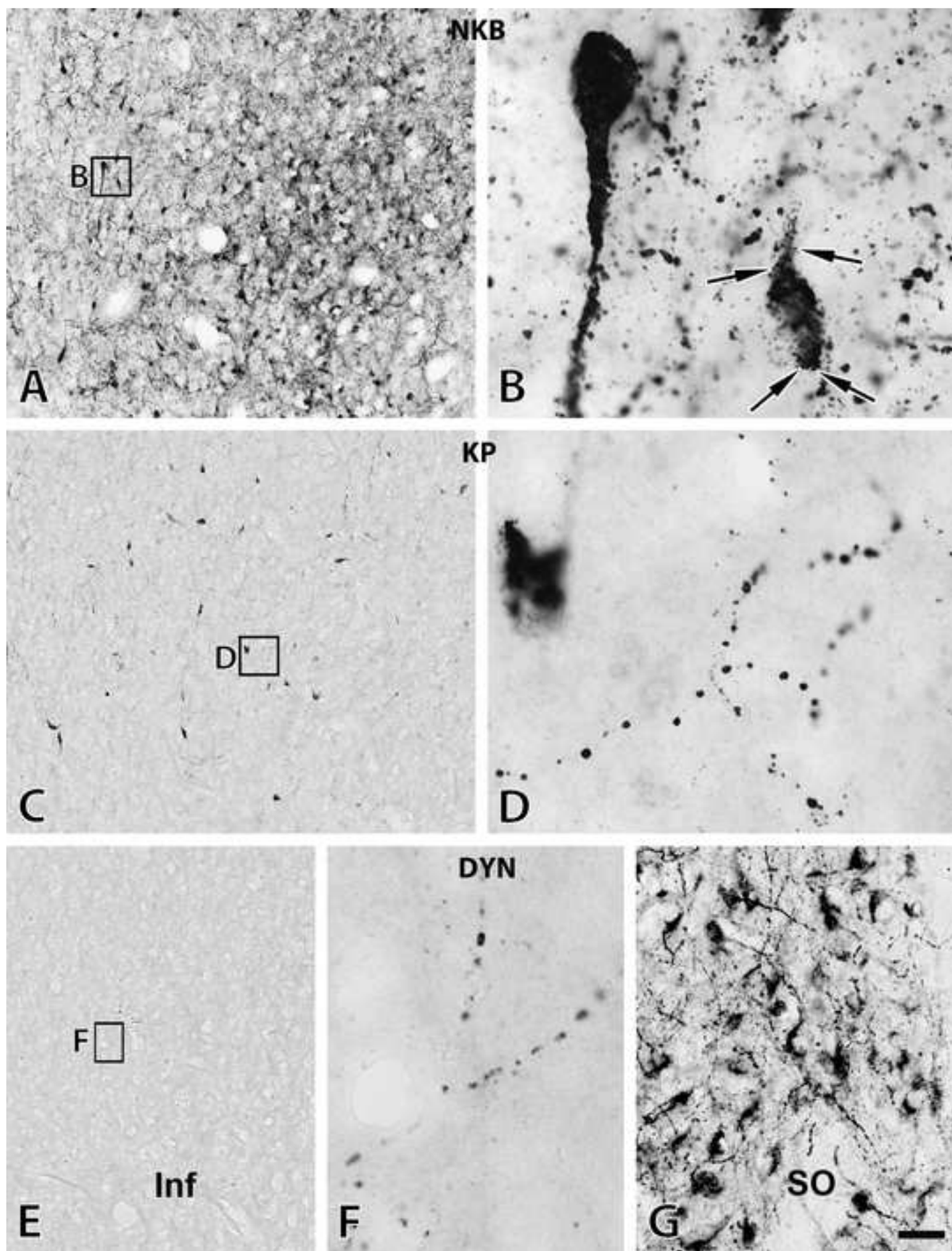


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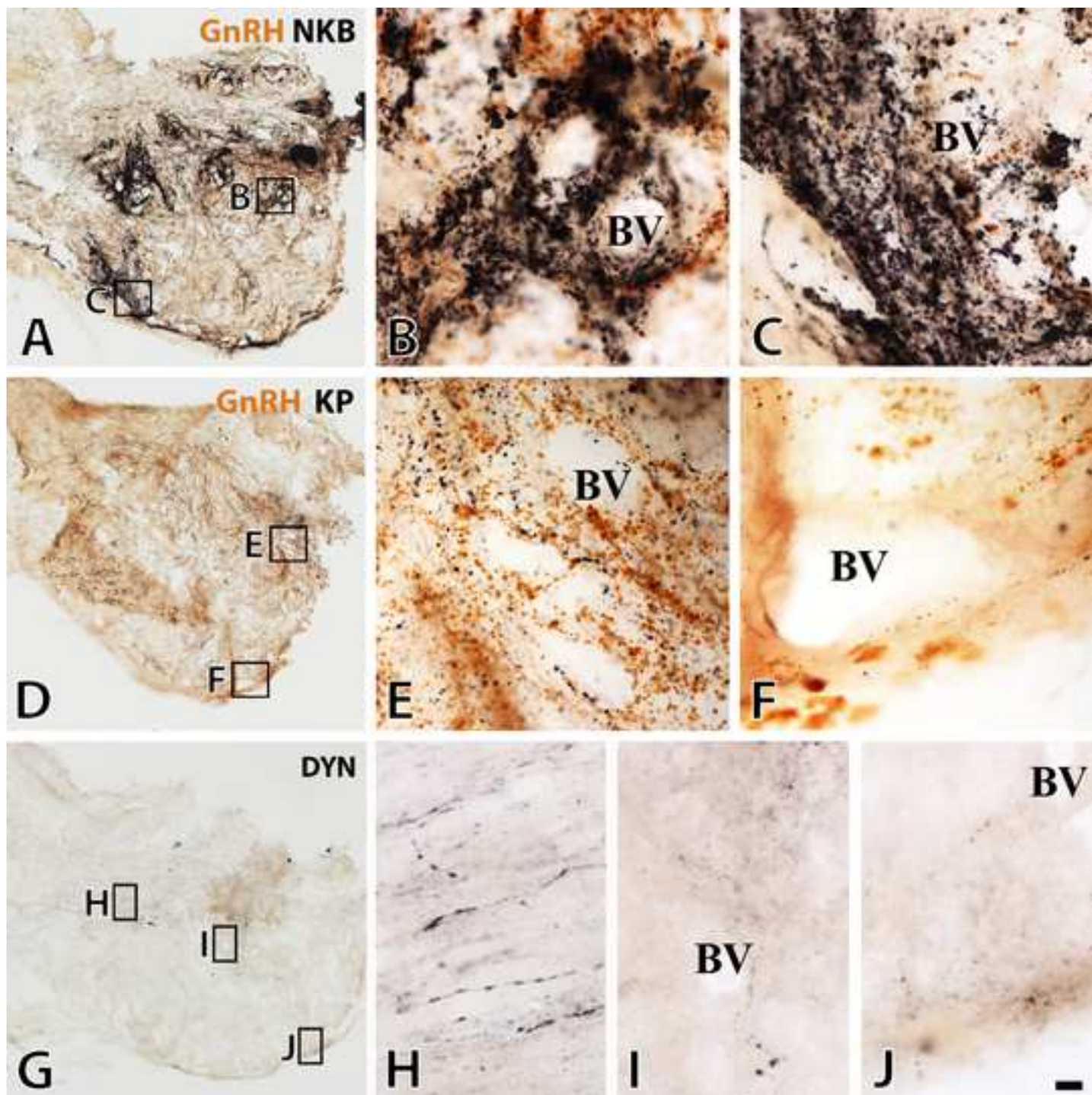
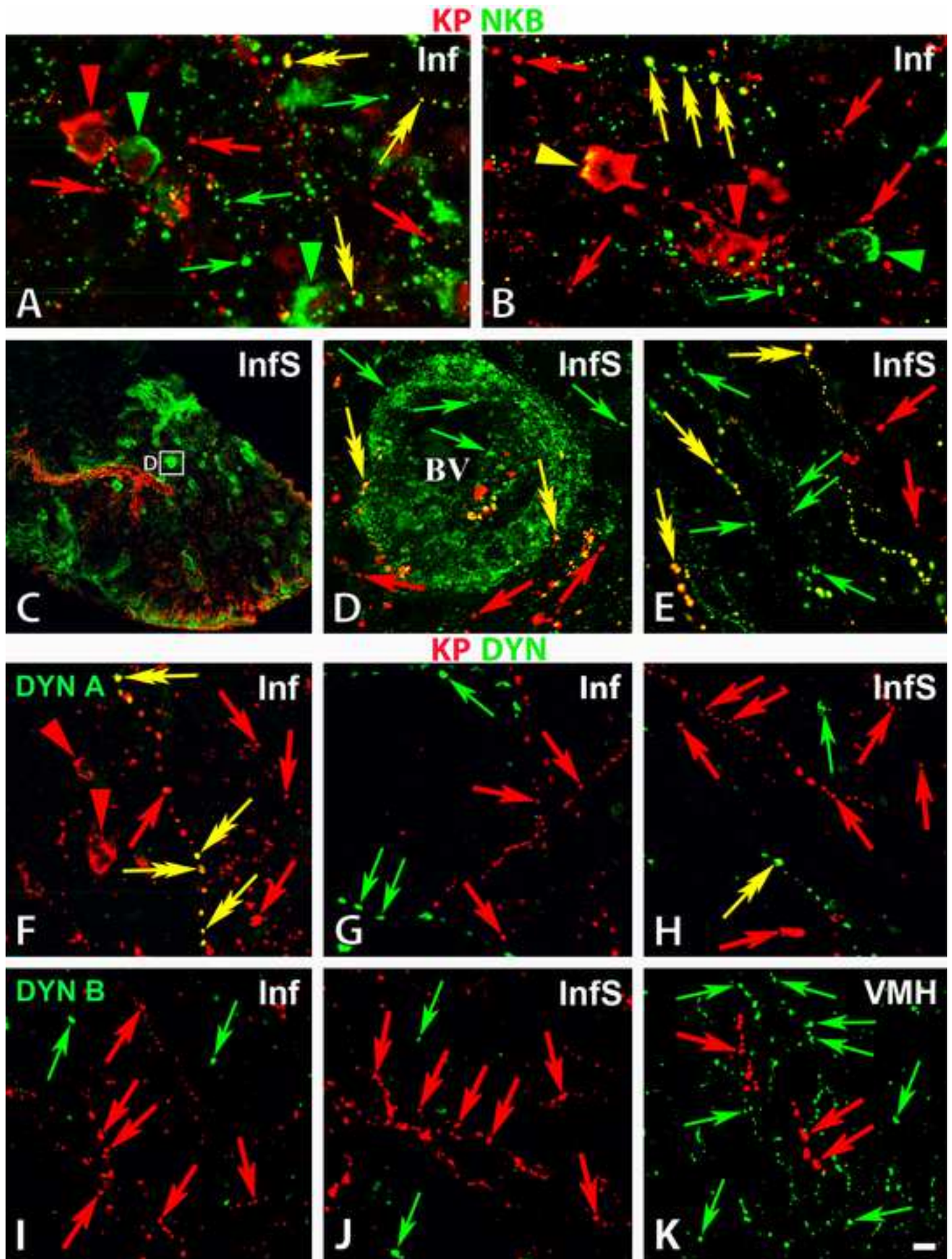
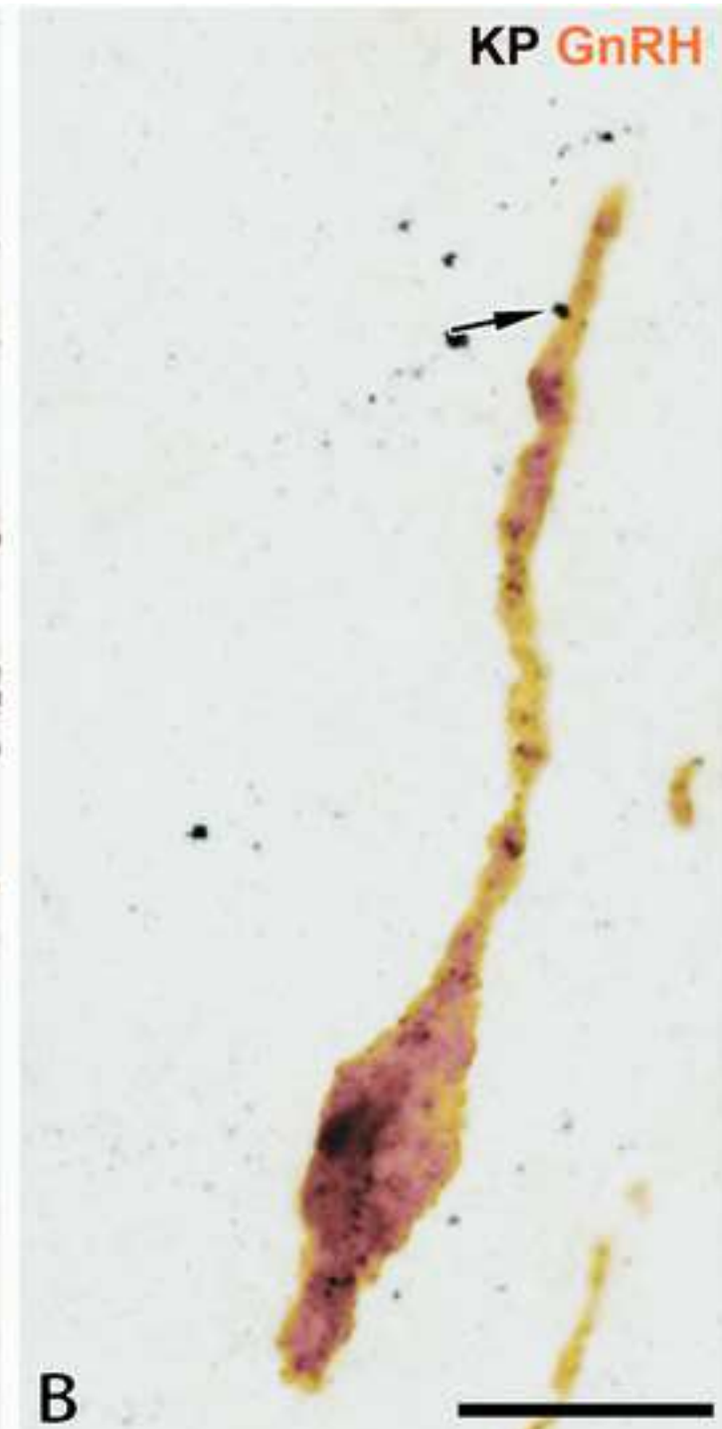
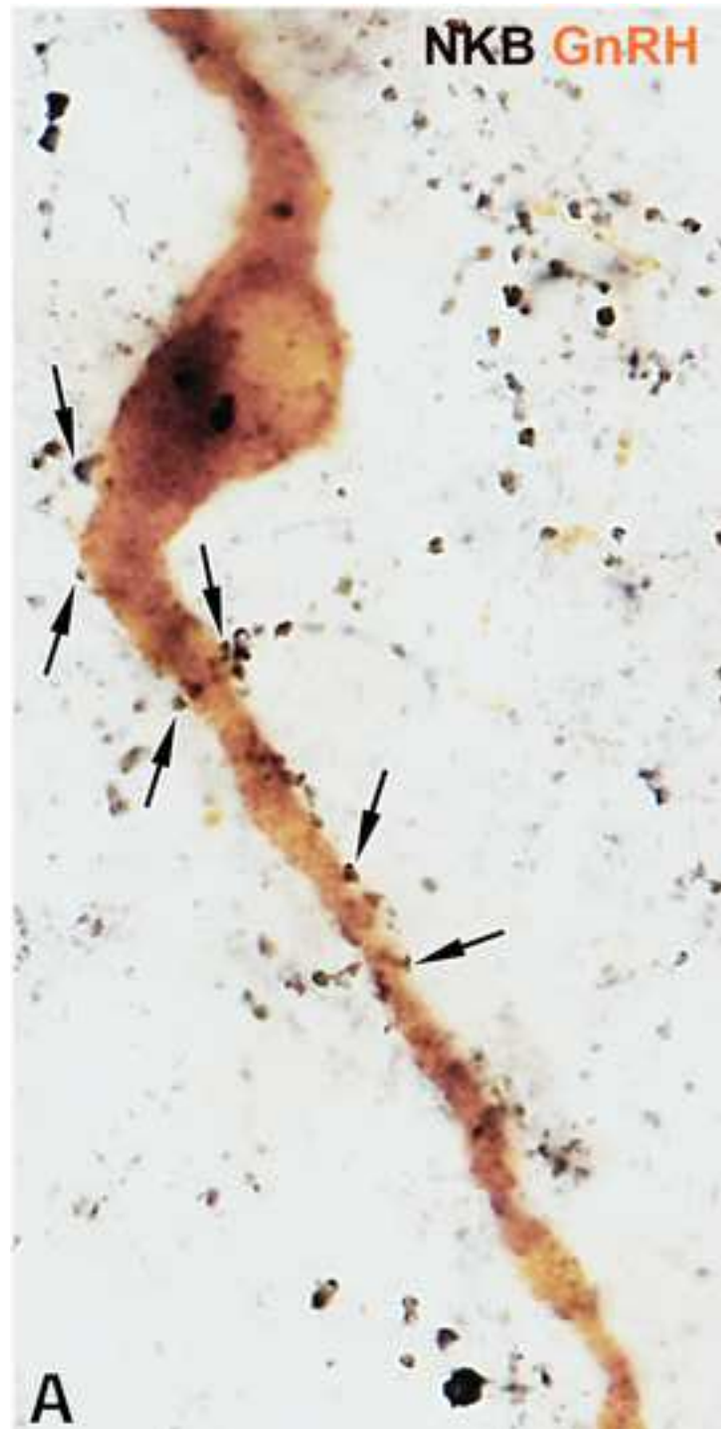


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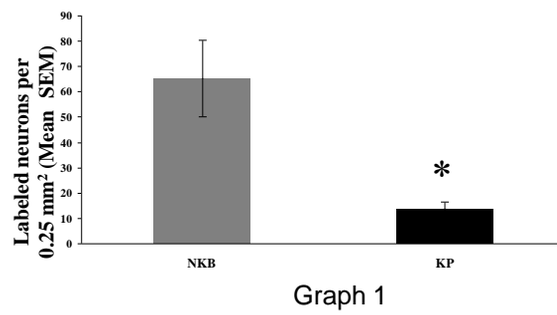
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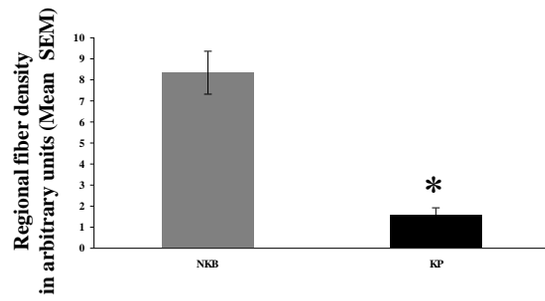
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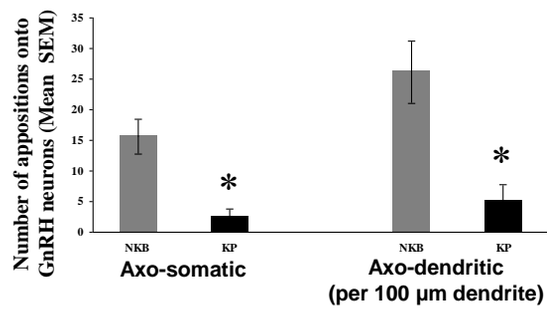


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Graph 2



Graph 3