

Erratum and Corrigendum

FEBS 21323

Erratum to: **Distribution of orexin receptor mRNA in the rat brain** (FEBS 21024)

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In the above-mentioned publication the second author was not given the credit for having contributed equally to the study with the first author. The publisher apologizes for this error.

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Corrigendum to: **Trp proteins form store-operated cation channels in human vascular endothelial cells** (FEBS 20791)

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The authors wish to correct the legend of Fig. 1, which is given below.

Fig. 1. HUVEC express multiple Trp isoforms. Trp genes (Trp1, 3 and 4, as indicated) were amplified by PCR, blotted and probed with digoxigenin-labeled oligonucleotides designed to bind to sequences internal to the respective PCR primers. For amplification of Trp4, two different sets of primers were used. One set was designed according to the sequence of mTrp4 (Trp4), the other set was based on a published hTrp4 sequence (Trp4*) [12]. PCR amplification was performed with reverse transcribed RNA from HUVEC and HEK 293 as well as with RNA (from HUVEC) which was not subjected to reverse transcription (negative control) and with the three available cDNA clones (hTrp1, hTrp3, and mTrp4) as template (positive control). The expected size of the PCR products is indicated.

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