

Listado de genes

ABCA1¹, ABCB1¹, ABCG1¹, ADCYAP1¹, ADORA1¹, ADORA2A¹, ADORA3¹, ADRA2A¹, ADRB2¹, AGTR2¹, ALOX12¹, AQP4¹, AR¹, ARNTL¹, ARRB2¹, ASIC1¹, ASIC3¹, ATAT1¹, ATF4¹, ATL1², ATL3³, AVIL¹, BACE1¹, BAX¹, BDKRB1¹, BDKRB2¹, BDNF¹, BEGAIN¹, BHLHA9¹, CACNA1B¹, CACNA2D1¹, CACNA2D3¹, CACNA2D3¹, CACNA2D3¹, CACNB3¹, CACNG2¹, CALCA¹, CAMK2A¹, CAMKIV¹, CAPN1¹, CCKBR¹, CCR2¹, CDK5R1¹, CDKL5¹, CHRM2¹, CLCN6¹, CNR1¹, CNR2¹, COMT¹, COX1¹, COX2¹, CRIP2¹, CSF1¹, CSF2¹, CTSB¹, CTSS¹, CYP19A1¹, CYP2D6⁴, CYP2D19⁵, DAN¹, DAO¹, DBH¹, DDO¹, DICER1¹, DLG2¹, DMPK¹, DPP4¹, EAR2¹, EDN1¹, EDNRB¹, EFN1¹, EFN2¹, EGR1¹, EHMT2¹, EIF2¹, EIF2AK2¹, EIF2AK3¹, EIF4E¹, EIF4EBP1¹, ESR1¹, F2RL1¹, FAAH¹, FKBP5¹, FPR1¹, GABBR1¹, GABRG2¹, GAL¹, GALR1¹, GALR2¹, GCH1¹, GCH1¹, GFRA2¹, GNA11¹, GNAO1¹, GNAQ¹, GNAZ¹, GPR34¹, GPR55¹, GRIK1¹, GRIN1¹, GRIN2A¹, GRIN2B¹, GRK2¹, GRK3¹, GRK5¹, GRM1¹, GRM5¹, HAMLET¹, HCN2¹, HDC¹, HMOX2¹, HOXB8¹, HRH1¹, HRH2¹, IAPP¹, ICA1¹, IFNGR1¹, IL10¹, IL10R1¹, IL1A¹, IL1B¹, IL1R1¹, IL23¹, IL23R¹, IL4¹, IL6¹, IL6ST¹, KCNA2¹, KCNJ3¹, KCNJ6¹, KCNK2¹, KCNS1¹, KLF11¹, KIF1A⁶, LEP¹, LMX1B¹, LPAR1¹, LPAR5¹, LRP1¹, MAPK1¹, MAPK10¹, MAPK3¹, MAPK8¹, MAPK9¹, MC1R¹, MDK¹, MECP2¹, MKK3¹, MKK6¹, MME¹, MMP1³, MMP24¹, MMP9¹, MNK1¹, MNK2¹, MPZ¹, MRGPRE¹, MTA1¹, NDN¹, NF1¹, NGF¹, NGFR¹, NGR2¹, NLGN2¹, NOS1¹, NOS2¹, NPTX1¹, NPY¹, NPY1R¹, NRG1¹, NTRK1¹, NTSR1¹, NTSR2¹, OPRD1¹, OPRK1¹, OPRM1¹, OSM¹, P2RX3¹, P2RX7¹, P2RY12¹, PANX1¹, PDE10A¹, PDYN¹, PENK1¹, PER2¹, PIP5K1C¹, PIRT¹, PLCB3¹, PLP1¹, PNOC¹, POMC¹, PPARA¹, PPP1R14C¹, PRDM12¹, PRKAR1B¹, PRKCG¹, PRKG1¹, PRLR¹, PROK2¹, PROKR1¹, PRRX1¹, PRX¹, PTGER1¹, PTGER3¹, PTGIR1¹, PTN¹, PTPN5¹, RAG1¹, REST¹, RGS9¹, RLUA-1¹, RLUA-2¹, RUNX1¹, S100A10¹, SARM1¹, SCH1¹, SCN10A¹, SCN11A¹, SCN3A¹, SCN9A¹, SHANK3¹, SLC12A2¹, SLC12A5¹, SLC17A6¹, SMAL¹, SORT1¹, SPARC¹, SPP1¹, ST8SIA1¹, SIRT1⁷, TAC1¹, TAC4¹, TACR1¹, TBK1¹, TDAG8¹, TLR4¹, TMEM16F¹, TMEM35A¹, TNF¹, TNFRSF1A¹, TNFRSF1B¹, TRPA1¹, TRPC5¹, TRPM2¹, TRPM8¹, TRPV1¹, TRPV4¹, TSC2¹, VGLUT2¹.

Anexo I- Bibliografía

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