

Sociedad Latinoamericana de Immunodeficiencias

Estatísticas - Registro de IDPs

Intravenous Immunoglobulin



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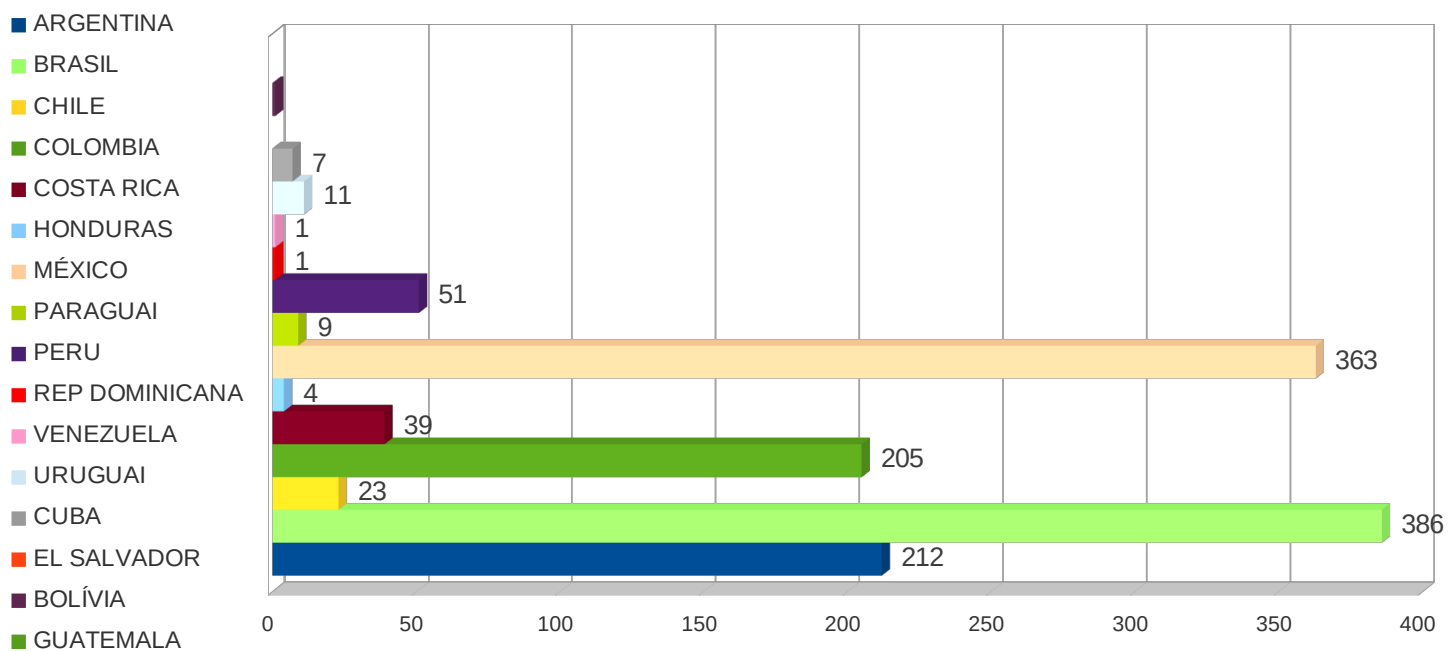
Objetivo

Este relatório apresenta dados estatísticos extraídos do Registro de Imuno Deficiências Primárias do LASID, referente ao mês de Junho de 2018.

São apresentados os casos registrados por IDP que fazem uso de '**intravenous immunoglobulin**' em sua terapia. Na ocasião da extração desses dados haviam **7.807** pacientes registrados no total.

1. Intravenous immunoglobulin por país

PAÍS	ABRIL / 2018	MAIO / 2018	JUNHO / 2018
ARGENTINA	212	212	212
BRASIL	371	381	386
CHILE	23	23	23
COLOMBIA	200	202	205
COSTA RICA	39	39	39
HONDURAS	4	4	4
MÉXICO	359	362	363
PARAGUAI	9	9	9
PERU	51	51	51
REP DOMINICANA	1	1	1
VENEZUELA	1	1	1
URUGUAI	11	11	11
CUBA	7	7	7
EL SALVADOR	0	0	0
BOLÍVIA	1	1	1
GUATEMALA	0	0	0
	1289	1304	1313



2. Intravenous immunoglobulin por PID

Uso intravenous immunoglobulin por PID	JUNHO / 2018
pdt; ada_ada - ADA deficiency	9
pad; ag_unknown - Agammaglobulinemias with unknown genetic cause	93
pad; ag_xla - XLA (Btk.)	181
aid; alps_fas - Fas (CD95)	1
aid; alps_unknown - ALPS with unknown genetic cause	6
pdt; cd3_cd3g - CD3 gamma deficiency	2
pdt; cd4_cd4 - Selective CD4 cell deficiency	2
pd; cgd_unknown - CGD with unknown genetic cause	2
pd; cgd_xlinkedcybb - CGD X-linked (CYBB)	1
owd; chh_pmrpcchh - PMRP/CHH	1
pd; chs_chs1 - Chediak-Higashi syndrome (CHS1/Lyst)	4
pd; chs_unknown - CHS with unknown genetic cause	2
pdt; cmc_apced - APECED (AIRE)	1
pdt; cmc_unknown - Other CMC	2
pad; csr_aid - AID deficiency (AICDA)	2
pad; csr_cd40 - CD40 deficiency	1
pad; csr_cd40l - CD40L deficiency (CD154)	18
pad; csr_uhigm - CSR defects and HIGM syndromes with unknown genetic cause	25
pdt; dgs_chrom22 - DiGeorge Syndrome	14
owd; dnab_atm - Ataxia telangiectasia (ATM)	76
owd; dnab_other - Other DNA-breakage disorder	1
pd; dwstmi_other - Susceptibility to mycobacterial infection and unknown genetic defect	2
pd; dwstmi_stat1 - STAT1 deficiency	2
pd; griscs_unknown - Griscelli syndrome with unknown genetic cause	6
pad; hg_baffr - BAFFR deficiency	1
pad; hg_cd19 - CD19 deficiency	6
pad; hg_cvid - CVID	374
pad; hg_cviddi - Secondary hypogammaglobulinemia	10
pad; hg_goods - Good-syndrome (associated with thymoma)	3
pad; hg_iggspecdef - Deficiency of specific IgG	66
pad; hg_iggsub - IgG subclass deficiency	30
pad; hg_siga - Selective IgA deficiency	16
pad; hg_sigm - Selective IgM deficiency	1
pad; hg_taci - TACI deficiency	1
pad; hg_transhypinf - Transient hypogammaglobulinemia of infancy	95

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pad; hg_unknown - Other Hypogammaglobulinemias	21
owd; hige_hies - HIES	33
pdt; hla_ciita - Bare lymphocyte syndrome (CII TA)	6
idouc; idouc_idouc - Unclassified immunodeficiencies	6
pd; lad_lad1 - LAD1 = CD11/CD18 (CD18/ITGB2)	4
pd; nedcn_el2 - ELA2 defective neutropenia	2
owd; ostp_clcn7 - Osteopetrosis (CLCN7)	1
pd; other_other - Other phagocytic disorders	8
pdt; outd_outd - Other unclassified T-cell disorders	35
pd; perfd_prf1 - PRF1 deficiency	1
pdt; pnp_pnp - PNP deficiency	4
pd; scn_scn - Severe congenital neutropenia	3
pdt; tbscid1_artemis - Artemis deficiency	1
pdt; tbscid1_rag1 - RAG 1 deficiency	3
pdt; tbscid1_unknown - T-B- SCID with unknown genetic cause	28
pdt; tbscid2_il7r - IL7R deficiency (IL-7R-alpha)	1
pdt; tbscid2_unknown - T-B+ SCID with unknown genetic cause	25
pdt; tbscid2_xlinkedgammac - SCID X-linked (gamma-c) (CD132)	12
owd; tlr_nemo - NEMO/IKK-gamma defiency (XED)	1
owd; was_unknown - WAS with unknown genetic cause	27
owd; was_wasp - WAS with mutations in WASP	25
owd; xlp_sh2d1a - XLP (SH2D1A)	5
owd; xlp_unknown - XLP with unknown genetic cause	3
pdt; zap_zap70 - ZAP 70	1
	1313