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Endokrinologie | Diabetologie | Osteologie | Stoffwechselerkrankungen · Dept.
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Units

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Publications (12)

Koster K, Rothermundt C, Binet F, Borovicka J, Bozinov O, Clerici T, Engeler D, Greiner J, Hader C, Heinimann K, Azzarello-Burri S, Lang C, Krull I, Stöckli S, Omlin A, Hundsberger T. Von-Hippel-Lindau-Erkrankung – Interdisziplinäre Betreuung und neue therapeutische Optionen. Swiss Med Forum 2022; 48:4-784-787.

Streuli R, Krull I, Brändle M, Kolb W, Stalla G, Theodoropoulou M, Enzler-Tschudy A, Bilz S. A rare case of an ACTH/CRH co-secreting midgut neuroendocrine tumor mimicking Cushing's disease. Endocrinol Diabetes Metab Case Rep 2017; 2017

Lipowsky C, Sze L, Krull I, Brändle M. Liraglutide as add-on therapy to insulin in type 2 diabetes mellitus: a retrospective, observational study from a daily clinical practice setting in Switzerland. Diabetes Ther 2015; 6:41-7.

Schmid S, Rothermundt C, Weber J, van Leyen K, Sulz M, Stöckli S, Krull I, Krek W, Kloos P, Heinimann K, Hader C, Greiner J, Engeler D, Brändle M, Binet F, Gillessen Sommer S, Hundsberger T. Management of von Hippel-Lindau Disease: An Interdisciplinary Review. Oncol Res Treat 2014; 37:761-771.

Krull I, Brändle M. Hyperthyreose: Diagnostik und Therapie. Swiss Medical Forum 2013; 13:954-960.

Krull I. Hyperthyreose: Diagnostik und Therapie. Swiss Medical Forum 2013; 13:954-960.

Krull I, Maier M, Bärlocher K, Koehler K, Huebner A, Brändle M. Two patients with an identical novel mutation in the AAAS gene and similar phenotype of triple A (Allgrove) syndrome. Exp Clin Endocrinol Diabetes 2010; 118:530-6.

Krull I, Christ E, Kamm C, Ganter C, Sahli R. Hyponatremia associated coma due to pituitary apoplexy in early pregnancy: a case report. Gynecol Endocrinol 2010; 26:197-200.

Krull I. Variability in cross-reactivity of novel insulin analogues in immunometric insulin assays. Diabetic Medicine 2009; 2009 Oct;26(10):1075-6:1075-1076.

Krull I. Nicht verzagen, den Pathologen fragen. Swiss Medical Forum 2009; 9:779-781.

Krull I. A novel succinate dehydrogenase subunit B gene mutation, H132P, causes familial malignant sympathetic extraadrenal paragangliomas. J Clin Endocrinol Metab 2004; 89(1):362-7:362-367.

Maier M, Schmid C, Galeazzi R, Krull I, Heitz P, Locher T, Schmid S, Saremaslani P, Komminoth P, Brändle M, Perren A. A novel succinate dehydrogenase subunit B gene mutation, H132P, causes familial malignant sympathetic extraadrenal paragangliomas. The Journal of clinical endocrinology and metabolism 2004; 89:362-7.

Projects (6)

CAHmelia SPR001-203 <i>Clinical Studies - Dec 1, 2021 - Dec 31, 2025</i> <i>Ongoing</i>
CAHmelia SPR001-204 <i>Clinical Studies - Dec 1, 2021 - Dec 31, 2025</i> <i>Ongoing</i>
Akromegalie-Register <i>Clinical Studies - Sep 6, 2019 - Dec 31, 2029</i> <i>Ongoing</i>
LX4211 310 <i>Clinical Studies - Oct 10, 2015 - Dec 31, 2017</i> <i>Completed</i>
LUNA Studie <i>Clinical Studies - Jan 21, 2014 - Jan 21, 2014</i> <i>Completed</i>
Sekretion von Katecholaminen und ihrer methoxylierten Metabolite durch die menschliche Nebenniere und durch Phäochromozytome <i>Clinical Studies - May 30, 2013 - Dec 31, 2020</i> <i>Automatically Closed</i>