Autosomal Dominant Hypocalcemia Type 1: A Systematic Review of the Genotypic and Phenotypic Spectrum, and Effects of Treatment ¹KL Roszko, ²LM Stapleton, ²AV Sridhar, ²MS Roberts, ¹RI Gafni, ¹MT Collins, ³EF Nemeth



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Introduction

- Autosomal dominant hypocalcemia type 1 (ADH1) is a rare form of hypoparathyroidism due to activating mutations of the calcium-sensing receptor (CaSR) gene (CASR) causing low parathyroid hormone (PTH) levels, hypocalcemia, hyperphosphatemia, and relative hypercalciuria.
- Conventional therapy includes calcium and active vitamin D; however, this treatment can worsen hypercalciuria and lead to renal complications.

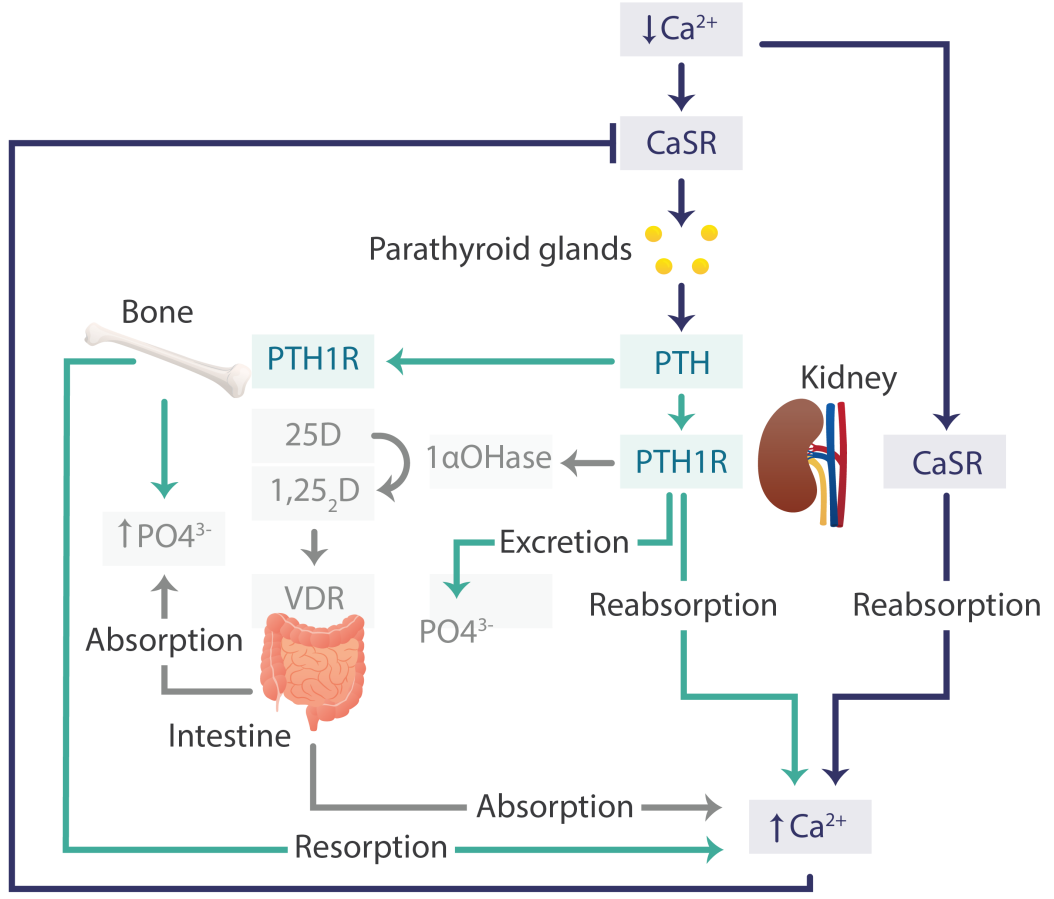
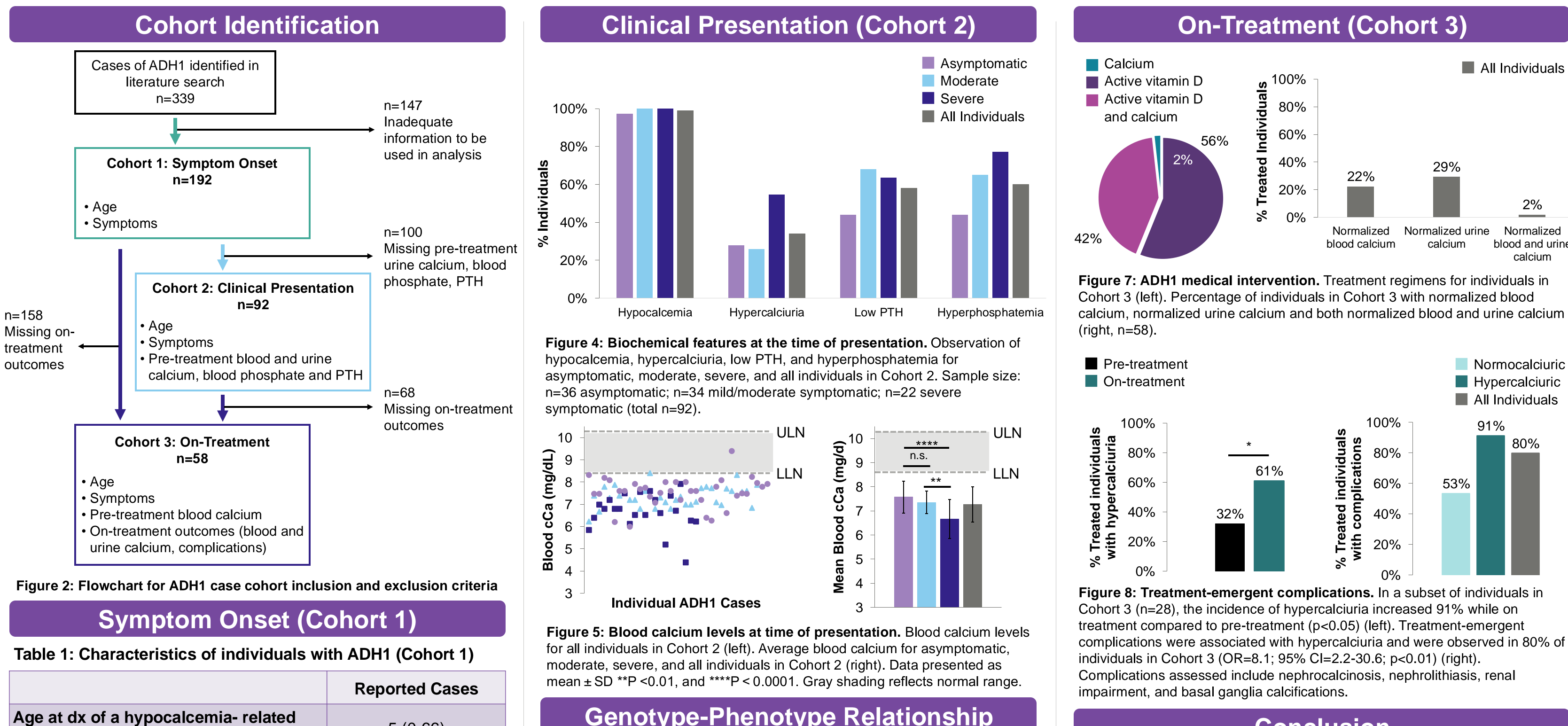


Figure 1: Regulation of systemic mineral metabolism. In the parathyroid, low levels of extracellular Ca²⁺ diminish signaling through the CaSR which leads to increased synthesis and secretion of PTH. PTH stimulates bone resorption to release Ca²⁺ and phosphate into the blood. It also acts in the kidney to increase the synthesis of 1,25-dihydroxyvitamin D_3 , which then increases the absorption of dietary calcium and phosphate in the small intestine. PTH decreases phosphate and increases Ca²⁺ reabsorption in the kidney. CaSRs in the kidney regulate Ca²⁺ and phosphate reabsorption and possibly 1,25-dihydroxyvitamin D synthesis, opposing the actions of PTH.

Systematic Review Methodology

- Search Criteria: autosomal dominant hypocalcemia; familial hypoparathyroidism; genetic/congenital hypoparathyroidism; hypercalciuric hypocalcemia; activating CASR variants
- Extracted Data: age at clinical presentation, mode of diagnosis, CASR variant, presenting symptoms, pre-treatment biochemical profile, on-treatment blood calcium, urine calcium, and complications
- Final Analysis Set: 339 ADH1 cases across 76 published reports spanning 1994-2021.

1. NIDCR, NIH, Bethesda, MD, USA, 20892; 2. Calcilytix Therapeutics, Inc, San Francisco, CA, USA, 94104; 3. MetisMedica, Toronto, ON, Canada, M4V 2M7

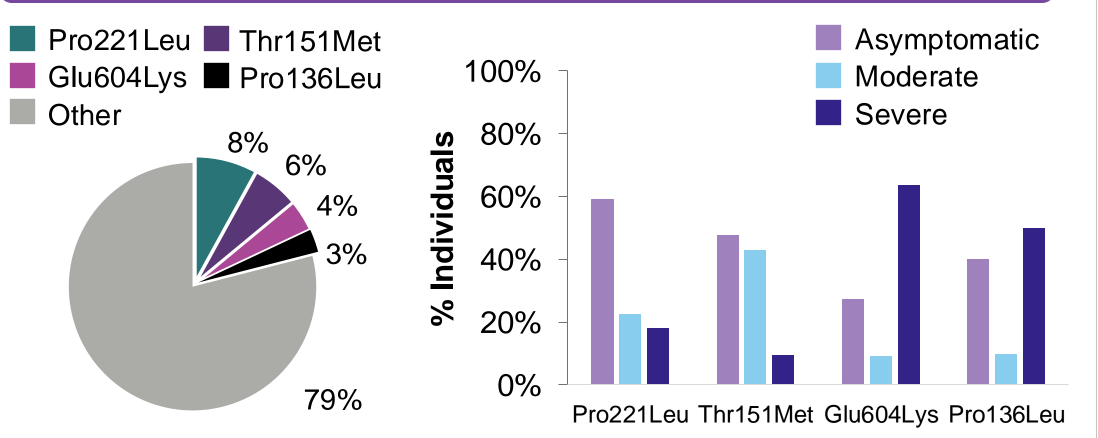


	Reported Cases
Age at dx of a hypocalcemia- related disorder, median (range)	5 (0-66)
Age of dx of ADH1, median (range)	25 (0-77)
Primary driver of dx (%)	
Symptom Driven	71%
Family Screen	23%
Incidental	6%
Symptom burden at presentation (%)	
Asymptomatic	27%
Moderate	32%
Severe	41%

Primary driver of dx refers to the main factor contributing to a diagnosis of a hypocalcemia-related disorder. Asymptomatic symptom burden refers to the absence of hypocalcemia-related symptoms. Moderate symptom burden includes muscle cramps/spasms, bone/joint pain, tetany, paresthesia, brain fog, and/or fatigue. Severe symptom burden includes seizures, loss of consciousness, and/or laryngospasm.

Figure 6: CASR variants. Four most common variants with reported symptoms as a proportion of all the variants identified the literature (n=339), regardless of kindred (left). Symptom severity for each common variant was analyzed (right). Overall, there was no direct correlation between CASR genotype and clinical phenotype. Unique variants (n=114) were present throughout the CaSR but most (55%) were in the extracellular domain, 33% were within the transmembrane domain, and 12% in the intracellular domain.

Genotype-Phenotype Relationship



blood and urine

bridge

Conclusion

This is the largest systemic literature review of ADH1 cases which analyzed 339 cases and uncovered >100 unique CASR variants

There was a wide range of clinical heterogeneity and a general lack of genotype/phenotype correlation in ADH1 individuals

Lower blood calcium levels were associated with more severe symptoms

Conventional treatment was associated with an increase in hypercalciuria, and the available data suggest an inability to simultaneously normalize blood and urine calcium

These findings underscore the limitations of the current standard of care for ADH1 patients

References

Roszko KL, et al. Front Physiol. 2016; 7:458. 2. Gorvin, CM, et al. J Mol Endocrinol. 2019; 63:2.