

# Resource Summary Report

Generated by [dkNET](#) on Aug 16, 2026

## Asper Biotech

RRID:SCR\_000700

Type: Tool

---

### Proper Citation

Asper Biotech (RRID:SCR\_000700)

---

### Resource Information

**URL:** <http://www.asperbio.com>

**Proper Citation:** Asper Biotech (RRID:SCR\_000700)

**Description:** A genetic testing company for rare and complex disorders and syndromes. The company specializes in retinal disorders, reproductive medicine and oncology. They also offer custom genotyping services.

**Synonyms:** AsperBio

**Resource Type:** material resource, instrument supplier

**Keywords:** genetic, test, syndrome, disorder, retinal, reproductive, medicine, oncology, genotyping, genotype, dna, blood, saliva, microarray, primer

**Resource Name:** Asper Biotech

**Resource ID:** SCR\_000700

**Alternate IDs:** nif-0000-30125

**Record Creation Time:** 20220129T080203+0000

**Record Last Update:** 20260815T182157+0000

---

### Ratings and Alerts

No rating or validation information has been found for Asper Biotech.

No alerts have been found for Asper Biotech.

---

### Data and Source Information

## Usage and Citation Metrics

We found 10 mentions in open access literature.

**Listed below are recent publications.** The full list is available at [dkNET](#) .

Wawrocka A, et al. (2018) Novel variants identified with next-generation sequencing in Polish patients with cone-rod dystrophy. *Molecular vision*, 24, 326.

Smaragda K, et al. (2018) Mutation Spectrum of the ABCA4 Gene in a Greek Cohort with Stargardt Disease: Identification of Novel Mutations and Evidence of Three Prevalent Mutated Alleles. *Journal of ophthalmology*, 2018, 5706142.

Van Cauwenbergh C, et al. (2017) Mutations in Splicing Factor Genes Are a Major Cause of Autosomal Dominant Retinitis Pigmentosa in Belgian Families. *PloS one*, 12(1), e0170038.

Coppieters F, et al. (2015) Hidden Genetic Variation in LCA9-Associated Congenital Blindness Explained by 5'UTR Mutations and Copy-Number Variations of NMNAT1. *Human mutation*, 36(12), 1188.

Verma A, et al. (2013) Mutational screening of LCA genes emphasizing RPE65 in South Indian cohort of patients. *PloS one*, 8(9), e73172.

Ferrari S, et al. (2011) Retinitis pigmentosa: genes and disease mechanisms. *Current genomics*, 12(4), 238.

Coppieters F, et al. (2010) Genetic screening of LCA in Belgium: predominance of CEP290 and identification of potential modifier alleles in AHI1 of CEP290-related phenotypes. *Human mutation*, 31(10), E1709.

Aguirre-Lamban J, et al. (2009) Molecular analysis of the ABCA4 gene for reliable detection of allelic variations in Spanish patients: identification of 21 novel variants. *The British journal of ophthalmology*, 93(5), 614.

Maia-Lopes S, et al. (2009) ABCA4 mutations in Portuguese Stargardt patients: identification of new mutations and their phenotypic analysis. *Molecular vision*, 15, 584.

Riveiro-Alvarez R, et al. (2008) Molecular analysis of ABCA4 and CRB1 genes in a Spanish family segregating both Stargardt disease and autosomal recessive retinitis pigmentosa. *Molecular vision*, 14, 262.