## **Resource Summary Report**

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# **GESND**

RRID:SCR\_005179 Type: Tool

**Proper Citation** 

GESND (RRID:SCR\_005179)

#### **Resource Information**

URL: http://sourceforge.net/projects/gesnd/

Proper Citation: GESND (RRID:SCR\_005179)

**Description:** A software package and a pipeline for identifying causal mutations for rare congenital diseases by next-generation sequencing. Features \* one-stop solution for identifying causal mutations of rare genetic diseases \* detect wide-spctrum variants, including medium and large sized indels, and tandem repeats \* annotate and filter variants \* prioritize candidate variants

Abbreviations: GESND

Synonyms: Genetic Screening and Diagnosis, GESND - Genetic Screening and Diagnosis

Resource Type: software resource

Keywords: next-generation sequencing, mutation, variant, indel, tandem repeat

Related Condition: Rare congenital disease

Funding:

Resource Name: GESND

Resource ID: SCR\_005179

Alternate IDs: OMICS\_00175

**Record Creation Time:** 20220129T080228+0000

#### **Ratings and Alerts**

No rating or validation information has been found for GESND.

No alerts have been found for GESND.

#### Data and Source Information

Source: <u>SciCrunch Registry</u>

### **Usage and Citation Metrics**

We have not found any literature mentions for this resource.