

## Kallman Syndrome

Kallman syndrome, or hypogonadotropic hypogonadism, is a genetic disorder leading to failure to start or complete puberty. This occurs because of defective migration of GnRH releasing-neurons to the hypothalamus, leading to hypogonadism. Patients display anosmia, delayed puberty and infertility.



PLAY PICMONIC

### Mutation in KAL-1 or FGFR-1 Gene

[Mutant who Calls-1 for FroG-Firefighter-\(1\) Wand](#)

Though most cases of this disorder have an unknown origin, there is evidence that FGFR-1 (fibroblast growth factor receptor 1) and KAL1 gene defects can lead to Kallman Syndrome. These genes help normal migration of GnRH neurons into the hypothalamus.

### Failure of GnRH Secreting Neurons

[Failure of Gonad-gopher Nerve Secretion](#)

This syndrome occurs from defective migration of GnRH releasing neurons to the hypothalamus. This leads to decreased synthesis and release of GnRH in the hypothalamus, leading to decreased gonadotropin release.

### Delayed Puberty

[Delayed Pubescent-teen](#)

Due to low serum testosterone and gonadotropins, LH and FSH, patients can show delayed puberty. This can also manifest as incomplete puberty, which is also a cause of hypogonadotropic hypogonadism.

### Anosmia

[Broken A-nose](#)

Anosmia occurs because normal GnRH neurons originate in the olfactory placode, pass through the olfactory bulb to the hypothalamus. If there is a neuron migration issue, where this path is no longer taken, the patient's smell becomes defective because of olfactory bulb development issues.

### Hypogonadism

[Hippo-gonads](#)

Due to improper migration of GnRH releasing cells, hypogonadism occurs. Patients have low levels of FSH, LH and resultingly, testosterone and estrogen.

### Infertility

[Infertile-plant](#)

Due to hormone abnormalities, adult patients are often infertile. Males can show low sperm counts, while many females will present with amenorrhea.