

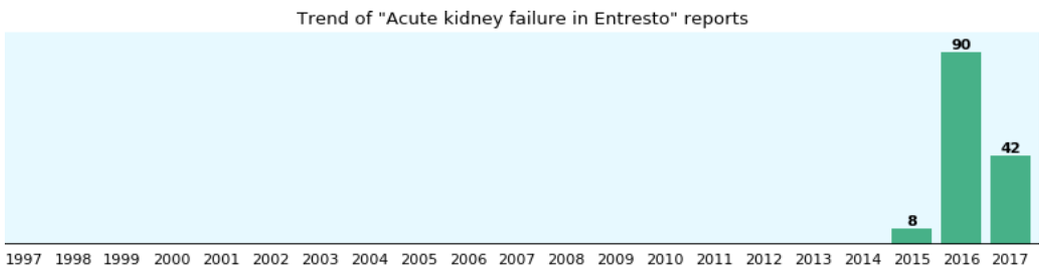
# Entresto and Acute kidney failure - from FDA reports

On Jul, 23, 2017

9,385 people reported to have side effects when taking Entresto.

Among them, 140 people (1.49%) have Acute kidney failure

## Number of reports submitted per year:



## Time on Entresto when people have Acute kidney failure \*:

- < 1 month: 53.66 %
- 1 - 6 months: 36.59 %
- 6 - 12 months: 9.76 %
- 1 - 2 years: 0.0 %
- 2 - 5 years: 0.0 %
- 5 - 10 years: 0.0 %
- 10+ years: 0.0 %

## Gender of people who have Acute kidney failure when taking Entresto \*:

- female: 22.12 %
- male: 77.88 %

## Age of people who have Acute kidney failure when taking Entresto \*:

- 0-1: 0.0 %
- 2-9: 0.0 %
- 10-19: 0.0 %
- 20-29: 0.0 %
- 30-39: 0.0 %
- 40-49: 8.2 %

- 50-59: 16.39 %
- 60+: 75.41 %

### Top conditions involved for these people \*:

- Cardiac Failure (37 people, 26.43%)
- Cardiac Failure Chronic (20 people, 14.29%)
- Left Ventricular Failure (11 people, 7.86%)
- Atrial Fibrillation/flutter (5 people, 3.57%)
- Cardiac Failure Congestive (4 people, 2.86%)

### Top co-used drugs for these people \*:

- Lasix (5 people, 3.57%)
- Coreg (5 people, 3.57%)
- Aldactone (5 people, 3.57%)
- Revatio (4 people, 2.86%)
- Eliquis (4 people, 2.86%)

### Top other side effects for these people \*:

- Hypotension (23 people, 16.43%)
- Cardiac Failure (21 people, 15.00%)
- Hyperkalemia (10 people, 7.14%)
- Death (10 people, 7.14%)
- Breathing Difficulty (9 people, 6.43%)

\* Approximation only. Some reports may have incomplete information.

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**NOTE:** The study is based on active ingredients and brand name. Other drugs that have the same active ingredients (e.g. generic drugs) are NOT considered.

**WARNING:** Please DO NOT STOP MEDICATIONS without first consulting a physician since doing so could be hazardous to your health.

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You may report adverse side effects to the FDA at <http://www.fda.gov/medwatch/> or 1-800-FDA-1088 (1-800-332-1088).

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