

# Person—bleeding episode status, code N

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# Person—bleeding episode status, code N

## Identifying and definitional attributes

<b>Metadata item type:</b>	Data Element
<b>Short name:</b>	Bleeding episode using TIMI criteria (status)
<b>METEOR identifier:</b>	284812
<b>Registration status:</b>	<a href="#">Health</a> , Superseded 01/10/2008
<b>Definition:</b>	A person's episode of bleeding as described by the Thrombolysis In Myocardial Infarction (TIMI) criteria, as represented by a code.
<b>Data Element Concept:</b>	<a href="#">Person—bleeding episode status</a>
<b>Value Domain:</b>	<a href="#">Bleeding episode status code N</a>

## Value domain attributes

## Representational attributes

<b>Representation class:</b>	Code	
<b>Data type:</b>	Number	
<b>Format:</b>	N	
<b>Maximum character length:</b>	1	
<b>Permissible values:</b>	<b>Value</b>	<b>Meaning</b>
	1	Major
	2	Minor
	3	Non TIMI bleeding
	4	None
<b>Supplementary values:</b>	9	Not stated/inadequately described

## Collection and usage attributes

<b>Guide for use:</b>	<p>Note in calculating the fall in haemoglobin or haematocrit, transfusion of whole blood or packed red blood cells is counted as 1g/dl (0.1g/l) haemoglobin or 3% absolute haematocrit.</p> <p>CODE 1    Major</p> <p>Overt clinical bleeding (or documented intracranial or retroperitoneal haemorrhage) associated with a drop in haemoglobin of greater than 5g/dl (0.5g/l) or a haematocrit of greater than 15% (absolute).</p> <p>CODE 2    Minor</p> <p>Overt clinical bleeding associated with a fall in haemoglobin of 3g/dl to less than or equal to 5g/dl (0.5g/l) or a haematocrit of 9% to less than or equal to 15% (absolute).</p> <p>CODE 3    Non TIMI Bleeding</p> <p>Bleeding event that does not meet the major or minor definition.</p> <p>CODE 4    None</p> <p>No bleeding event</p>
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## Source and reference attributes

**Submitting organisation:** Australian Institute of Health and Welfare

## Data element attributes

### Source and reference attributes


**Submitting organisation:** Acute coronary syndrome data working group

**Steward:** [The National Heart Foundation of Australia and The Cardiac Society of Australia and New Zealand](#)

**Origin:** Rao AK, Pratt C, Berke A, et al. Thrombolysis in Myocardial Infarction (TIMI) Trial, phase I: hemorrhagic manifestations and changes in plasma fibrinogen and the fibrinolytic system in patients with recombinant tissue plasminogen activator and streptokinase. J Am Coll Cardiol 1988; 11:1-11.

### Relational attributes

**Related metadata references:** Has been superseded by [Person—bleeding episode status, Thrombolysis in Myocardial Infarction \(TIMI\) code N](#)  
[Health](#), Standard 01/10/2008

Is re-engineered from  [Bleeding episode using TIMI criteria - status, version 1, DE, NHDD, NHIMG, Superseded 01/03/2005.pdf](#) (15.3 KB)  
*No registration status*

**Implementation in Data Set Specifications:** [Acute coronary syndrome \(clinical\) DSS](#)  
[Health](#), Superseded 01/10/2008

***DSS specific information:***

Can be collected at any time point during the management of the current event (i.e. at the time of triage, at times during the admission, or at the time of discharge).

[Acute coronary syndrome \(clinical\) DSS](#)  
[Health](#), Superseded 07/12/2005

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