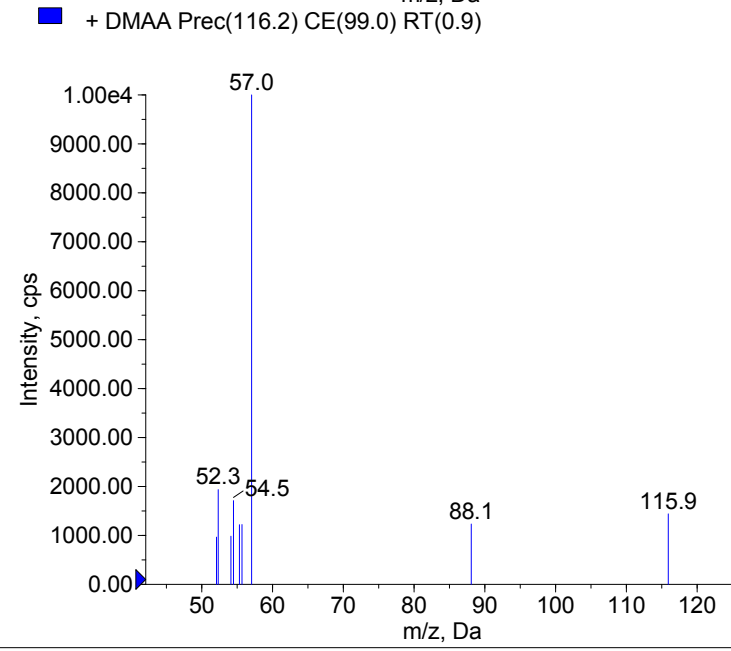
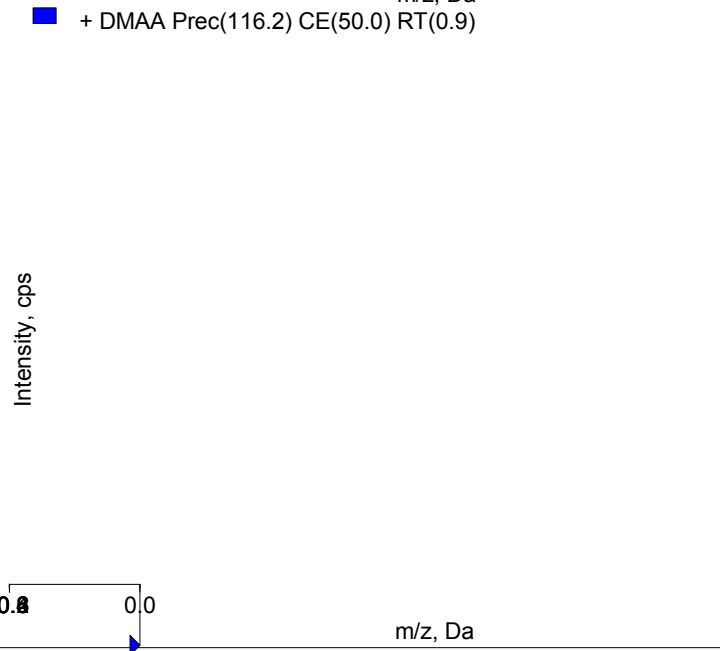
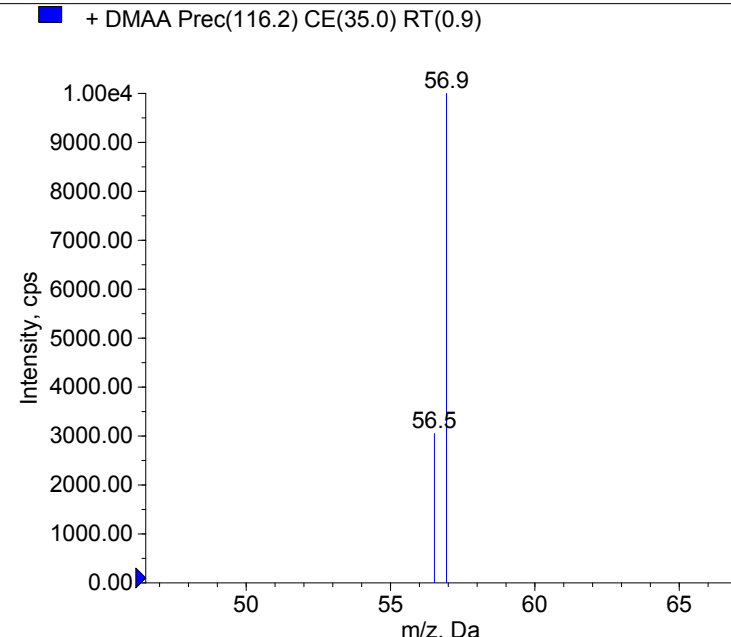
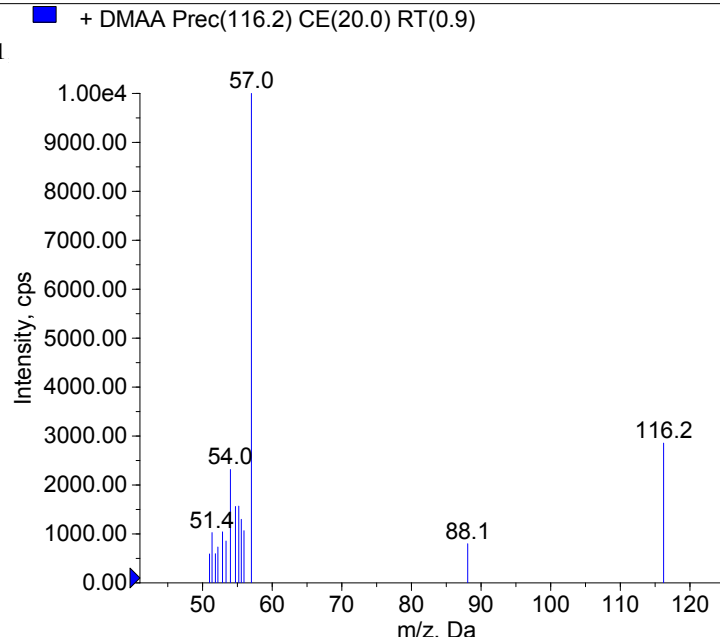


Compound Name: DMAA
 Synonyms: geranamine; methylhexanamine; 1
 Formula: C7H17N
 CAS Number: 105-41-9
 Molecular weight(Da): 115.1361
 Compound Class: Others
 Compound ID:
 User Value: 0.0000
 Keyword:
 Internal Standard: No
 Comment1: 115.136100



Instrument Model: 3200 Q TRAP
 Ion Source: Turbo Spray
 Polarity: Positive
 Scan Type: EPI
 1st Precursor m/z: 116.2000
 Collision Energy1(V): 20.0000
 Charge State1: 1
 CAD Gas Type:
 CAD Gas Value: Level 4
 Retention Time(min): 0.90
 Resolution: unit
 Comment2:
 Comment3:

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 www.irm.unibe.ch

Comment1: monoisotopic mass
 "CE(99.0)": CE spread 35 +/- 15 eV

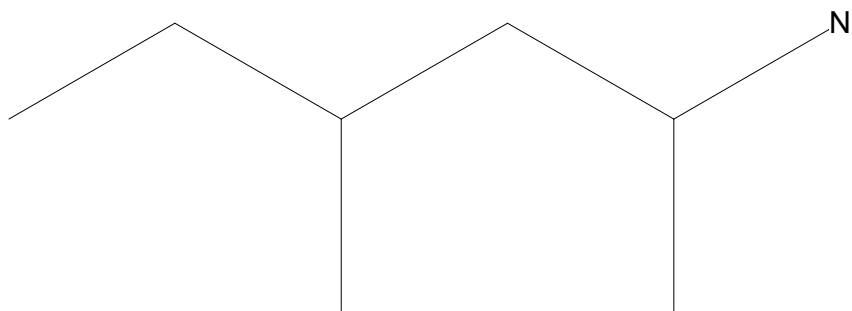
C7H17N

Chemist Name:

Project Name:

Workbook Name:

Workbook Page:



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Comment1: monoisotopic mass
"CE(99.0)": CE spread 35 +/- 15 eV