

# Molecular Origami of C2H2

given information

ElementNames	[ (C) (C) (H) ]
dotted	F
distance	102.468
distance	117.572
angle	179.999
	220.
dopage	T
AutoAlign	F
showboth	F

C<sup>1</sup>-H<sup>1</sup>  
C<sup>1</sup>-C<sup>2</sup>  
H<sup>1</sup>-C<sup>1</sup>-C<sup>2</sup>  
H<sup>1</sup>-C<sup>2</sup>

structure type: XAB

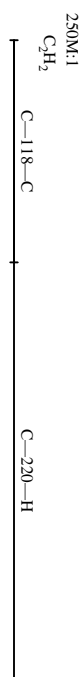
# Molecular Origami of C2H2

!C1  
C2  
H1  
C2H2

linear

scale 250,000,000 : 1  
units: pm  
offsetx -0.05 offsety 0.9

View -1



Current: (centerx 4.25) (centery 5.90) (scale 250)

%%BoundingBox: 296 176 316 435 actual: 306 186 306 425

Better: (centerx 4.25) (centery 7.16) (scale 250)

%%BoundingBox: 292 331 312 590 actual: 302 341 302 580

center: 306 305

actual size: 0 239

center: 302 461

actual size: 0 239