# Molecular Coding Format manual

## Akira Yamaji

September 20, 2021

Located at http://www.ctan.org/pkg/mcf2graph
Suggestion or request mail to: mcf2graph@gmail.com

H <sub>2</sub> N COOH	H <sub>2</sub> N COOH		H <sub>2</sub> N COOH
//,,	0H H <sub>2</sub> N C00H	H <sub>2</sub> N COOH	SH H <sub>2</sub> N COOH
H <sub>2</sub> N COOH	H <sub>2</sub> N COOH	H <sub>2</sub> N COOH	H <sub>2</sub> N COOH
COOH	H <sub>2</sub> N COOH	NH <sub>2</sub>	COOH H <sub>2</sub> N COOH
COOH H <sub>2</sub> N COOH	H <sub>2</sub> N COOH	H <sub>2</sub> N COOH	H <sub>2</sub> N COOH
OH OH H <sub>2</sub> N COOH	NH <sub>2</sub> H <sub>2</sub> N COOH	H <sub>2</sub> N COOH 0	H <sub>2</sub> N COOH
H <sub>2</sub> N COOH	COOH H <sub>2</sub> N	COOH COOH H <sub>2</sub> N COOH	HO NH

# Contents

## 1 Introduction

Molecular Coding Format(MCF) is new linear notation represent chemical structure diagrams. This Coding is named from programing technique such as operator, array, scope, macro, adressing, etc. mcf2graph convert from MCF to pk font, PNG, SVG, EPS, MOL file. It is also able to calculate molecular weight, exact mass, molecular formula.

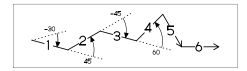
# 2 MCF syntax

#### 2.1 Make bond

#### 2.1.1 Chain

real number plus (+): counterclockwize
real number minus(-): clockwize
\$n (0<=n<360): absolute angle</pre>

<10,-30,45,-45,60,\$300,\$0

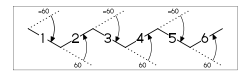


#### 2.1.2 Chain use !,!n

! : take value 60 or -60 depend on current angle and environment

!6: !,!,!,!,!,!

<-30,!6

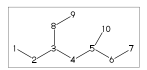


#### 2.1.3 Jump to atom

On : Jump to An

\*\* An: atom number(-999<=n<=4095)

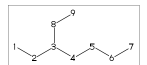
<-30,!6,@3,0,!,@5,-30



#### 2.1.4 Branch bond

\:0

<-30,!6,@3,\,!



#### 2.1.5 Branch modified bond

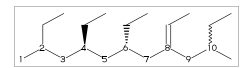
\ : 0 \*\ : 0~wf

\\* : 0~zf \\ : 0~dm

\*\\* : 0~wv

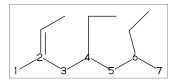
<30,!8,

02,\,!,04,\*\,!,06,\\*,!,08,\\,!,010,\*\\*,!



<30,!6,

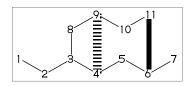
\^dr,!, : 0~dr,! \'1.5,-90 : 0'1.5,-90 \^15,-60 : 0^15,-60



#### 2.1.6 Connect atom

&n : Connect to An

<-30,!6,@3,\,!3,&6~bd,@9,&4~bz



#### 2.1.7 Ring

?n : n membered ring(3<=n<=20) ?6 : <-120,60,60,60,60,60,&1

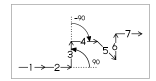
?6



#### 2.1.8 Rotate current angle

<angle : rotate current angle</pre>

0,0,<90,0,<-90,0,<\$315,0,<\$90,0,<\$0,0



#### 2.2 Change bond type

#### 2.2.1 Double, triple, wedge, vector

db : double left or right side

tm : triple

!! : !~db / !!! : !~tm

<-30,!~dm,!,!~dl,!,!~dr,!~db,!~db,!,!~tm <-30,!~dm,!,!~dl,!,!~dr,!! ,!! ,!,!!!



(Wedge, Vector)

wf: wedge forward
wb: wedge backward

zf: hashed(zebra stripe) wedge foward
zb: hashed(zebra stripe) wedge backward

vf:vector forward
vb:vector backward

<-30,

!~wf,!,!~wb,!,!~zf,!,!~zb,!,!~vf,!~vb



(Dotted, wave)

Bn=bond type : change bond type at Bn

dt : dotted / wv : wave

bd : broad / bz : broad dotted

<-30,!7,1=dt,3=wv,5=bd,7=bz



#### 2.2.2 Over line

si\_ : single over line

wf\_ : wedge forward over line
wb\_ : wedge backward over line

zf\_ : hashed wedge forward over line zb\_ : hashed wedge backward over line

bd\_ : broad over line
dl\_ : duble left over line
dr\_ : duble right over line

 ${\tt dm}\_$  : duble over line

<30,!8,!,60,90'18,
{2~si\_,4~wf\_,6~wb\_,8~zf\_,10~zb\_,
 12~bd\_,14~dl\_,16~dr\_,18~dm\_}:/\_'2</pre>



#### 2.2.3 Steric ring

wf\_r : wedge foward (half width)
bd\_r : broad (half width, rounded)
wb\_r : wedge backward (half width)

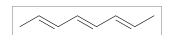
#1.25,-30~wf\_r,30~bd\_r'1,30~wb\_r, 120,0,30,&1,##,6^\$90:/!OH'-.5, {1^\$-90,2^\$90,3^\$-90,4^\$90}:/OH'-.5,



#### 2.2.4 Change multiple bond type

 ${2,4,6}=dr : 2=dr,4=dr,6=dr$ 

<30,!7,{2,4,6}=dr

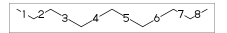


#### 2.3 Change bond length

#### 2.3.1 Chain length

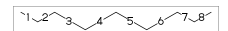
(!,!n)'length : change length of !,!n

<-30,!2,!4'1.2,!2



#n : bond length=n
## : reset bond length

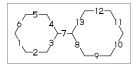
<-30,!2,#1.2,!4,##,!2



#### 2.3.2 Ring length

?n'length : change ring length

?6,04,\,?6'1.2



#### 2.4 Change atom

#### 2.4.1 Insert atom

Insert hetero atom

<-30,!2,0,!2,N,!2

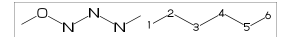


#### 2.4.2 Addressed atom

2:0 : change A2 C to O

 ${3,4}:N$ : change A3,A4 C to N

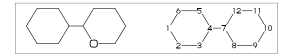
<30,!4,2:0,{3,4}:N



#### 2.4.3 Brock address

| : divide brock

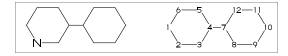
?6,04,\,|,?6,2:0



#### 2.4.4 Reset brock address

|| : reset brock adress

?6,04,\,|,?6,||,2:N



#### 2.4.5 Absolute address

\$2:N : change A\$2 C to N \*\*1<=n<=3095

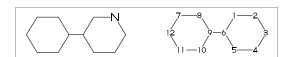
?6,@4,\,|,?6,\$2:N



#### 2.4.6 Relative address

-2:N: change A(-2) C to N \*\*-999<=n<=-1

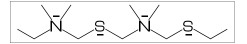
?6,@4,\,?6,-2:N



#### 2.4.7 Charged atom

p\_ : positive / n\_ : negative

<-30,!2,N,??,p\_,!2,S,n\_^180, !6,7:N,7:??,9:S,7:n\_,9:n\_^180



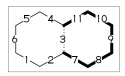
#### 2.5 Fuse ring

(Attached 1 bond)

?6,3=?6 : fuse ?6 at B3

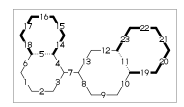
\*\*  $Bn(n:-999 \le n \le 4095)$ : bond number

?6,3=?6



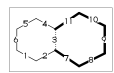
\*\* fused ring size depend on attached bond length

?6,04,\,?6'1.2,5=?6,11=?6

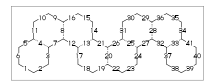


?6,3=?6[13] : fuse ?6[13] at B3
?6[13]: 6 membered ring scaled 13/10
\*\* ?m[n] (5<=m<=8,11<=n<=15)</pre>

?6,3=?6[13]



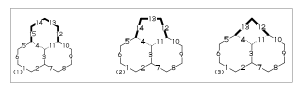
?6,{-3,-4,-4,-2,-2,-4,-4}=?6 ?6,{4,8,13,20,25,28,33}=?6



#### (Attached 2 bond)

(4,11)=?6[4] : fuse 4/6 ring to B11..B4 (4,11)=?5[3] : fuse 3/5 ring to B11..B4 (4,11)=?4[2] : fuse 2/4 ring to B11..B4 \*\* ?m[n] (4<=m<=6,n=m-2)

1:<30,?6,3=?6,(11,4)=?6[4] 2:<30,?6,3=?6,(11,4)=?5[3] 3:<30,?6,3=?6,(11,4)=?4[2]

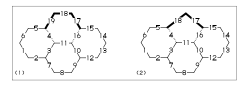


#### (Attached 3 bond)

(16,4)=?6[3]: fuse 3/6 ring to B16..B4 (16,4)=?5[2]: fuse 2/5 ring to B16..B4

\*\* ?m[n] (5<=m<=6,n=m-3)

1:?6,{3,10}=?6,(16,4)=?6[3] 2:?6,{3,10}=?6,(16,4)=?5[2]

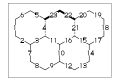


#### (Attached 4 bond)

(21,4)=?6[2]: fuse 2/6 ring to B21..B4

 $MC(<-30,?6,{3,10,15}=?6,(21,4)=?6[2])$ 

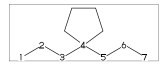
\*\* ?m[n] (m=6,n=2)



## 2.6 Spiro ring

04,?5 : add ?5 at A4

<30,!6,@4,?5



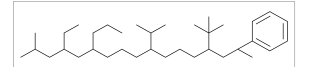
#### 2.7 Group

#### 2.7.1 Insert group

/ : group start single bond

/\_ : methyl
/! : ethyl
/!2 : propyl
/?! : isopropyl
/??! : tert-butyl
/Ph : phenyl

<30,!,/\_,!2,/!,!2,/!2,!4,/?!, !4,/??!,!2,/Ph^-60,!



#### 2.7.2 Insert modified group

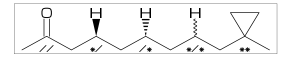
// : double (double middle)

\*/ : wedge forward

/\* : hashed wedge forward

\*/\* : wave \*\* : direct

<30,!,//0,!2,\*/H,!2,/\*H,!2,\*/\*H,!2,\*\*?3,!



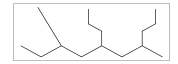
~ : change type

^ : change angle

': change length

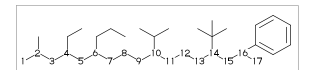
> : change environment

<-30, ''1,!, /\_'2^30,!2,/!2>lr,!2,/!2>rl,!)



#### 2.7.3 Add group

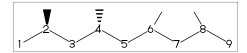
<30,!17,2:/\_,4:/!,6:/!2, 10:/?!,14:/??!,16:/Ph^-60



#### 2.7.4 Add modified group

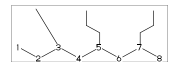
~,^,' : change type,angle,length

<30,!6,{2~wf,4~zf,6^-30,8^\$120}:/\_



^,',> : change angle,length,environment

<-30,!7'1,3:/\_'2^30,5:/!2>lr,7:/!2>rl



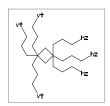
#### 2.8 Chain environment

#### 2.8.1 Horizontal, vertical

>hz : horizontal environment (default)

>vt : vertical environment

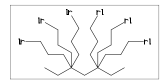
?4, {3^-90,3^-30,3^90}:/'(!3,"{hz}")>hz, {1^-60,1,1^60}:/'(!3,"{vt}")>vt



#### 2.8.2 Left-right,right-left

>lr : left-right environment
>rl : right-left environment

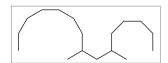
<-30,!6, {3^-30,3,3^30}:/'(!3,"{lr}")>lr, {5^-30,5,5^30}:/'(!3,"{rl}")>rl



#### 2.8.3 Fixed rotate angle

>n : rotate n

<30,!4, 2:/!6>30, % 2:\,30,30,30,30,30,30 4:/!4>-45 % 4:\,-45,-45,-45



#### 2.8.4 Multi rotate angle

>'(90,-90,...) : rotate 90,-90,...

<30,!6,6>'(90,-90,90,-90,90):/!5

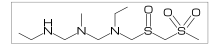


#### 2.9 Miscellaneous

#### 2.9.1 Abbreviated parts

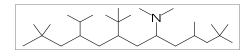
NH : N,/H~nl N! : N,/\_ N!2 : N,/! SO : S,//O SOO : S,//O^35,//^-35

<-30,!2,NH,!2,N!,!2,N!2,S0,!2,S00,!



/N?! : dimethylamino

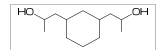
<30, !9'1,?!,!,??,!,2:??,4:/??,6:/??!,8:/N?!



#### 2.9.2 Parts definition

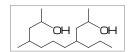
'(..): user defined parts

iBuOH:='(!,/\_,!,OH); MC(<30,?6,{4,6}:/iBuOH)



#### 2.9.3 Parts inline definition

<30,!8,{2,6}:/'(!,/\_,!,OH)

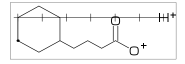


#### 2.9.4 Move position

Q(x,y): Move (1\*x,1\*y) from current position Q(x,y): Move (1\*x,1\*y) from origin Q(x,y)

\*\* l=bond length of ring

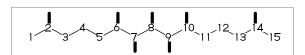
<30,?6,@3,!4,//0,!,0,n\_^60,@\$(6,1),H,p\_^15



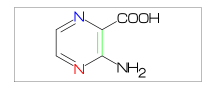
#### 2.9.5 Serial number

6--10 : 6,7,8,9,10

<30,!14,{2,6--10,14}:/\_~bd\_r'0.5

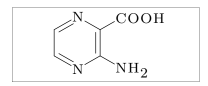


#### 2.9.6 Change color



#### 2.9.7 Change font

```
(use with metapost only)
beginfont()
    %------
    atomfont:="cmr8";
    %-----
    MC(<30,Ph,{2,5}:N,3:/NH2,4:/COOH)
endfont</pre>
```

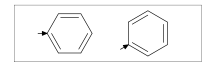


# 3 Option parameter

#### 3.1 Angle parameter

mangle=0 \*\* default

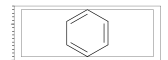
MCat(0.2,0.5)(Ph)
mangle:=30;
MCat(0.8,0.5)(Ph)



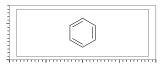
#### 3.2 Size/Ratio parameter

#### 3.2.1 Bond length

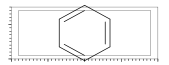
(fit to font size)
blength=0 \*\* default



(ratio bond/font width)
blength=0.1 \*\* (0<blength<=1)
blength=60mm(width)\*0.1=6mm</pre>

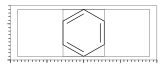


(bond length)
blength=9mm
\*\* (blength>1) ignore msize(w,h)

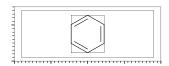


#### 3.2.2 Molecular size

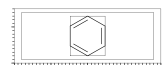
msize=(1,1) \*\* default



msize=(0.25,1)
msize=40mm-4mm\*0.25=9mm

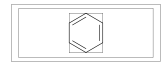


msize=(11mm,11mm)



#### 3.2.3 Molecular position

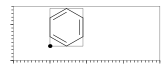
mposition=(0.5,0.5) \*\* default



mposition=(1,0)



mposition=(10mm,4mm)

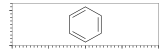


#### 3.3 Size parameter

#### 3.3.1 Font size

fsize=(font width,font height)
\*\* default: (30mm,20mm)

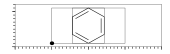
fsize=(40mm,15mm)



#### 3.3.2 Font margin

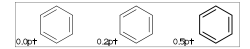
fmargin=(margin left rigth,top bottom)
\*\* default: (0.4mm,0.4mm)

fmargin=(10mm,2mm)



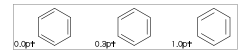
#### 3.3.3 Offset thickness of bond

default: offset\_thickness=0.2pt



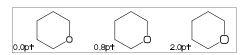
#### 3.3.4 Offset of double bond gap

default: offset\_bond\_gap=0.3pt



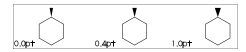
#### 3.3.5 Offset of atom width

default: offset\_atom=0.8pt



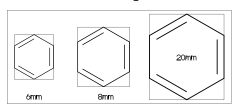
#### 3.3.6 Offset of wedge width

default: offset\_wedge=0.4pt



#### 3.3.7 Max bond length

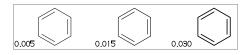
default: max\_blength=10mm



#### 3.4 Ratio parameter

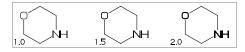
#### 3.4.1 Thickness/bond length

default: ratio\_thickness\_bond=0.015



#### 3.4.2 Char/bond thickness

default: ratio\_char\_bond=1.5



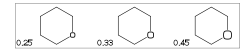
#### 3.4.3 Bond gap/bond length

default: ratio\_bondgap\_bond= 0.15



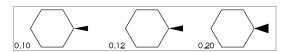
#### 3.4.4 Atom/bond length

default: ratio\_atom\_bond= 0.36



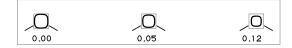
#### 3.4.5 Wedge/bond length

default: ratio\_wedge\_bond=0.12



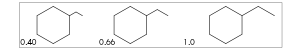
#### 3.4.6 Font atom gap/atom length

default: ratio\_atomgap\_atom= 0.050



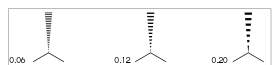
#### 3.4.7 Chain/ring length

default: ratio\_chain\_ring= 0.66



#### 3.4.8 Hash gap/bond length

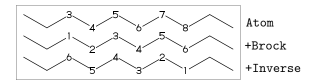
default: ratio\_hashgap\_bond=0.12



## 3.5 Drawing mode

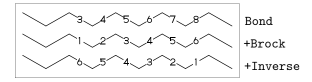
#### 3.5.1 Numbering atom

sw\_numbering=Atom
numbering\_start:=3; numbering\_end:=8;
default: sw\_numbering=0 :



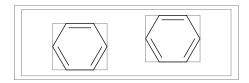
#### 3.5.2 Numbering bond

sw\_numbering=Bond
numbering\_start:=3; numbering\_end:=8;
default: sw\_numbering=0 :

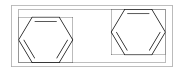


#### 3.5.3 Trimming mode

sw\_trimming:=0; \*\* default
msize:=(1,0.7);
MCat(0.2,0.3)(Ph)
MCat(0.8,0.7)(Ph)

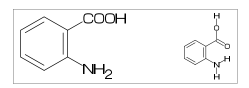


sw\_trimming:=1;
MCat(0.2,0.3)(Ph)
MCat(0.8,0.7)(Ph)



#### 3.5.4 Expand mode

MCat(0, .5)(<30,Ph,4:/COOH,3:/NH2)
sw\_expand:=1;
MCat(1, .5)(<30,Ph,4:/COOH,3:/NH2)
\*\* default: sw\_expand=0</pre>



#### 3.5.5 Abbreviate group

\*\* default: sw\_abbreviate=Group



#### 3.5.6 Abbreviate bond type

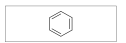
\*\* default: sw\_abbreviate=Bond



#### 3.6 Frame

#### 3.6.1 Font frame

\*\* default:sw\_frame=0
(Draw font frame)
fmargin:=(5mm,2mm);
sw\_frame=Outside



(Frame inside margin) sw\_frame=Inside

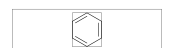


(Draw both frame)
sw\_frame=Bothside=Inside+Outside



#### 3.6.2 Molecular frame

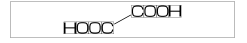
sw\_frame=Mol
\*\* default:sw\_frame=0



#### 3.6.3 Atom frame

sw\_frame=Atom
\*\* default: sw\_frame=0

MC(<30,COOH,!,COOH)



#### 3.7 Parameter setting

#### 3.7.1 Local parameter setting

```
beginfont()
  MC(Ph)
endfont
beginfont()
  %-----
  ratio_thickness_bond:=0.05;
  %------
  MC(Ph)
endfont
beginfont()
  MC(Ph)
endfont
```



#### 3.7.2 Global parameter setting

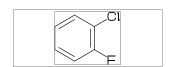
```
beginfont()
   MC(Ph)
endfont
%------
ratio_thickness_bond:=0.05;
%------
beginfont()
   MC(Ph)
endfont
beginfont()
   MC(Ph)
endfont
```



#### 4 Function

#### 4.1 Function MC()

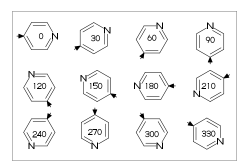
(Draw molecule)



endfont

#### 4.2 Function MCat()

```
(Draw molecule at mposition)
MCat(c,d)(...) :
mposition:=(c,d); MC(....)
c: x axis position
d: y axis position
defaultsize:=5bp;
fsize:=(60mm,40mm); fmargin:=(3mm,3mm);
blength:=0.07; sw_frame:=Outside;
mangle:=0;
for i=1 step -0.5 until 0:
  for j=0 step 0.33 until 1:
    MCat(j,i)(Ph,4:N)
    add(drawarrow((A1+A1up**aw)..A1);
        label(decimal(mangle),
              p0+(0.5w,0.5h));
    )
    mangle:=mangle+30;
  endfor
endfor
```



#### 4.3 Function check()

(immediately compile)

```
beginfont("EN:Pyridine")
  MC(<30,Ph,2:N)
endfont

(check mcf and compile)

** check(mc) : error count

beginfont("EN:Pyridine",
  ":<30,Ph,}2:N") % ** extra '}'
  if check(mc)=0: MC(scantokens(mc)) fi
endfont</pre>
```





check(mc)=0 check(mc)>=1

#### 4.4 Function add() drawarrow A1{A1down}..A6; dotlabel.urt( "B3e",B3e); (Add label to molecule) label.rt( "An="&decimal(An)& " Bn="&decimal(Bn)& molecular width " aw="&decimal(aw)& molecular height h: " em="&decimal(em), atom font size p0+(-9em,-1.5em));label font size label.rt( "w="&decimal(w)& p0: origin of molecular structure " h="&decimal(h)& bond length 1: " l="&decimal(1), p0+(-9em,-3em));An: atom number ) A[m]: atom position endfont A[m]ang: branch angle of A[m] A[m]up: dir A[m]ang •p0+(w,h) A[m]left: dir A[m]ang+90 ВЗе A[m]right: dir A[m]ang-90 A[m]down: dir A[m]ang+180 R3m Bn: bond number **B3s** B[m]: bond(path) р0 B[m]s: bond start position An=6 Bn=6 aw=11.00174 em=8 B[m]m: bond middle position w=49.09756 h=67.69455 l=28.3464 B[m]e: bond end position B[m]ang: bond angle beginfont("EN:add() 2") B[m]up: dir B[m]ang fsize:=(60mm,20mm); B[m]left: dir B[m]ang+90 msize:=(1,0.85);B[m]right: dir B[m]ang-90 %-----B[m]down: dir B[m]ang+180 MCat(0,0)(<30,Ph,3=d1,4:/NH2)%----plus : '+' circled add( minus : '-' circled labeloffset:=.7aw; circlediam = 0.6aw (default) label.top(lone\_pair 90,A7); circlepen = 0.2bp (default) drawarrow (A7+up\*\*1.2aw){A7left} lonepair r: ':' rotated r ..{B7right}B7/\*0.3; lonepairdiam = 0.3aw (default) drawarrow lonepairspace = 0.7aw (default) B3m..A3+B2up\*\*1.5aw..{A3down}A3; \*\* : scaled << : rotated $MCat(1,0)(<30,?6,{1,5}=d1,4://NH2)$ a /\* b : point b of a %----add( beginfont("EN:add() 1") labeloffset:=.7aw; fsize:=(70mm,40mm);label.top(plus,A7); sw\_frame:=Bothside; label.urt(minus,A3); max\_blength:=10mm; label(lonepair A3ang,A3+A3up\*\*.7aw); msize:=(.91,.9); $MCat(.5,.85)(<30,?6,\{2,5\}:0)$ %----add( ext(drawdblarrow (.4w,.4h)..(.55w,.4h);) defaultscale:=.8; %----labeloffset:=.3aw; endfont dotlabel.lft("p0",p0); dotlabel.rt( "p0+(w,h)",p0+(w,h)); dotlabel.ulft("A1",A1); drawarrow A1..A1+\_\_\*l<<A1ang;</pre> dotlabel.lrt( "B3s",B3s);

dotlabel.rt("B3m",B3m);

dotlabel.ulft("A6",A6);

drawarrow B3m..B3m+\_\_\*1<<(B3ang+90);

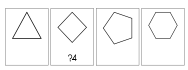
#### 4.5 Function ext()

```
(Extra label to font)
     font width
w:
     font height
h:
w0:
     font width-2xpart(fmargin)
     font height-2ypart(fmargin)
     atom font size
aw:
     label font size
em:
p0:
     fmargin
     molecular number
n:
p[m]: molecular origin position
w[m]: molecular width
h[m]: molecular height
ratio_thickness_char:
pen thickness / char width
%-----
beginfont()
fsize:=(70mm,30mm;);
 blength:=0.065;
%-----
MCat(0.1,0.5)(
  <-210,60'1,60'1,60'1,{1,3}=d1,
  1:/R1,4:/R2^-60
  add(
    defaultscale:=0.6;
    label.bot("Diene",p0+(0.5w,0));
  )
MCat(0.4,0.5)(
  <-30,-60'1,1=d1,1:/R3,2:/R4<sup>60</sup>)
  add(defaultscale:=0.6;
  label.bot("Dienophile",p0+(.5w,0));
 )
MCat(0.9, 0.5)
   <30,?6,6=d1,2:/R2,3:/R4,4:/R3,5:/R1
 %-----
 drawarrow (.52w,.5h)..(.6w,.5h);
 defaultscale:=0.7;
 label("+",(0.25w,0.5h));
 ratio_thickness_char:=0.125;
 label.bot("Diels-Alder Reaction",
           (.5w,h));
endfont
```

# Diets-Alder Reaction R1 + R3 + Dienophite Diene

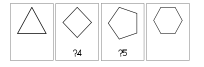
#### 4.5.1 Local ext() setting

```
beginfont("EN:?3")
  fsize:=(12mm,15mm);
  MCat(0.5,1)(<30,?3)
endfont
beginfont("EN:?4")
  fsize:=(12mm, 15mm);
  MCat(0.5,1)(?4)
  ext(label.top(inf_EN,(0.5w,0));)
endfont
beginfont("EN:?5")
  fsize:=(12mm,15mm);
  MCat(0.5,1)(?5)
endfont
beginfont("EN:?5")
  fsize:=(12mm,15mm);
  MCat(0.5,1)(?6)
endfont
```



#### 4.5.2 Global ext() setting

ext\_clear: reset global ext() beginfont("EN:?3") fsize:=(12mm,15mm); MCat(0.5,1)(<30,?3)endfont ext(label.top(inf\_EN,(0.5w,0));) %----beginfont("EN:?4") fsize:=(12mm,15mm); MCat(0.5,1)(?4)endfont beginfont("EN:?5") fsize:=(12mm, 15mm); MCat(0.5,1)(?5)endfont %----ext\_clear; %----beginfont("EN:?6") fsize:=(12mm, 15mm); MCat(0.5,1)(?6)endfont



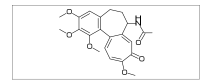
#### MCF example 5

#### Luciferin 5.1

(use data base file 'mcf\_data\_base') beginfont("f:mcf\_data\_base", "t:EN", "v:Luciferin") fsize:=(50mm,15mm); if check(mc)=0: MC(scantokens(mc)) fi endfont

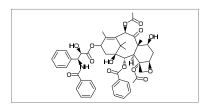
#### Colchicine 5.2

beginfont("EN:Colchicine", "MW:385.41", %-----": <30,Ph,{1,2,6}:/0!,-4=?7,-5=?7, ", ":  $\{-1,-4,-6\}=d1,-2://0,-3:/0!,$ ": @9,\,NH,!,//O,! fsize:=(50mm,20mm); if check(mc)=0: MC(scantokens(mc)) fi endfont



#### Paclitaxel 5.3

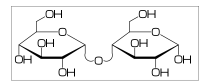
beginfont("EN:Paclitaxel", "MW:853.91", %-----": ?6,5=d1,@3,#1,36,45,45,45,45,##, ", ": -4=?6, -4=?4, -1=wb, -3=wf, -1:0, ||, ": 4:??,6:/\_,{3^-60,15}:\*/OH, ": 8:/\*H^-60,", ": 9:\*/\_^60,10://0, ": @1,\,0,!,//0,!,\*/OH,!,/Ph,60~wf, ": NH,-60,//0,60,Ph, ":  $@7,\*,0,-45,//0,60,Ph,$ ": @11,\*\,0,-60,//0,60, ": @12,\\*^-15,0,60,//0,-60 %----fsize:=(50mm,25mm); if check(mc)=0: MC(scantokens(mc)) fi endfont.



#### 5.4 Maltose

(bond type for glycan) arc\_lb : arc left > bottom arc\_br : arc bottom right

beginfont("EN:Maltose", "MW:342.3", ": #1.25,-30~wf\_r,30~bd\_r'1,30~wb\_r, ": 120,0,30,&1,##, ": {1^\$-90,2^\$90,3^\$-90}:/OH'-.5, ": 6^\$90:/!OH'-.5, ": @4,\$-50~arc\_lb'1,0,\$50~arc\_br'1,<\$0, ": |,#1.25,-30~wf\_r,30~bd\_r'1,30~wb\_r, 120,0,30,&1,##, ": {2^\$90,3^\$-90,4^\$-90}:/OH'-.5, ": 6^\$90:/!OH'-.5 fsize:=(50mm,20mm); if check(mc)=0: MC(scantokens(mc)) fi endfont



#### Cellobiose 5.5

(bond type for glycan)

arc\_lbr : arc left > bottom > right arc\_ltr : arc left > top > right

beginfont("EN:Cellobiose","MW:342.3",

```
": #1.25,-30~wf_r,30~bd_r'1,30~wb_r,
": 120,0,30,&1,##,
": {1^$-90,2^$90,3^$-90}:/OH'-.5,
": 6^$90:/!OH'-.5,
": @4,$0~arc_ltr,0,$0~arc_lbr,
": |,#1.25,-30~wf_r,30~bd_r'1,30~wb_r,
     120,0,30,&1,##,
": {2^$90,3^$-90,4^$-90}:/OH'-.5,
": 6^$90:/!OH'-.5
%-----
fsize:=(50mm,20mm);
if check(mc)=0: MC(scantokens(mc)) fi
```

endfont

# 6 Example to use mcf2graph

#### 6.1 Metafont/Metapost souce file

```
%-----
input mcf2graph.mf;
                                       > input main macro
sw_output:=Info; % aux(information) file output on > global setting
                                       > report output
%%%% sw_output:=Report;
%%%% sw_output:=MOL2k;
                                       > MOL file output
fsize:=(60mm,40mm); % (font width,font height)
tag1:="J";
                                       > jobname
tag2:="C";
                                       > char No
% calculated molecular formula >
outputformat:="png"; hppp:=vppp:=0.1;
outputtemplate:="%j-%3c.png";
%-----
beginfont("EN:Ampicillin", "MW:349.405")
                                      > information
 MC(<45,?4,-3=?5,2:N,7:S,
                                       > immediately compile
  3<sup>45</sup>:/*H,1://0<sup>15</sup>,5:/*COOH<sup>-18</sup>,6:??,
  @4,*\^15,NH,!,//O,!,/*NH2,!,Ph)
endfont
beginfont("EN:Cholesterol", "MW:386.65",
                                      >information
 %-----
 ": <30,?6,{-4,-2}=?6,-4=?5,7=d1, ",
                                       > mc1
 ": 10:/*H^180,11:/*H^-60,17:/*H^-54, ",
                                       > mc2
 ": {4,12}:*/_^60,
                                      > mc3
 ": @-1,18,/*_,-60,!3,?!
                           ")
                                      > mc4
 if check(mc)=0: MC(scantokens(mc)) fi
                                       > mc=mc1 - mc4
%-----
beginfont("f:mcf_data_base.mcf","t:EN","v:Adenine") > from mcf_data_base.mcf
 if check(mc)=0: MC(scantokens(mc)) fi
                                       > select EN="Adenine"
endfont
%-----
beginfont("t:EN","v:Guanine")
                                       > select EN="Guanine"
 if check(mc)=0: MC(scantokens(mc)) fi
endfont.
%-----
beginfont("t:EN","v:Cytosine")
                                       > select EN="Cytosine"
 if check(mc)=0: MC(scantokens(mc)) fi
endfont
%-----
                                     > v+:4 = select No.4
beginfont("t:n","v+:4")
 if check(mc)=0: MC(scantokens(mc)) fi
                                      > keep file open
endfont
%-----
forever:
%%%%%%%%% beginfont("f:mcf_data_base","v+:*") > select all beginfont("f:mcf_data_base","t:EXA","v+:1") > 'v+:1'= select EXA=1
   if f_EOF=0: if check(mc)=0: MC(scantokens(mc)) fi fi > keep file open
 exitif (f_EOF=1)or(f_close=1);
                                       > exit if file end
%-----
bye
```

#### 6.2 Molecular data base file

```
% molecular data base file mcf_data_base.mcf by Akira Yamaji 2021.04.18
% tag1:var1;tag2:var2;tag3:var3 ....
% first character of line '%' comment out
\% first character of line '+' begin MCF , end MCF
Cat:biological; EN:Adenine; MW:135.13
<30,?6,3=?5,{1,3,5,9}=d1,{2,6,9}:N,5:/NH2,7:NH
Cat:biological; EN:Guanine; MW:151.13
<30,?6,3=?5,{1,3,9}=d1,{2,9}:N,{6,7}:NH,5://0,1:/NH2
Cat:biological; EN:Cytosine; MW:111.10
<30,?6,\{4,6\}=d1,4:N,3://0,2:NH,5:/NH2
Cat:biological; EN:Thymine; MW:126.11
<30,?6,3=d1,{2,6}:NH,{1,5}://0,4:/_
Cat:biological; EN:Uracil; MW:112.09
<30,?6,6=d1,{3,5}://0,{2,4}:NH
Cat:biological; EN:Geraniol; MW:154.25
<30,!8,OH,{2,6}=dr,{2,6}:/_
Cat:biological; EN:Limonene; MW:136.24
<30,?6,2=d1,2:/_,@5,*\,/_,!!
Cat:biological; EN:1-Menthol; MW:156.27
<30,?6,2:/*?!,5:*/_,3:*/OH
+-----
Cat:biological; EN:Vanillin; MW:152.15
<30,Ph,2:/OH,3:/O!,5:/CHO
Cat:biological; EN:Allicin; MW:162.28
<-30,!!,!2,S0,!,S,!2,!!
Cat:biological; EN:Stearic acid; MW:284.48
<30,!17,COOH
+-----
Cat:biological; EN:Linoleic acid; MW:280.45
<30,!5,-30,-30,!,-30,-30,!7,COOH,{6,9}=dr
```

## 6.3 Function query()

```
(Example)
```

```
%-----
% query()
% "f:filename" : input file name (default "mcf_data_base.mcf")
% "o:filename" : output file name (default "temp.mcf")
% "s:sort-key" : sort by sort-key
% operator : = , <> , <= , >= , < , >
% filter 1 : Cat=biological
% filter 2 : MW>=285
\% filter 3 : MW<=295
query("s:EN",
      "f:mcf_data_base.mcf","o:temp.mcf","s:EN",
    "Cat=biological","MW>=285","MW<=295");
%-----
 if f_EOF=0: if check(mc)=0: MC(scantokens(mc)) fi fi
 {\tt endfont}
 exitif f_EOF=1;
endfor
%-----
(output)
Cat:biological; EN:Atoropin; MW:289.375; EXA:1
<30,0,!,//0,!,!,Ph,@$1,\~zb^-60,|,?7'1.1,@6,*\^190'1.25,N!,&3~wb,$3:/!OH~wv
Cat:biological; EN:Cianidanol; MW:290.27; EXA:1
<30,Ph,3=?6,@8,*\,Ph,7:0,{1,5,13,14}:/OH,9:/*OH
Cat:biological; EN:Lycorine; MW:287.315; EXA:1
<30,Ph,-4=?6,-2=?6,6=?5,(9,12)=?5[3],13=d1,8:N,{15,17}:0,
9:/*H^180,10:*/H^60,13:*/OH,14:/*OH
Cat:biological; EN:Morphine; MW:285.343; EXA:1
<30,Ph,2=?6,-4=?6,(1,12)=?5[2],-1:0,-1=zb,
@7,60 wf `0.75,70 si_`1.3,45,N!,&9 wb,15=dl,6:/OH,8^180:*/H,12:/*OH
Cat:biological;EN:Piperine;MW:285.343;EXA:1
<30,Ph,|,-1=?5,{1,3}:0,0$4,\,!!,!!,!,!,//0,!,|,?6,1:N
```

#### 6.4 Information aux file output

```
(Insert option parameter setting)
```

```
sw_output:=Info;
                             %% tag1:var1;tag2:var2
 sw_output:=Info+Table;
                             %% tag1;tag2 var1;var2
 sw_output:=Info+Temp;
                             %% tag1:var1;tag2:var2 / output 'temp-info.aux'
                             %% output jobname&'.aux'
 sw_output:=Info+Mcode;
 sw_output:=Info+Mcode+Temp; %% output 'temp-info.aux','temp-mc.aux'
 (Command line)
 >mpost -s ahlength=1 FILENAME (sw_output=Info)
 >mpost -s ahlength=2 FILENAME (sw_output=Info+Table)
(Sourse)
beginfont("EN:Ampicillin")
                             .... endfont
beginfont("EN:Cholesterol")
                           .... endfont
beginfont("EN:Limonin")
                         .... endfont
beginfont("EN:beta-Carotene") .... endfont
(Setting)
tag1:="J"; tag2:="C"; tag3:="mw"; tag4:="fm"; tag5:="EN";
(Output)
(sw_output=Info)
F:mcf_man_soc;C:1;mw:349.40462;fm:C16H19N304S;EN:Ampicillin
F:mcf_man_soc;C:2;mw:386.6532;fm:C27H460;EN:Cholesterol
F:mcf_exa_soc;C:3;mw:470.5113;fm:C26H3008;EN:Limonin
F:mcf_exa_soc; C:4; mw:536.8722; fm:C40H56; EN:beta-Carotene
(sw_output=Info+Table)
F;C;mw;fm
mcf_man_soc;1;349.40462;C16H19N3O4S;Ampicillin
mcf_man_soc;2;386.6532;C27H460;Cholesterol
mcf_exa_soc;3;470.5113;C26H3008;Limonin
mcf_exa_soc;4;536.8722;C40H56;beta-Carotene
(aux_delimiter="/")
F:mcf_man_soc/C:1/mw:349.40462/fm:C16H19N3O4S/EN:Ampicillin
F:mcf_man_soc/C:2/mw:386.6532/fm:C27H460/EN:Cholesterol
F:mcf_exa_soc/C:3/mw:470.5113/fm:C26H3008/EN:Limonin
F:mcf_exa_soc/C:4/mw:536.8722/fm:C40H56/EN:beta-Carotene
(Tag)
J
   : jobname
   : char number
NO : serial number
EN : english name
JN : japanese name
FM : formula from literature data
MW : molecular weight from literature data
MI : monoisotopic mass from literature data
USE : the use
mw : molecular weight calculated
mi : monoisotopic mass calculated
fm : molecular formula calculated
w : font width
h
  : font height
```

#### 6.5 Metafont aux file output

```
(Insert option parameter setting)
sw_output:=Mfont;
(Command line)
```

```
>mpost -s ahlength=7 FILENAME (sw_output=Mfont)
```

#### (Output)

```
beginfont("Cat:biological","EN:Adenine","MW:135.13",
": <30,?6,3=?5,{1,3,5,9}=d1,{2,6,9}:N,5:/NH2,7:NH")
if check(mc)=0: MC(scantokens(mc)) fi
endfont
beginfont("Cat:biological", "EN:Guanine", "MW:151.13",
": <30,?6,3=?5,{1,3,9}=d1,{2,9}:N,{6,7}:NH,5://0,1:/NH2")
if check(mc)=0: MC(scantokens(mc)) fi
endfont
beginfont("Cat:biological", "EN:Cytosine", "MW:111.10",
": <30,?6,\{4,6\}=d1,4:N,3://0,2:NH,5:/NH2")
if check(mc)=0: MC(scantokens(mc)) fi
beginfont("Cat:biological", "EN:Thymine", "MW:126.11",
": <30,?6,3=d1,{2,6}:NH,{1,5}://0,4:/_")
if check(mc)=0: MC(scantokens(mc)) fi
beginfont("Cat:biological", "EN:Adenine", "MW:135.13",
": <30,?6,3=?5,{1,3,5,9}=d1,{2,6,9}:N,5:/NH2,7:NH")
if check(mc)=0: MC(scantokens(mc)) fi
endfont
beginfont("Cat:biological", "EN:Guanine", "MW:151.13",
": <30,?6,3=?5,\{1,3,9\}=d1,\{2,9\}:N,\{6,7\}:NH,5://0,1:/NH2")
if check(mc)=0: MC(scantokens(mc)) fi
endfont
beginfont("Cat:biological","EN:Cytosine","MW:111.10",
": <30,?6,\{4,6\}=d1,4:N,3://0,2:NH,5:/NH2")
if check(mc)=0: MC(scantokens(mc)) fi
endfont
beginfont("Cat:biological", "EN:Thymine", "MW:126.11",
": <30,?6,3=d1,{2,6}:NH,{1,5}://0,4:/_")
if check(mc)=0: MC(scantokens(mc)) fi
endfont.
beginfont("Cat:biological", "EN:Uracil", "MW:112.09",
": <30,?6,6=d1,{3,5}://0,{2,4}:NH")
if check(mc)=0: MC(scantokens(mc)) fi
endfont
beginfont("Cat:biological", "EN:Geraniol", "MW:154.25",
": <30,!8,OH,{2,6}=dr,{2,6}:/_")
if check(mc)=0: MC(scantokens(mc)) fi
endfont
beginfont("Cat:biological", "EN:Limonene", "MW:136.24",
": <30,?6,2=d1,2:/_,@5,*\,/_,!!")
if check(mc)=0: MC(scantokens(mc)) fi
beginfont("Cat:biological", "EN:1-Menthol", "MW:156.27",
": <30,?6,2:/*?!,5:*/_,3:*/OH")
if check(mc)=0: MC(scantokens(mc)) fi
endfont
```

#### 6.6 MCF aux file output

#### (Insert option parameter setting)

```
%% output 'jobname-nnn-EN-mc.aux'
sw_output:=Mcode;
                              %% output 'temp-mc.aux'
sw_output:=Mcode+Temp;
sw_output:=Info+Mcode;
                              %% output 'jobname-data.aux'
                              \% output 'temp-info.aux', 'temp-mc.aux'
sw_output:=Info+Mcode+Temp;
                              %% output font,'temp-mc.aux'
sw_output:=Font+Mcode+Temp;
                              %% output font,'temp-info.aux','temp-mc.aux'
sw_output:=Font+Info+Mcode+Temp;
(Command line)
 >mpost -s ahlength=8 FILENAME (sw_output=Info+Mcode)
(Output temporary file)
sw_output=Mcode
                   ** file name = 'jobname-nnn-EN-mc.aux'
sw_output=Mcode+Temp ** file name = 'temp-mc.aux'
(result)
<30,?6,3=?5,{1,3,5,9}=d1,{2,6,9}:N,5:/NH2,7:NH
(Output data-base file)
sw_output=Mcode+Info ** file name = 'jobname-data.aux'
(result)
Cat:biological; EN:Adenine; MW:135.13; EXA:1
<30,?6,3=?5,{1,3,5,9}=d1,{2,6,9}:N,5:/NH2,7:NH
(Lualatex example)
beginfont("t:EN","v:Adenine")
 sw_output:=Mcode+Temp;
endfont
%-----
\begin{mplibcode}
 beginfont("t:EN","v:Vancomycin")
   sw_output:=Mcode+Temp; %%%% output temp-mc.aux %%%%
 endfont;
\end{mplibcode}
%-----
\verbatiminput{temp-mc.aux}
%-----
(result)
file name = 'temp-mc.aux'
<30,?6,@4,?6,@-4,\,!3,<-12,?5,@-3,<-12,?6,-3=?6,@-3,*\,!3,
?6,0-4,?6,06,\,!,/*Me^-40,*/OH^20,!,//0,!1,OH,
3=wb,11=d1,15=dr,17=wf,19=wf,38=wb,\{5,7,16,24,25,33,42\}:0,
32:*/H^60,10:/Me,{12,31}:*/_,27://_,37:/*_,28:/OH,{3,29}:/*OH
```

#### Report output

```
(Insert option parameter setting)
                       ** file name = 'jobname-report.aux'
  sw_output:=Report;
```

```
(Command line)
```

>mpost -s ahlength=3 FILENAME

Fomula Calc: C4H5N3O / Input:

sw\_output:=Report+Temp; \*\* file name = 'temp-report.aux'

#### (Output)

```
No. 3 / Name = Cytosine
 <30,?6,{4,6}=d1,4:N,3://0,2:NH,5:/NH2
_____
row= 1 / length= 37 / commands= 7
 {} = 1 / {}: X = 0 / '() = 0 / 0 = 0 / & = 0 / < = 1
 Warnings = 0 / Code= 60
 Width * Height = 34.68852 *
                                    47.4036
Shift width * height = 0 * -14.46167
Bond length = 12.75589 Atom size = 5.38914
Atom count= 9 Bond count= 9 Ring count= 1 Hide H count= 2
______
< NO. ><atom(s) >( x axis , y axis )<bond><hideH><chg>
       C ( 0, 0) 3
N ( 0.866, -0.5) 3
C ( 1.732, 0) 4
N ( 1.732, 1) 3
 A1
A2 N ( 0.866 ,
A3 C ( 1.732 ,
A4 N ( 1.732 ,
A5 C ( 0.866 ,
A6 C ( 0 ,
A7 O ( 2.508 , -0
A8 H ( 0.866 ,
A9 NH2 ( 0.866 ,
 A2
                                        1.5 )
1 )
                                                   3
                                     -0.448 )
                                                   2
                                      -0.922 )
                                                      1
                                        2.371 )
                                                      1
< NO. >< bond (sdt)><angle +( +- )><length ( pt )>
     1 -> 2 ( 1) 330 ( -30) 1 ( 12.76)
B1
      2 -> 3 ( 1) 30 ( 30) 1 ( 12.76)

3 -> 4 ( 1) 90 ( 90) 1 ( 12.76)

4 -> 5 ( 2) 150 ( 150) 1 ( 12.76)

5 -> 6 ( 1) 210 ( -150) 1 ( 12.76)

6 -> 1 ( 2) 270 ( -90) 1 ( 12.76)

3 -> 7 ( 2) 330 ( -30) 0.66 ( 8.42)

2 -> 8 ( 1) 270 ( -90) 0.36 ( 4.59)
B2
ВЗ
В4
B5
B6
B7
      2 -> 8 ( 1) 270 ( -90)
5 -> 9 ( 1) 90 ( 90)
                                               0.36 (
В8
                                                            4.59)
В9
                                              0.66 (
                                                            8.42)
______
<atom>( atom wt )[ mi wt ] < cnt > < sum wt >[ sum mi wt ]
C ( 12.0107)[ 12] * 4 48.04279[ 48]
H ( 1.00793)[ 1.00783] * 5 5.03967[ 5.03914]
N ( 14.0067)[ 14.00307] * 3 42.0201[ 42.0092]
O ( 15.9994)[ 15.99492] * 1 15.9994[ 15.99492]
Molecular Weight [Mono Isotopic] = 111.1019[ 111.04326]
 Weight Calc: 111.1019 / Input: 111.10 / weight gap= 0.00195
```

\_\_\_\_\_

#### 6.8 MOL file output

#### (Insert option parameter setting)

#### (Command line)

#### (Output)

```
14 15 0 0 0 0 0 0 0 0999 V2000
      0
           0
                     0 C
                               0
           -0.5
                     O N
  0.86603
                         0 0 0 0
                   0 C
  1.73206
            0
                         0 0 0 0
                    0 C
  1.73206
             1
                         0 0 0
                               0
  0.86603
           1.5
                    0 C
                         0 0 0
      0
            1
                    O N
                         0 0 0
                               0
  2.6831 -0.30902
                    O N
                         0 0 0 0
                   0 0 0 0 0
  3.27089 0.5
  2.6831 1.30902
                   0 N 0 0 0 0
                   0 0 0 0 0
  0.86603 -1.36383
 -0.76894 1.44394
                   0 0 0 0 0
 -0.76894 -0.44394
                   0 0 0 0 0 0
  0.86603 2.36383
                    000000
  2.95299 2.1396
                  0 0 0 0 0
         0 0
 1 2 1 0
 2 3 1 0
            0 0
 3 4 2 0
            0 0
 4 5 1 0
           0 0
 5 6 1 0
           0 0
 6
  1 1 0
            0 0
 3
   7 1 0
            0 0
 7 8 2 0
           0 0
 8 9 1 0
           0 0
 9 4 1 0
           0 0
 2 10 1 0
           0 0
 6 11 1 0
           0 0
 1 12 2 0
           0 0
 5 13
     2 0
            0 0
 9 14
     1 0
            0 0
M END
```

#### 6.9 LuaTeX file example

```
\documentclass{article}
\usepackage{luamplib}%
\usepackage[T1]{fontenc}%
\usepackage{textcomp}%
\mplibcodeinherit{enable}%
\mplibverbatim{enable}%
\mplibnumbersystem{double}%
\everymplib{%
 if unknown Ph1: input mcf2graph.mf; fi
 sw_output:=Font; max_blength:=4.5mm;
 defaultfont:="uhvr8r"; defaultsize:=8bp; defaultscale:=1;
}%
\begin{document}
\noindent%
%-----
               _____
\begin{mplibcode}
 fsize:=(50mm,50mm);
 beginfont("NO:1", "EN:Limonin", "MW:470.51",
   %-----
   ": <30,?6,{-3,-4}=?6,
   ": -5=?3, -2=wf, -1=wb, 6=?5, -4=?6, -5=wf, ",
   ": {13,15,17,20}:0,{3,12,21}://0, ",
   ": {4~wf^60,8~zf^60,18^35,18^-35}:/_, ",
   ": {1^60,5^180,16^60}:/*H,
   ": @14,\*,|,?5,{1,4}=d1,3:0
 if check(mc)=0: MC(scantokens(mc)) fi
 {\tt endfont}
\end{mplibcode}\\
%-----
\begin{mplibcode}
 fsize:=(80mm,50mm);
 beginfont("NO:2", "EN:beta-carotene", "MW:536.87",
   %-----
   ": <30,?6,3=d1,{3,5^35,5^-35}:/_, ",
   ": 04,\,|,!18,{1,3,5,7,9,11,13,15,17}=dr, ",
   ": {3,7,12,16}:/_,
   ": |,?6,6=d1,{6,2^35,2^-35}:/_
   %-----
 if check(mc)=0: MC(scantokens(mc)) fi
 endfont
\end{mplibcode}\\
%-----
\begin{mplibcode}
 fsize:=(50mm,50mm);
 beginfont("NO:3", "EN:Gibberellin A3", "MW:346.37",
   ": <18,?5,3=?7,5=?6[12], ",
   ": @8,160'1.3,&3,13=d1,6=wf,8=wb, ",
   ": @5,40~zf'1,0,60,//0~180,&14~zb, ",
   ": 2:/COOH,7://_,13:*/OH,8:/*OH, ",
   ": 14:*/_,{1^60,4^60}:*/H
   %-----
 if check(mc)=0: MC(scantokens(mc)) fi
endfont;
%-----
\end{mplibcode}\\
\end{document}
```

#### 6.10 LaTeX file example

```
_____
\documentclass[a4paper]{article}
\usepackage{graphicx}
\makeatletter%
\usepackage{mcf_setup}
%-----
\pagestyle{empty}
%-----
\def\put@char{%
 \begin{picture}(84,42)%
    \put(0,38){\bf [\MOLnum]\EN{ }\small\tt/FM:\fm/MW:\mw}%
    \put(10,0){\font\@strufont=\File\relax%
             \hbox{\@strufont\char\Char}}%
 \end{picture}%
}%
\def\INFO#1{\@for\@temp:=#1\do{\tag@var\@temp}\put@char}%
\makeatother
\begin{document}
\unitlength=1mm%
\INFO{J:mcf_man_soc,C:141,NO:1,mw:349.40462,fm:C16H19N304S,EN:Ampicillin}%
\INFO{J:mcf_man_soc,C:142,NO:2,mw:386.6532,fm:C27H460,EN:Cholesterol}%
\end{document}
%-----
```

#### [1]Ampicillin

FM:C16H19N3O4S MW:349.40462

#### [2]Cholesterol

FM:C27H460 MW:386.6532

```
NO = 144
EN = Cholesterol
MW(D) = 386.65
MW(C) =
FM(C) =
w = 204.09412
h = 240.94444
n = 1
ratio_thickness_bond = 0.015
ratio_atom_bond = 0.36
ratio_bondgap_bond = 0.15
ratio_chain_ring = 0.66
sw_trimming = 0
mc1 = <30,?6,{-4,-2}=?6,-4=?5,7=dl,
mc2= 10:/*HA180,11:/*HA-60,17:/*HA-54,
mc3= 1:*/OH,{4,12}:*/_^60,@-1,18,/*_,-60,!3,?!
```