MCF example for luamplib(LualATEX)

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FM(C): molecular formula calculated by mcf2graphMW(C): molecular weight calculated by mcf2graphMW(D): molecular weight from literature data

(Chlorophyll a)

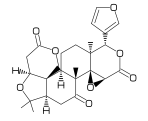
FM(C): C55H72MgN4O5 MW(C): 893.4884 MW(D): 893.49 |<, ''1,?5,{2,5}=d1,4:N,3:\,54~d1,
|,?5,{2,4}=d1,5:N,
-2:\,54~d1,|,?5,2=d1,5:N,
-2:\~d1,54,|,?5,5=d1,5:N,-2:\~d1,\$5:&,
-1:@,24,/*C00!^15,72,//0,\$1:&,>|,
4:\'1.45,Mg,17:&,-1:@,11~vb:&,-1:@,23~vb:&,
{2,9,15,20~zf}:/_,8:/!,14:\,!!,
21:@,-6~wf,!2,//0,!,0,!2,!!,
|,!13,{1,5,9,13}:/_,

(beta-Carotene)

FM(C): C40H56 MW(C): 536.8722 MW(D): 536.888 <30,?6,3=d1,{3,5^35,5^-35}:/_, 4:\,|,!18, {1,3,5,7,9,11,13,15,17}=dr, {3,7,12,16}:/_, |,?6,6=d1,{6,2^35,2^-35}:/_

(Limonin)

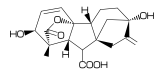
FM(C): C26H30O8 MW(C): 470.5113 MW(D): 470.51



<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</pre>

(Gibberellin A3)

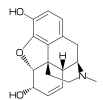
FM(C): C19H22O6 MW(C): 346.3742 MW(D): 346.37



<18,75,3=?7,5=?6[12],8:0,160'1.3,3:&, 13=dl,6=wf,8=wb, 5:0,40~zf'1,0,60,//0^180,14~zb:&, 2:/C00H,7://_,13:*/OH,8:/*OH, 14:*/_,{1^60,4^60}:*/H)

(Morphine)

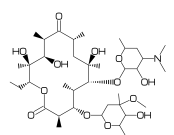
FM(C): C17H19NO3 MW(C): 285.3375 MW(D): 285.343



<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

(Erythromycin)

FM(C): C37H67NO13 MW(C): 733.9263 MW(D): 733.93



<30, |<, '1, <-120,60,60,60,-60,60, 60,-60,60,60,60,-60,60,60,>|,1:&, 14:0,13:/*Et,{1,9}://0,{2,10}:*/_, {4,6^-35,8,12^35}:/*_, {6^35,11,12^-35}:*/0H, \$3:*,0,30,|,?6'.7,2:0, {3,5^35}:/_,4:/OH,5^-35:/0!, \$5:*^30'1.7,0,!,|,?6'.7,6:0, 5:/_,2:/OH,3:/NMeMe

(Paclitaxel)

FM(C): C47H51NO14 MW(C): 853.9057

MW(D): 853.918

?6,5=d,3:0,<''1,36,45,45,45,45,>|,\$5:&,-4=?6, -4=?4, ||, -1=wb, -3=wf, -1:0,

{4^35,4^-35,6}:/_,{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)

(Vancomycin)

FM(C): C66H75Cl2N9O24

MW(C): 1449.253

<30, |<, ''1, !12, {1,3,12}=zf,7=wf,

/H^-60,60,*/OH,60,Ph,-4:/Cl,

-3:\,0,!,Ph,-4:\,0,!,Ph,-1^15:/Cl,

-3:\,/*OH,*/H^-60,\$1:&,

\$7:0,\$26:&,\$1:0,120,//0,60,NH,60,

/*H,*/COOH^180,-60,

 $Ph, \{-2, -4\}: /OH, -1: \, Ph, -5: /OH, -2: @, $4: &, >|,$

{3^40,6,9,12}://0,{2,5,8,11}:NH,

{1^180,4^180}:*/H,

 $\{7^-60, 10^60, 14^60\}: /*H,$

\$10:*\^-60,60,//0,!,NH2,\$13:*\,NH,!,//0,!,

/!iPr^-35>60,*/H^60 ,!~zf,NH,!,

\$23:\,0,!,|,?6'.7,2:0,3^10:/!OH,{4,5}:/OH,

 $-1:\,0,!,|,?6'.7,6:0,$

{3^35,5}:/_,3^-35:/NH2,4:/OH

FM(C): C164H256Na2O68S2

MW(C): 3425.854 MW(D): 3425.86

<55.8,?6,-4=?7,{-4,-3,-3,-3}=?6,

 $-3:\,!3,?6,\{-4,-3,-3,-3\}=?6,-3:\,?6,-3=?6,-3:\,!3,60,<-30,?6,-3=?6,$

-3:0,30,<30,?6,{-3,-3}=?6,-3=?7,{-4,-3,-3}=?6,

 $-2:\,?6,-3=?6,-3=?7,\{-3,-3\}=?6,-3=?8,-3=d1,\{-5,-3,-3,-3\}=?6,$

{5,7,15,16,23,24,32,40,41,48,49,58,59,72,73,82,83,90,91,99,

100,107,113,114,122,123,130,131,140,141,148,149}:0,

{1^60,2,26,28,29,51,54,61,63,68,75^60,78,109}:*/OH,

 $\{11,20,35,45,52,55,65,69,86\}:/*OH,\{47,57,71\}:/*H^60,$

 $\{3,8,13,17,21,33,38,42,56,70,84,92,101,106,111,128,138,142,146,150\}:/*H^-60,$

{4,14,22,34,39,43,81,89,98,102,116,121,125,129,133}:*/H^60,

{6,46,50,53,60,67,74}:*/H^-60,{9,18,85,93,112,139,143,147}:*/_'1^60,

{80,88,97,115,120,124}:/*_'1^-60,108:*/_'1^-60,

\$6:\,|,!11,60~dr,-60,60,0H,2:/*OH,{7,10}:*/OH,{1,3}:*/_,{8~zf,11~dm,12}:/_,

 $6:\,0,30,S00,30,"0{Na}",$

\$36:@,-45~zf,0,30,S00,30,"O{Na}",\$150:\,|,!7,{1,2}:/*OH,4:*/_,5:/*_,7=dl

```
НÓ
                             Oxaloacetate
    L-Malate
            NAD+
                                                                                cis-Aconitate
                                            CoA-SH
                                                                        H<sub>2</sub>O
                                                        Citrate
                  NADH2<sup>+</sup>
                                     Acetyl-CoA
                                                                               H_2O
                                       TCA-cycle
    Fumarate
                                                                                 Isocitrate
                                                                              NAD+
        FADH2
                                            NAD+,CoA-SH
                                                                            NADH2+
                     GDP,Pi
         ΑD
                                       NADH2<sup>+</sup>,CO<sub>2</sub>
                                                                     CO_2
             GTP,CoA-SH
    Succinate
                           Succinyl-CoA
                                                    alfa-Ketoglutarate
                                                                               Oxalosuccinate
beginfont("EN:TCA cycle")
font_wd:=160mm;
font_ht:=75mm;
max_bond_length:=5mm;
Om := '("{O^-}");
MCa(0.33,
            1)(<30,0m,!0,//0,!,//0,!2,//0,!,0m)
MCa(0.66,
             1) (<30,0m,!0,//0,!4,//0,!,0m,-4'1:\,,//0,!,0m,4:/0H^-165)
MCa(1,
             1) (<30,0m,!0,//0,!2,!~dr,!,//0,!,0m,-4'1:\,,//0,!,0m)
MCa(1.
         0.55) (<30,0m,!0,//0,!4,//0,!,0m,-4:\'1,//0,!,0m)
MCa(1,
         0.05) (<30,0m,!0,//0,!3,//0,!,//0,!,0m,-4:\'1,//0,!,0m)
MCa(0.66,0.05)(<30,0m,!0,//0,!3,//0,!,//0,!,0m)
MCa(0.33,0.05)(<30,0m,!0,//0,!3,//0,!,"{S-CoA}")
MCa(0,
         0.05) (<30,0m,!0,//0,!3,//0,!,0m)
         0.55) (<30,0m,!0,//0,!,!~dr,!,//0,!,0m)
MCa(0,
             1)(<30,0m,!0,//0,!3,//0,!,0m,3:/OH)
MCa(0,
EXT(
defaultfont:="uhvr8r";
defaultscale:=0.75;
ext_setup;
save dx; pair dx; dx := (12mm, 0);
label.bot("Oxaloacetate",p1+dx);
                                     label.bot("Citrate",p2+dx);
label.bot("cis-Aconitate",p3+dx);
                                     label.bot("Isocitrate",p4+dx);
label.bot("Oxalosuccinate",p5+dx); label.bot("alfa-Ketoglutarate",p6+dx);
label.bot("Succinyl-CoA",p7+dx);
                                     label.bot("Succinate",p8+dx);
label.bot("Fumarate",p9+dx);
                                     label.bot("L-Malate",p10+dx);
sw_label_emu:=1;
ext_setup;
r_arrow(10mm)( 0)(p1+(1.1w1,.3h1))("",0)("",0)("Acetyl-CoA",1.5)(" CoA-SH",1);
r_arrow(10mm)( 0)(p2+(1.1w2,.4h2))("",0)("",0)("",0)("H_2_0",1);
r_arrow( 8mm)(270)(p3+(.5w3,-.4h3))("",0)("",0)("H_2_0",1)("",0);
r_arrow( 8mm)(270)(p4+(.5w4,-.4h4))("",0)("",0)("NAD^++",1)("NADH2^+",1);
r_arrow(10mm)(180)(p5+(-.1w5,.4h5))("",0)("",0)("",0)("CO_2_",1);
r_arrow(10mm)(180)(p6+(-.1w6,.5h6))("",0)("",0)("NAD^+^,CoA-SH",1.7)("NADH2^+^,CO_2",1);
r_arrow(10mm)(180)(p7+(-.1w7,.5h7))("",0)("",0)("GDP,Pi",1.7)("GTP,CoA-SH",1);
r_arrow( 8mm)( 90)(p8+(.4w8,1.2h8))("",0)("",0)("FAD",1)("FADH2",1);
r_arrow( 8mm)( 90)(p9+(.4w9,1.2h9))("",0)("",0)("H_2_0",1)("",0);
r_arrow(10mm)( 0)(p10+(1.1w10,.3h10))("",0)("",0)("NAD^++",1)("NADH2^+",1.5);
defaultscale:=1.5;
label("TCA-cycle",(0.5w,0.5h));
)
endfont
```