

The Chemical Mechanism of MECCA

KPP version: 2.2.3_rs3

MECCA version: 4.4.2

Date: 2023-08-25

MECCA config (*.ini) file: delhi.ini

Integrator: rosenbrock_posdef

Gas equation file: gas.eqn

Replacement file:

Selected reactions:

“Tr and (G or (Aa or Mb1)) and not I and not Hg”

Number of aerosol phases: 1

Number of species in selected mechanism:

Gas phase: 2761

Aqueous phase: 1703

All species: 4464

Number of reactions in selected mechanism:

Gas phase (Gnnn): 1742

Aqueous phase (Annn): 386

Henry (Hnnn): 715

Photolysis (Jnnn): 348

Aqueous phase photolysis (PHnnn): 26

Heterogeneous (HETnnn): 0

Equilibria (EQnn): 112

Isotope exchange (IEXnnn): 0

Tagging equations (TAGnnn): 0

Dummy (Dnn): 1

All equations: 3330

Table 1: Gas phase reactions

#	labels	reaction	rate coefficient	reference
G1000	UpStTrG	$O_2 + O(^1D) \rightarrow O(^3P) + O_2$	3.3E-11*EXP(55./temp)	Burkholder et al. (2015)
G1001	UpStTrG	$O_2 + O(^3P) \rightarrow O_3$	6.0E-34*((temp/300.)**(-2.4)) *cair	Burkholder et al. (2015)
G2100	UpStTrG	$H + O_2 \rightarrow HO_2$	k_3rd(temp,cair,4.4E-32,1.3, 7.5E-11,-0.2,0.6)	Burkholder et al. (2015)
G2104	UpStTrG	$OH + O_3 \rightarrow HO_2 + O_2$	1.7E-12*EXP(-940./temp)	Burkholder et al. (2015)
G2105	UpStTrG	$OH + H_2 \rightarrow H_2O + H$	2.8E-12*EXP(-1800./temp)	Burkholder et al. (2015)
G2107	UpStTrG	$HO_2 + O_3 \rightarrow OH + 2 O_2$	1.E-14*EXP(-490./temp)	Burkholder et al. (2015)
G2109	UpStTrG	$HO_2 + OH \rightarrow H_2O + O_2$	4.8E-11*EXP(250./temp)	Burkholder et al. (2015)
G2110	UpStTrG	$HO_2 + HO_2 \rightarrow H_2O_2 + O_2$	k_HO2_HO2	Burkholder et al. (2015)*
G2111	UpStTrG	$H_2O + O(^1D) \rightarrow 2 OH$	1.63E-10*EXP(60./temp)	Burkholder et al. (2015)
G2112	UpStTrG	$H_2O_2 + OH \rightarrow H_2O + HO_2$	1.8E-12	Burkholder et al. (2015)
G2117	UpStTrG	$H_2O + H_2O \rightarrow (H_2O)_2$	6.521E-26*temp*EXP(1851.09/temp) *EXP(-5.10485E-3*temp)	Scribano et al. (2006)*
G2118	UpStTrG	$(H_2O)_2 \rightarrow H_2O + H_2O$	1.E0	see note*
G3101	UpStTrGN	$N_2 + O(^1D) \rightarrow O(^3P) + N_2$	2.15E-11*EXP(110./temp)	Burkholder et al. (2015)
G3103	UpStTrGN	$NO + O_3 \rightarrow NO_2 + O_2$	3.0E-12*EXP(-1500./temp)	Burkholder et al. (2015)
G3106	StTrGN	$NO_2 + O_3 \rightarrow NO_3 + O_2$	1.2E-13*EXP(-2450./temp)	Burkholder et al. (2015)
G3108	StTrGN	$NO_3 + NO \rightarrow 2 NO_2$	1.5E-11*EXP(170./temp)	Burkholder et al. (2015)
G3109	UpStTrGN	$NO_3 + NO_2 \rightarrow N_2O_5$	k_NO3_NO2	Burkholder et al. (2015)*
G3110	StTrGN	$N_2O_5 \rightarrow NO_2 + NO_3$	k_NO3_NO2/(5.8E-27*EXP(10840./ temp))	Burkholder et al. (2015)*
G3200	TrGN	$NO + OH \rightarrow HONO$	k_3rd(temp,cair,7.0E-31,2.6, 3.6E-11,0.1,0.6)	Burkholder et al. (2015)
G3201	UpStTrGN	$NO + HO_2 \rightarrow NO_2 + OH$	3.3E-12*EXP(270./temp)	Burkholder et al. (2015)
G3202a	UpStTrGN	$NO_2 + OH \rightarrow HNO_3$	(1.-alpha_HOONO) * k_NO2_OH	Amedro et al. (2020)
G3202b	UpStTrGN	$NO_2 + OH \rightarrow HOONO$	alpha_HOONO * k_NO2_OH	Amedro et al. (2020)
G3203	StTrGN	$NO_2 + HO_2 \rightarrow HNO_4$	k_NO2_HO2	Burkholder et al. (2015)*
G3204	TrGN	$NO_3 + HO_2 \rightarrow NO_2 + OH + O_2$	3.5E-12	Burkholder et al. (2015)
G3205	TrGN	$HONO + OH \rightarrow NO_2 + H_2O$	1.8E-11*EXP(-390./temp)	Burkholder et al. (2015)
G3206	StTrGN	$HNO_3 + OH \rightarrow H_2O + NO_3$	k_HN03_OH	Dulitz et al. (2018)*
G3207	StTrGN	$HNO_4 \rightarrow NO_2 + HO_2$	k_NO2_HO2/(2.1E-27*EXP(10900./ temp))	Burkholder et al. (2015)*
G3208	StTrGN	$HNO_4 + OH \rightarrow NO_2 + H_2O$	1.3E-12*EXP(380./temp)	Burkholder et al. (2015)

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G3209	TrGN	$\text{NH}_3 + \text{OH} \rightarrow \text{NH}_2 + \text{H}_2\text{O}$	1.7E-12*EXP(-710./temp)	Kohlmann and Poppe (1999)
G3210	TrGN	$\text{NH}_2 + \text{O}_3 \rightarrow \text{NH}_2\text{O} + \text{O}_2$	4.3E-12*EXP(-930./temp)	Kohlmann and Poppe (1999)
G3211	TrGN	$\text{NH}_2 + \text{HO}_2 \rightarrow \text{NH}_2\text{O} + \text{OH}$	4.8E-07*EXP(-628./temp)*(temp) **(-1.32)	Kohlmann and Poppe (1999)
G3212	TrGN	$\text{NH}_2 + \text{HO}_2 \rightarrow \text{HNO} + \text{H}_2\text{O}$	9.4E-09*EXP(-356./temp)*(temp) **(-1.12)	Kohlmann and Poppe (1999)
G3213	TrGN	$\text{NH}_2 + \text{NO} \rightarrow \text{HO}_2 + \text{OH} + \text{N}_2$	1.92E-12*((temp/298.)**(-1.5))	Kohlmann and Poppe (1999)
G3214	TrGN	$\text{NH}_2 + \text{NO} \rightarrow \text{N}_2 + \text{H}_2\text{O}$	1.41E-11*((temp/298.)**(-1.5))	Kohlmann and Poppe (1999)
G3215	TrGN	$\text{NH}_2 + \text{NO}_2 \rightarrow \text{N}_2\text{O} + \text{H}_2\text{O}$	1.2E-11*((temp/298.)**(-2.0))	Kohlmann and Poppe (1999)
G3216	TrGN	$\text{NH}_2 + \text{NO}_2 \rightarrow \text{NH}_2\text{O} + \text{NO}$	0.8E-11*((temp/298.)**(-2.0))	Kohlmann and Poppe (1999)
G3217	TrGN	$\text{NH}_2\text{O} + \text{O}_3 \rightarrow \text{NH}_2 + \text{O}_2$	1.2E-14	Kohlmann and Poppe (1999)
G3218	TrGN	$\text{NH}_2\text{O} \rightarrow \text{NHOH}$	1.3E3	Kohlmann and Poppe (1999)
G3219	TrGN	$\text{HNO} + \text{OH} \rightarrow \text{NO} + \text{H}_2\text{O}$	8.0E-11*EXP(-500./temp)	Kohlmann and Poppe (1999)
G3220	TrGN	$\text{HNO} + \text{NHOH} \rightarrow \text{NH}_2\text{OH} + \text{NO}$	1.66E-12*EXP(-1500./temp)	Kohlmann and Poppe (1999)
G3221	TrGN	$\text{HNO} + \text{NO}_2 \rightarrow \text{HONO} + \text{NO}$	1.0E-12*EXP(-1000./temp)	Kohlmann and Poppe (1999)
G3222	TrGN	$\text{NHOH} + \text{OH} \rightarrow \text{HNO} + \text{H}_2\text{O}$	1.66E-12	Kohlmann and Poppe (1999)
G3223	TrGN	$\text{NH}_2\text{OH} + \text{OH} \rightarrow \text{NHOH} + \text{H}_2\text{O}$	4.13E-11*EXP(-2138./temp)	Kohlmann and Poppe (1999)
G3224	TrGN	$\text{HNO} + \text{O}_2 \rightarrow \text{HO}_2 + \text{NO}$	3.65E-14*EXP(-4600./temp)	Kohlmann and Poppe (1999)
G3227	UpStTrGN	$\text{HOONO} \rightarrow \text{NO}_2 + \text{OH}$	(alpha_HOONO*k_NO2_OH) /(3.5E-27*EXP(10135./temp))	see note*
G3228	UpStTrGN	$\text{HOONO} + \text{OH} \rightarrow \text{H}_2\text{O} + \text{NO}_3$	1.3E-12*EXP(380./temp)	Burkholder et al. (2015)*
G4101	StTrG	$\text{CH}_4 + \text{OH} \rightarrow \text{CH}_3 + \text{H}_2\text{O}$	1.85E-20*EXP(2.82*LOG(temp) -987./temp)	Atkinson (2003)
G4102	TrG	$\text{CH}_3\text{OH} + \text{OH} \rightarrow .85 \text{ HCHO} + .85 \text{ HO}_2 + .15 \text{ CH}_3\text{O} + \text{H}_2\text{O}$	6.38E-18*((temp)**2)*EXP(144./temp)	Atkinson et al. (2006)
G4103a	StTrG	$\text{CH}_3\text{O}_2 + \text{HO}_2 \rightarrow \text{CH}_3\text{OOH} + \text{O}_2$	3.8E-13*EXP(780./temp)/(1.+1./498.*EXP(1160./temp))	Atkinson et al. (2006)
G4103b	StTrG	$\text{CH}_3\text{O}_2 + \text{HO}_2 \rightarrow \text{HCHO} + \text{H}_2\text{O} + \text{O}_2$	3.8E-13*EXP(780./temp)/(1.+498.*EXP(-1160./temp))	Atkinson et al. (2006)
G4104a	StTrGN	$\text{CH}_3\text{O}_2 + \text{NO} \rightarrow \text{CH}_3\text{O} + \text{NO}_2$	2.3E-12*EXP(360./temp)*(1.-beta_CH3NO3)	Atkinson et al. (2006), Butkovskaya et al. (2012), Flocke et al. (1998)
G4104b	StTrGN	$\text{CH}_3\text{O}_2 + \text{NO} \rightarrow \text{CH}_3\text{ONO}_2$	2.3E-12*EXP(360./temp)*beta_CH3NO3	Atkinson et al. (2006), Butkovskaya et al. (2012), Flocke et al. (1998)*

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G4105	TrGN	$\text{CH}_3\text{O}_2 + \text{NO}_3 \rightarrow \text{CH}_3\text{O} + \text{NO}_2 + \text{O}_2$	1.2E-12	Atkinson et al. (2006)
G4106a	StTrG	$\text{CH}_3\text{O}_2 \rightarrow \text{CH}_3\text{O} + .5 \text{ O}_2$	$7.4\text{E}-13*\text{EXP}(-520./\text{temp})*\text{R02*2.}$	Atkinson et al. (2006)
G4106b	StTrG	$\text{CH}_3\text{O}_2 \rightarrow .5 \text{ HCHO} + .5 \text{ CH}_3\text{OH} + .5 \text{ O}_2$	$(k_{\text{CH3O}_2}-7.4\text{E}-13*\text{EXP}(-520./\text{temp}))$ $*\text{R02*2.}$	Atkinson et al. (2006)
G4107	StTrG	$\text{CH}_3\text{OOH} + \text{OH} \rightarrow .6 \text{ CH}_3\text{O}_2 + .4 \text{ HCHO} + .4 \text{ OH} + \text{H}_2\text{O}$	$k_{\text{CH3OOH_OH}}$	Wallington et al. (2018)
G4108	StTrG	$\text{HCHO} + \text{OH} \rightarrow \text{CO} + \text{H}_2\text{O} + \text{HO}_2$	$9.52\text{E}-18*\text{EXP}(2.03*\text{LOG}(\text{temp})$ $+636./\text{temp})$	Sivakumaran et al. (2003)
G4109	TrGN	$\text{HCHO} + \text{NO}_3 \rightarrow \text{HNO}_3 + \text{CO} + \text{HO}_2$	$3.4\text{E}-13*\text{EXP}(-1900./\text{temp})$	Burkholder et al. (2015)*
G4110	UpStTrG	$\text{CO} + \text{OH} \rightarrow \text{H} + \text{CO}_2$	$(1.57\text{E}-13+\text{cair}*3.54\text{E}-33)$	McCabe et al. (2001)
G4111	TrG	$\text{HCOOH} + \text{OH} \rightarrow \text{CO}_2 + \text{HO}_2 + \text{H}_2\text{O}$	$2.94\text{E}-14*\text{exp}(786./\text{temp})$ $+9.85\text{E}-13*\text{EXP}(-1036./\text{temp})$	Paulot et al. (2011)
G4114	StTrGN	$\text{CH}_3\text{O}_2 + \text{NO}_2 \rightarrow \text{CH}_3\text{O}_2\text{NO}_2$	$k_{\text{NO}_2\text{-CH3O}_2}$	Burkholder et al. (2015)
G4115	StTrGN	$\text{CH}_3\text{O}_2\text{NO}_2 \rightarrow \text{CH}_3\text{O}_2 + \text{NO}_2$	$k_{\text{NO}_2\text{-CH3O}_2}/(9.5\text{E}-29*\text{EXP}(11234./\text{temp}))$	Burkholder et al. (2015)*
G4116	StTrGN	$\text{CH}_3\text{O}_2\text{NO}_2 + \text{OH} \rightarrow \text{HCHO} + \text{NO}_3 + \text{H}_2\text{O}$	3.00E-14	see note*
G4117	StTrGN	$\text{CH}_3\text{ONO}_2 + \text{OH} \rightarrow \text{H}_2\text{O} + \text{HCHO} + \text{NO}_2$	$4.0\text{E}-13*\text{EXP}(-845./\text{temp})$	Atkinson et al. (2006)
G4118	StTrG	$\text{CH}_3\text{O} \rightarrow \text{HO}_2 + \text{HCHO}$	$1.3\text{E}-14*\text{exp}(-663./\text{temp})*\text{c(ind_02)}$	Chai et al. (2014)
G4119a	StTrGN	$\text{CH}_3\text{O} + \text{NO}_2 \rightarrow \text{CH}_3\text{ONO}_2$	$k_{\text{3rd_iupac}}(\text{temp}, \text{cair}, 8.1\text{E}-29,$ $4.5, 2.1\text{E}-11, 0., 0.44)$	Atkinson et al. (2006)
G4119b	StTrGN	$\text{CH}_3\text{O} + \text{NO}_2 \rightarrow \text{HCHO} + \text{HONO}$	$9.6\text{E}-12*\text{EXP}(-1150./\text{temp})$	Atkinson et al. (2006)
G4120a	StTrGN	$\text{CH}_3\text{O} + \text{NO} \rightarrow \text{CH}_3\text{ONO}$	$k_{\text{3rd_iupac}}(\text{temp}, \text{cair}, 2.6\text{E}-29,$ $2.8, 3.3\text{E}-11, 0.6, \text{REAL}(\text{EXP}(-\text{temp}/$ $900.), \text{SP}))$	Atkinson et al. (2006)
G4120b	StTrGN	$\text{CH}_3\text{O} + \text{NO} \rightarrow \text{HCHO} + \text{HNO}$	$2.3\text{E}-12*(\text{temp}/300.)**(0.7)$	Atkinson et al. (2006)
G4121	StTrG	$\text{CH}_3\text{O}_2 + \text{O}_3 \rightarrow \text{CH}_3\text{O} + 2 \text{ O}_2$	$2.9\text{E}-16*\text{exp}(-1000./\text{temp})$	Burkholder et al. (2015)
G4122	StTrGN	$\text{CH}_3\text{ONO} + \text{OH} \rightarrow \text{H}_2\text{O} + \text{HCHO} + \text{NO}$	$1.\text{E}-10*\text{exp}(-1764./\text{temp})$	Nielsen et al. (1991)
G4123	StTrG	$\text{HCHO} + \text{HO}_2 \rightarrow \text{HOCH}_2\text{O}_2$	$9.7\text{E}-15*\text{EXP}(625./\text{temp})$	Atkinson et al. (2006)
G4124	StTrG	$\text{HOCH}_2\text{O}_2 \rightarrow \text{HCHO} + \text{HO}_2$	$2.4\text{E}12*\text{EXP}(-7000./\text{temp})$	Atkinson et al. (2006)
G4125	StTrG	$\text{HOCH}_2\text{O}_2 + \text{HO}_2 \rightarrow .5 \text{ HOCH}_2\text{OOH} + .5 \text{ HCOOH} + .2 \text{ OH} + .2 \text{ HO}_2 + .3 \text{ H}_2\text{O} + .8 \text{ O}_2$	$5.6\text{E}-15*\text{EXP}(2300./\text{temp})$	Atkinson et al. (2006)
G4126	StTrGN	$\text{HOCH}_2\text{O}_2 + \text{NO} \rightarrow \text{NO}_2 + \text{HO}_2 + \text{HCOOH}$	$0.7275*2.3\text{E}-12*\text{EXP}(360./\text{temp})$	Atkinson et al. (2006)*
G4127	StTrGN	$\text{HOCH}_2\text{O}_2 + \text{NO}_3 \rightarrow \text{NO}_2 + \text{HO}_2 + \text{HCOOH}$	1.2E-12	see note*
G4129a	StTrG	$\text{HOCH}_2\text{O}_2 \rightarrow \text{HCOOH} + \text{HO}_2$	$(k_{\text{CH3O}_2}*5.5\text{E}-12)**(0.5)*\text{R02*2.}$	Atkinson et al. (2006)
G4129b	StTrG	$\text{HOCH}_2\text{O}_2 \rightarrow .5 \text{ HCOOH} + .5 \text{ HOCH}_2\text{OH} + .5 \text{ O}_2$	$(k_{\text{CH3O}_2}*5.7\text{E}-14*\text{EXP}(750./\text{temp}))$ $**(0.5)*\text{R02*2.}$	Atkinson et al. (2006)

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G4130a	StTrG	$\text{HOCH}_2\text{OOH} + \text{OH} \rightarrow \text{HOCH}_2\text{O}_2 + \text{H}_2\text{O}$	k_{ROOHRO}	Taraborrelli (2010)*
G4130b	StTrG	$\text{HOCH}_2\text{OOH} + \text{OH} \rightarrow \text{HCOOH} + \text{H}_2\text{O} + \text{OH}$	$k_{\text{ROHRO}} + k_{\text{s*f_sOOH*f_sOH}}$	Taraborrelli (2010)*
G4132	StTrG	$\text{HOCH}_2\text{OH} + \text{OH} \rightarrow \text{HO}_2 + \text{HCOOH} + \text{H}_2\text{O}$	$2.*k_{\text{ROHRO}} + k_{\text{s*f_sOH*f_sOH}}$	Taraborrelli (2010)*
G4133	StTrG	$\text{CH}_3\text{O}_2 + \text{OH} \rightarrow \text{CH}_3\text{O} + \text{HO}_2$	1.4E-10	Bossolasco et al. (2014)*
G4134	StTrG	$\text{CH}_2\text{OO} \rightarrow \text{CO} + \text{HO}_2 + \text{OH}$	$1.124\text{E+14*EXP(-10000./temp)}$	see note*
G4135	StTrG	$\text{CH}_2\text{OO} + \text{H}_2\text{O} \rightarrow \text{HOCH}_2\text{OOH}$	$k_{\text{CH2OO_NO2}} * 3.6\text{E-6}$	Ouyang et al. (2013)*
G4136	StTrG	$\text{CH}_2\text{OO} + (\text{H}_2\text{O})_2 \rightarrow \text{HOCH}_2\text{OOH} + \text{H}_2\text{O}$	5.2E-12	Chao et al. (2015), Lewis et al. (2015)*
G4137	StTrGN	$\text{CH}_2\text{OO} + \text{NO} \rightarrow \text{HCHO} + \text{NO}_2$	6.E-14	Welz et al. (2012)*
G4138	StTrGN	$\text{CH}_2\text{OO} + \text{NO}_2 \rightarrow \text{HCHO} + \text{NO}_3$	$k_{\text{CH2OO_NO2}}$	Welz et al. (2012), Stone et al. (2014)*
G4140	StTrG	$\text{CH}_2\text{OO} + \text{CO} \rightarrow \text{HCHO} + \text{CO}_2$	3.6E-14	Vereecken et al. (2012)
G4141	StTrG	$\text{CH}_2\text{OO} + \text{HCOOH} \rightarrow 2 \text{HCOOH}$	1.E-10	Welz et al. (2014)*
G4142	StTrG	$\text{CH}_2\text{OO} + \text{HCHO} \rightarrow 2 \text{LCARBON}$	1.7E-12	Stone et al. (2014)*
G4143	StTrG	$\text{CH}_2\text{OO} + \text{CH}_3\text{OH} \rightarrow 2 \text{LCARBON}$	5.E-12	Vereecken et al. (2012)*
G4144	StTrG	$\text{CH}_2\text{OO} + \text{CH}_3\text{O}_2 \rightarrow 2 \text{LCARBON}$	5.E-12	Vereecken et al. (2012)*
G4145	StTrG	$\text{CH}_2\text{OO} + \text{HO}_2 \rightarrow \text{LCARBON}$	5.E-12	Vereecken et al. (2012)
G4146	StTrG	$\text{CH}_2\text{OO} + \text{O}_3 \rightarrow \text{HCHO} + 2 \text{O}_2$	1.E-12	Vereecken et al. (2014)
G4147	StTrG	$\text{CH}_2\text{OO} + \text{CH}_2\text{OO} \rightarrow 2 \text{HCHO} + \text{O}_2$	6.E-11	Buras et al. (2014)
G4148	StTrGN	$\text{HOCH}_2\text{O}_2 + \text{NO}_2 \rightarrow \text{HOCH}_2\text{O}_2\text{NO}_2$	$k_{\text{NO2_CH3O2}}$	see note*
G4149	StTrGN	$\text{HOCH}_2\text{O}_2\text{NO}_2 \rightarrow \text{HOCH}_2\text{O}_2 + \text{NO}_2$	$k_{\text{NO2_CH3O2}} / (9.5\text{E-29*EXP(11234./temp)})$	Barnes et al. (1985)*
G4150	StTrGN	$\text{HOCH}_2\text{O}_2\text{NO}_2 + \text{OH} \rightarrow \text{HCOOH} + \text{NO}_3 + \text{H}_2\text{O}$	$9.50\text{E-13*EXP(-650./temp)} * f_{\text{sOH}}$	see note*
G4151	StTrG	$\text{CH}_3 + \text{O}_2 \rightarrow \text{CH}_3\text{O}_2$	$k_{\text{3rd_iupac}}(\text{temp}, \text{cair}, 7.0\text{E-31}, 3., 1.8\text{E-12}, -1.1, 0.33)$	Atkinson et al. (2006)
G4152	StTrG	$\text{CH}_3 + \text{O}_3 \rightarrow .956 \text{HCHO} + .956 \text{H} + .044 \text{CH}_3\text{O} + \text{O}_2$	$5.1\text{E-12*exp(-210./temp)}$	Albaladejo et al. (2002), Ogryzlo et al. (1981)
G4153	StTrG	$\text{CH}_3 + \text{O}({}^3\text{P}) \rightarrow .83 \text{HCHO} + .83 \text{H} + .17 \text{CO} + .17 \text{H}_2 + .17 \text{H}$	1.3E-10	Atkinson et al. (2006)
G4154	StTrG	$\text{CH}_3\text{O} + \text{O}_3 \rightarrow \text{CH}_3\text{O}_2 + \text{O}_2$	2.53E-14	Albaladejo et al. (2002)*
G4155	StTrG	$\text{CH}_3\text{O} + \text{O}({}^3\text{P}) \rightarrow .75 \text{CH}_3 + .75 \text{O}_2 + .25 \text{HCHO} + .25 \text{OH}$	2.5E-11	Baulch et al. (2005)
G4156	StTrG	$\text{CH}_3\text{O}_2 + \text{O}({}^3\text{P}) \rightarrow \text{CH}_3\text{O} + \text{O}_2$	4.3E-11	Zellner et al. (1988)
G4157	StTrG	$\text{HCHO} + \text{O}({}^3\text{P}) \rightarrow .7 \text{OH} + .7 \text{CO} + .3 \text{H} + .3 \text{CO}_2 + \text{HO}_2$	$3.4\text{E-11*EXP(-1600./temp)}$	Burkholder et al. (2015)

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G4158	TrG	$\text{CH}_2\text{OO}^* \rightarrow .37 \text{ CH}_2\text{OO} + .47 \text{ CO} + .47 \text{ H}_2\text{O} + .16 \text{ HO}_2 + .16 \text{ CO} + .16 \text{ OH}$	KDEC	Atkinson et al. (2006)
G4159	TrGN	$\text{HCN} + \text{OH} \rightarrow \text{H}_2\text{O} + \text{CN}$	$k_{\text{3rd}}(\text{temp}, \text{cair}, 4.28\text{E}-33, 1.0, \text{REAL}(4.25\text{E}-13*\text{EXP}(-1150./\text{temp}), \text{SP}), 1.0, 0.8)$	Kleinböhl et al. (2006)
G4160a	TrGN	$\text{HCN} + \text{O}({}^1\text{D}) \rightarrow \text{O}({}^3\text{P}) + \text{HCN}$	$1.08\text{E}-10*\text{EXP}(105./\text{temp}) * 0.15*\text{EXP}(200./\text{temp})$	Strekowski et al. (2010)
G4160b	TrGN	$\text{HCN} + \text{O}({}^1\text{D}) \rightarrow \text{H} + \text{NCO}$	$1.08\text{E}-10*\text{EXP}(105./\text{temp}) * 0.68/2.$	Strekowski et al. (2010)*
G4160c	TrGN	$\text{HCN} + \text{O}({}^1\text{D}) \rightarrow \text{OH} + \text{CN}$	$1.08\text{E}-10*\text{EXP}(105./\text{temp}) * (1. - (0.68/2. + 0.15*\text{EXP}(200./\text{temp})))$	Strekowski et al. (2010)*
G4161	TrGN	$\text{HCN} + \text{O}({}^3\text{P}) \rightarrow \text{H} + \text{NCO}$	$1.0\text{E}-11*\text{EXP}(-4000./\text{temp})$	Burkholder et al. (2015)*
G4162	TrGN	$\text{CN} + \text{O}_2 \rightarrow \text{NCO} + \text{O}({}^3\text{P})$	$1.2\text{E}-11*\text{EXP}(210./\text{temp}) * 0.75$	Baulch et al. (2005)
G4163	TrGN	$\text{CN} + \text{O}_2 \rightarrow \text{CO} + \text{NO}$	$1.2\text{E}-11*\text{EXP}(210./\text{temp}) * 0.25$	Baulch et al. (2005)
G4164	TrGN	$\text{NCO} + \text{O}_2 \rightarrow \text{CO}_2 + \text{NO}$	$7.\text{E}-15$	Becker et al. (2000)*
G42000	TrGC	$\text{C}_2\text{H}_6 + \text{OH} \rightarrow \text{C}_2\text{H}_5\text{O}_2 + \text{H}_2\text{O}$	$1.49\text{E}-17*\text{temp}*\text{temp}*\text{EXP}(-499./\text{temp})$	Atkinson et al. (2006)
G42001	TrGC	$\text{C}_2\text{H}_4 + \text{O}_3 \rightarrow \text{HCHO} + \text{CH}_2\text{OO}^*$	$9.1\text{E}-15*\text{EXP}(-2580./\text{temp})$	Atkinson et al. (2006)*
G42002	TrGC	$\text{C}_2\text{H}_4 + \text{OH} \rightarrow \text{HOCH}_2\text{CH}_2\text{O}_2$	$k_{\text{3rd_iupac}}(\text{temp}, \text{cair}, 8.6\text{E}-29, 3.1, 9.\text{E}-12, 0.85, 0.48)$	Atkinson et al. (2006), Rickard (2022)
G42003	TrGC	$\text{C}_2\text{H}_5\text{O}_2 + \text{HO}_2 \rightarrow \text{C}_2\text{H}_5\text{OOH}$	$7.5\text{E}-13*\text{EXP}(700./\text{temp})$	Burkholder et al. (2015)
G42004a	TrGCN	$\text{C}_2\text{H}_5\text{O}_2 + \text{NO} \rightarrow \text{CH}_3\text{CHO} + \text{HO}_2 + \text{NO}_2$	$2.55\text{E}-12*\text{EXP}(380./\text{temp}) * (1. - \beta_{\text{C}_2\text{H}_5\text{NO}_3})$	Atkinson et al. (2006), Butkovskaya et al. (2010)
G42004b	TrGCN	$\text{C}_2\text{H}_5\text{O}_2 + \text{NO} \rightarrow \text{C}_2\text{H}_5\text{ONO}_2$	$2.55\text{E}-12*\text{EXP}(380./\text{temp}) * \beta_{\text{C}_2\text{H}_5\text{NO}_3}$	Atkinson et al. (2006), Butkovskaya et al. (2010)
G42005	TrGCN	$\text{C}_2\text{H}_5\text{O}_2 + \text{NO}_3 \rightarrow \text{CH}_3\text{CHO} + \text{HO}_2 + \text{NO}_2$	$2.3\text{E}-12$	Wallington et al. (2018)
G42006	TrGC	$\text{C}_2\text{H}_5\text{O}_2 \rightarrow .8 \text{ CH}_3\text{CHO} + .6 \text{ HO}_2 + .2 \text{ C}_2\text{H}_5\text{OH}$	$2.*(.76\text{E}-14*k_{\text{CH3O2}})**(.5)*R02$	Sander et al. (2019), Atkinson et al. (2006)
G42007a	TrGC	$\text{C}_2\text{H}_5\text{OOH} + \text{OH} \rightarrow \text{C}_2\text{H}_5\text{O}_2 + \text{H}_2\text{O}$	k_{ROOHRO}	Sander et al. (2019)
G42007b	TrGC	$\text{C}_2\text{H}_5\text{OOH} + \text{OH} \rightarrow \text{CH}_3\text{CHO} + \text{OH}$	$k_{\text{s*f_sooh}}$	Sander et al. (2019)
G42008a	TrGC	$\text{CH}_3\text{CHO} + \text{OH} \rightarrow \text{CH}_3\text{C(O)} + \text{H}_2\text{O}$	$4.4\text{E}-12*\text{EXP}(365./\text{temp}) * 0.95$	Atkinson et al. (2006)
G42008b	TrGC	$\text{CH}_3\text{CHO} + \text{OH} \rightarrow \text{HCOCH}_2\text{O}_2 + \text{H}_2\text{O}$	$4.4\text{E}-12*\text{EXP}(365./\text{temp}) * 0.05$	Atkinson et al. (2006)
G42009	TrGCN	$\text{CH}_3\text{CHO} + \text{NO}_3 \rightarrow \text{CH}_3\text{C(O)} + \text{HNO}_3$	KN03AL	Rickard (2022)
G42010	TrGC	$\text{CH}_3\text{COOH} + \text{OH} \rightarrow \text{CH}_3 + \text{CO}_2 + \text{H}_2\text{O}$	$k_{\text{CH3CO2H_OH}}$	Atkinson et al. (2006)*
G42011a	TrGC	$\text{CH}_3\text{C(O)OO} + \text{HO}_2 \rightarrow \text{OH} + \text{CH}_3 + \text{CO}_2$	$5.20\text{E}-13*\text{EXP}(980./\text{temp}) * 1.507 * 0.61$	Groß et al. (2014)
G42011b	TrGC	$\text{CH}_3\text{C(O)OO} + \text{HO}_2 \rightarrow \text{CH}_3\text{C(O)OOH}$	$5.20\text{E}-13*\text{EXP}(980./\text{temp}) * 1.507 * 0.23$	Groß et al. (2014)

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G42011c	TrGC	$\text{CH}_3\text{C}(\text{O})\text{OO} + \text{HO}_2 \rightarrow \text{CH}_3\text{COOH} + \text{O}_3$	5.20E-13*EXP(980./temp)*1.507*0.16	Groß et al. (2014)
G42012	TrGCN	$\text{CH}_3\text{C}(\text{O})\text{OO} + \text{NO} \rightarrow \text{CH}_3 + \text{CO}_2 + \text{NO}_2$	8.1E-12*EXP(270./temp)	Tyndall et al. (2001a)
G42013	TrGCN	$\text{CH}_3\text{C}(\text{O})\text{OO} + \text{NO}_2 \rightarrow \text{PAN}$	k_CH3C03_N02	Burkholder et al. (2015)*
G42014	TrGCN	$\text{CH}_3\text{C}(\text{O})\text{OO} + \text{NO}_3 \rightarrow \text{CH}_3 + \text{NO}_2 + \text{CO}_2$	4.E-12	Canosa-Mas et al. (1996)
G42017a	TrGC	$\text{CH}_3\text{C}(\text{O})\text{OO} \rightarrow \text{CH}_3 + \text{CO}_2$	k1_R02RC03*0.9	Sander et al. (2019)
G42017b	TrGC	$\text{CH}_3\text{C}(\text{O})\text{OO} \rightarrow \text{CH}_3\text{COOH}$	k1_R02RC03*0.1	Sander et al. (2019)
G42018	TrGC	$\text{CH}_3\text{C}(\text{O})\text{OOH} + \text{OH} \rightarrow \text{CH}_3\text{C}(\text{O})\text{OO} + \text{H}_2\text{O}$	k_ROOHRO	Rickard (2022)*
G42020	TrGCN	$\text{PAN} + \text{OH} \rightarrow \text{HCHO} + \text{CO} + \text{NO}_2 + \text{H}_2\text{O}$	3.00E-14	Rickard (2022)
G42021	TrGCN	$\text{PAN} \rightarrow \text{CH}_3\text{C}(\text{O})\text{OO} + \text{NO}_2$	k_PAN_M	Burkholder et al. (2015)*
G42022a	TrGC	$\text{C}_2\text{H}_2 + \text{OH} \rightarrow \text{GLYOX} + \text{OH}$	k_3rd(temp,cair,5.5e-30,0.0, 8.3e-13,-2.,0.6)*0.71	Burkholder et al. (2015)*
G42022b	TrGC	$\text{C}_2\text{H}_2 + \text{OH} \rightarrow \text{HCOOH} + \text{CO} + \text{HO}_2$	k_3rd(temp,cair,5.5e-30,0.0, 8.3e-13,-2.,0.6)*0.29	Burkholder et al. (2015)*
G42023a	TrGC	$\text{HOCH}_2\text{CHO} + \text{OH} \rightarrow \text{HOCH}_2\text{CO} + \text{H}_2\text{O}$	8.00E-12*0.80	Atkinson et al. (2006)
G42023b	TrGC	$\text{HOCH}_2\text{CHO} + \text{OH} \rightarrow \text{HOCHCHO} + \text{H}_2\text{O}$	8.00E-12*0.20	Atkinson et al. (2006)
G42024a	TrGC	$\text{HOCH}_2\text{CO} + \text{O}_2 \rightarrow \text{HOCH}_2\text{CO}_3$	5.1E-12*(1.-1./(1+1.85E-18*cair))	Atkinson et al. (2006), Beyersdorf et al. (2010)*
G42024b	TrGC	$\text{HOCH}_2\text{CO} + \text{O}_2 \rightarrow \text{OH} + \text{HCHO} + \text{CO}_2$	5.1E-12*1./(1+1.85E-18*cair)	Atkinson et al. (2006), Beyersdorf et al. (2010)*
G42025	TrGC	$\text{HOCHCHO} \rightarrow \text{GLYOX} + \text{HO}_2$	KDEC	Sander et al. (2019)
G42026	TrGCN	$\text{HOCH}_2\text{CHO} + \text{NO}_3 \rightarrow \text{HOCH}_2\text{CO} + \text{HNO}_3$	KN03AL	Rickard (2022)
G42027a	TrGC	$\text{HOCH}_2\text{CO}_3 \rightarrow \text{HCHO} + \text{CO}_2 + \text{HO}_2$	k1_R02RC03*0.9	Sander et al. (2019)
G42027b	TrGC	$\text{HOCH}_2\text{CO}_3 \rightarrow \text{HOCH}_2\text{CO}_2\text{H}$	k1_R02RC03*0.1	Sander et al. (2019)
G42028a	TrGC	$\text{HOCH}_2\text{CO}_3 + \text{HO}_2 \rightarrow \text{HCHO} + \text{HO}_2 + \text{OH} + \text{CO}_2$	KAPH02*r_CO3_OH	Sander et al. (2019), Groß et al. (2014)
G42028b	TrGC	$\text{HOCH}_2\text{CO}_3 + \text{HO}_2 \rightarrow \text{HOCH}_2\text{CO}_3\text{H}$	KAPH02*r_CO3_OOH	Sander et al. (2019), Groß et al. (2014)
G42028c	TrGC	$\text{HOCH}_2\text{CO}_3 + \text{HO}_2 \rightarrow \text{HOCH}_2\text{CO}_2\text{H} + \text{O}_3$	KAPH02*r_CO3_O3	Sander et al. (2019), Groß et al. (2014)
G42029	TrGCN	$\text{HOCH}_2\text{CO}_3 + \text{NO} \rightarrow \text{NO}_2 + \text{HO}_2 + \text{HCHO} + \text{CO}_2$	KAPNO	Rickard (2022)
G42030	TrGCN	$\text{HOCH}_2\text{CO}_3 + \text{NO}_2 \rightarrow \text{PHAN}$	k_CH3C03_N02	Rickard (2022)
G42031	TrGCN	$\text{HOCH}_2\text{CO}_3 + \text{NO}_3 \rightarrow \text{NO}_2 + \text{HO}_2 + \text{HCHO} + \text{CO}_2$	KR02N03*1.74	Rickard (2022)
G42032	TrGC	$\text{HOCH}_2\text{CO}_2\text{H} + \text{OH} \rightarrow .09 \text{ HCHO} + .09 \text{ CO}_2 + .91 \text{ HCOCO}_2\text{H} + \text{HO}_2 + \text{H}_2\text{O}$	k_CO2H+k_s*f_sOH*f_CO2H	Sander et al. (2019)
G42033a	TrGC	$\text{HOCH}_2\text{CO}_3\text{H} + \text{OH} \rightarrow \text{HOCH}_2\text{CO}_3 + \text{H}_2\text{O}$	k_ROOHRO	Sander et al. (2019)

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G42033b	TrGC	$\text{HOCH}_2\text{CO}_3\text{H} + \text{OH} \rightarrow \text{HCOCO}_3\text{H} + \text{HO}_2$	$k_{\text{s}} * f_{\text{sOH}} * f_{\text{CO2H}}$	Sander et al. (2019)
G42034	TrGCN	$\text{PHAN} \rightarrow \text{HOCH}_2\text{CO}_3 + \text{NO}_2$	$k_{\text{PAN_M}}$	Rickard (2022)
G42035	TrGCN	$\text{PHAN} + \text{OH} \rightarrow \text{HCHO} + \text{CO} + \text{NO}_2 + \text{H}_2\text{O}$	$k_{\text{s}} * f_{\text{sOH}} * f_{\text{cpa}} + k_{\text{ROHRO}}$	Sander et al. (2019)
G42036	TrGC	$\text{GLYOX} + \text{OH} \rightarrow \text{HCOCO} + \text{H}_2\text{O}$	$3.1\text{E-12} * \text{EXP}(340./\text{temp})$	Atkinson et al. (2006), Orlando and Tyndall (2001), Lockhart et al. (2013)
G42037	TrGCN	$\text{GLYOX} + \text{NO}_3 \rightarrow \text{HCOCO} + \text{HNO}_3$	k_{NO3AL}	Rickard (2022)
G42038a	TrGC	$\text{HCOCO} \rightarrow \text{CO} + \text{CO} + \text{HO}_2$	$7.\text{E}11 * \text{EXP}(-3160./\text{temp}) + 5.\text{E}-12 * c(\text{ind_O2})$	Orlando and Tyndall (2001), Lockhart et al. (2013), Rickard (2022)
G42037b	TrGC	$\text{HCOCO} \rightarrow \text{HCOCO}_3$	$5.\text{E}-12 * c(\text{ind_O2}) * 3.2 * \text{exp}(-550./\text{temp})$	Lockhart et al. (2013), Rickard (2022)
G42037c	TrGC	$\text{HCOCO} \rightarrow \text{OH} + \text{CO} + \text{CO}_2$	$5.\text{E}-12 * c(\text{ind_O2}) * (1. - 3.2 * \text{exp}(-550./\text{temp}))$	Lockhart et al. (2013), Rickard (2022)
G42039a	TrGC	$\text{HCOCO}_3 \rightarrow \text{CO} + \text{HO}_2 + \text{CO}_2$	$k_1_{\text{RO2RC03}} * 0.9$	Sander et al. (2019)
G42039b	TrGC	$\text{HCOCO}_3 \rightarrow \text{HCOCO}_2\text{H}$	$k_1_{\text{RO2RC03}} * 0.1$	Sander et al. (2019)
G42040	TrGC	$\text{HCOCO}_3 + \text{HO}_2 \rightarrow \text{HO}_2 + \text{CO} + \text{CO}_2 + \text{OH}$	$KAPHO2$	Feierabend et al. (2008), Sander et al. (2019)
G42041	TrGCN	$\text{HCOCO}_3 + \text{NO} \rightarrow \text{HO}_2 + \text{CO} + \text{NO}_2 + \text{CO}_2$	$KAPNO$	Rickard (2022)
G42042	TrGCN	$\text{HCOCO}_3 + \text{NO}_3 \rightarrow \text{HO}_2 + \text{CO} + \text{NO}_2 + \text{CO}_2$	$KR02N03 * 1.74$	Rickard (2022)
G42043	TrGCN	$\text{HCOCO}_3 + \text{NO}_2 \rightarrow \text{HO}_2 + \text{CO} + \text{NO}_3 + \text{CO}_2$	$k_{\text{CH3C03_NO2}}$	Orlando and Tyndall (2001), Sander et al. (2019)
G42044	TrGC	$\text{HCOCO}_2\text{H} + \text{OH} \rightarrow \text{CO} + \text{HO}_2 + \text{CO}_2 + \text{H}_2\text{O}$	$k_{\text{CO2H}} + k_t * f_0 * f_{\text{CO2H}}$	Sander et al. (2019)
G42045a	TrGC	$\text{HCOCO}_3\text{H} + \text{OH} \rightarrow \text{HCOCO}_3 + \text{H}_2\text{O}$	k_{ROOHRO}	Sander et al. (2019)
G42045b	TrGC	$\text{HCOCO}_3\text{H} + \text{OH} \rightarrow \text{CO} + \text{CO}_2 + \text{H}_2\text{O} + \text{OH}$	$k_t * f_0 * f_{\text{CO2H}}$	Sander et al. (2019)
G42046	TrGC	$\text{HOCH}_2\text{CH}_2\text{O}_2 \rightarrow .6 \text{ HOCH}_2\text{CH}_2\text{O} + .2 \text{ HOCH}_2\text{CHO} + .2 \text{ ETHGLY}$	$2.* (7.8\text{E-14} * \text{EXP}(1000./\text{temp}) * k_{\text{CH3O2}}) ** (.5) * R02$	Atkinson et al. (2006), Rickard (2022)
G42047	TrGCN	$\text{HOCH}_2\text{CH}_2\text{O}_2 + \text{NO} \rightarrow .25 \text{ HO}_2 + .5 \text{ HCHO} + .75 \text{ HOCH}_2\text{CH}_2\text{O} + \text{NO}_2$	$KR02NO * (1. - \alpha_{\text{AN}}(3, 1, 0, 0, 0, \text{temp}, \text{cair}))$	Rickard (2022)*
G42048	TrGCN	$\text{HOCH}_2\text{CH}_2\text{O}_2 + \text{NO} \rightarrow \text{ETHOHN03}$	$KR02NO * \alpha_{\text{AN}}(3, 1, 0, 0, 0, \text{temp}, \text{cair})$	Sander et al. (2019)
G42049a	TrGC	$\text{HOCH}_2\text{CH}_2\text{O}_2 + \text{HO}_2 \rightarrow \text{HYETHO2H}$	$1.53\text{E-13} * \text{EXP}(1300./\text{temp}) * (1. - r_{\text{CHOHCH202_OH}})$	Rickard (2022)
G42049b	TrGC	$\text{HOCH}_2\text{CH}_2\text{O}_2 + \text{HO}_2 \rightarrow \text{HOCH}_2\text{CH}_2\text{O} + \text{OH}$	$1.53\text{E-13} * \text{EXP}(1300./\text{temp}) * r_{\text{CHOHCH202_OH}}$	Rickard (2022)

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G42050	TrGCN	$\text{ETHOHN}3 + \text{OH} \rightarrow .93 \text{NO}_3\text{CH}_2\text{CHO} + .93 \text{HO}_2 + .07 \text{HOCH}_2\text{CHO} + .07 \text{NO}_2 + \text{H}_2\text{O}$	$k_{\text{s}} * (f_{\text{sOH}} * f_{\text{CH}_2\text{NO}_2} + f_{\text{ON}_2} * f_{\text{pCH}_2\text{OH}}) + k_{\text{ROHRO}}$	Sander et al. (2019)
G42051a	TrGC	$\text{HYETHO}_2\text{H} + \text{OH} \rightarrow \text{HOCH}_2\text{CH}_2\text{O}_2 + \text{H}_2\text{O}$	k_{ROOHR}_0	Rickard (2022)*
G42051b	TrGC	$\text{HYETHO}_2\text{H} + \text{OH} \rightarrow \text{HOCH}_2\text{CHO} + \text{OH} + \text{H}_2\text{O}$	$k_{\text{s}} * f_{\text{sOH}} * f_{\text{pCH}_2\text{OH}}$	Sander et al. (2019)
G42051c	TrGC	$\text{HYETHO}_2\text{H} + \text{OH} \rightarrow \text{HOOCH}_2\text{CHO} + \text{HO}_2 + \text{H}_2\text{O}$	$k_{\text{s}} * f_{\text{sOH}} * f_{\text{pCH}_2\text{OH}} + k_{\text{ROHRO}}$	Sander et al. (2019)
G42052a	TrGC	$\text{HOCH}_2\text{CH}_2\text{O} \rightarrow \text{HO}_2 + \text{HOCH}_2\text{CHO}$	$6.00E-14 * \exp(-550./\text{temp}) * C(\text{ind_O}_2)$	Rickard (2022)
G42052b	TrGC	$\text{HOCH}_2\text{CH}_2\text{O} \rightarrow \text{HO}_2 + \text{HCHO} + \text{HCHO}$	$9.50E13 * \exp(-5988./\text{temp})$	Rickard (2022)
G42053	TrGC	$\text{ETHGLY} + \text{OH} \rightarrow \text{HOCH}_2\text{CHO} + \text{HO}_2 + \text{H}_2\text{O}$	$2.*k_{\text{s}} * f_{\text{sOH}} * f_{\text{pCH}_2\text{OH}} + 2.*k_{\text{ROHRO}}$	Sander et al. (2019)
G42054	TrGC	$\text{HCOCH}_2\text{O}_2 \rightarrow .6 \text{HCHO} + .6 \text{CO} + .6 \text{HO}_2 + .2 \text{GLYOX} + .2 \text{HOCH}_2\text{CHO}$	$k1_{\text{RO2pORO2}}$	Sander et al. (2019)
G42055a	TrGC	$\text{HCOCH}_2\text{O}_2 + \text{HO}_2 \rightarrow \text{HOOCH}_2\text{CHO}$	$k_{\text{RO2_H02}}(\text{temp}, 2) * r_{\text{COCH202_00H}}$	Sander et al. (2019)
G42055b	TrGC	$\text{HCOCH}_2\text{O}_2 + \text{HO}_2 \rightarrow \text{HCHO} + \text{CO} + \text{HO}_2 + \text{OH}$	$k_{\text{RO2_H02}}(\text{temp}, 2) * r_{\text{COCH202_OH}}$	Sander et al. (2019)
G42056a	TrGCN	$\text{HCOCH}_2\text{O}_2 + \text{NO} \rightarrow \text{NO}_2 + \text{HCHO} + \text{CO} + \text{HO}_2$	$\text{KR02NO} * (\text{1.-alpha_AN}(3, 1, 1, 0, 0, temp, cair))$	Sander et al. (2019)
G42056b	TrGCN	$\text{HCOCH}_2\text{O}_2 + \text{NO} \rightarrow \text{NO}_3\text{CH}_2\text{CHO}$	$\text{KR02NO} * \text{alpha_AN}(3, 1, 1, 0, 0, temp, cair)$	Sander et al. (2019)
G42057	TrGCN	$\text{HCOCH}_2\text{O}_2 + \text{NO}_3 \rightarrow \text{HCHO} + \text{CO} + \text{HO}_2 + \text{NO}_2$	KR02NO3	Sander et al. (2019)
G42058a	TrGC	$\text{HOOCH}_2\text{CHO} + \text{OH} \rightarrow \text{HCOCH}_2\text{O}_2$	k_{ROOHR}_0	Sander et al. (2019)
G42058b	TrGC	$\text{HOOCH}_2\text{CHO} + \text{OH} \rightarrow \text{HCHO} + \text{CO} + \text{OH}$	$0.8 * 8.E-12$	Sander et al. (2019)*
G42058c	TrGC	$\text{HOOCH}_2\text{CHO} + \text{OH} \rightarrow \text{GLYOX} + \text{OH}$	$k_{\text{s}} * f_{\text{sOH}} * f_{\text{CHO}}$	Sander et al. (2019)
G42059	TrGCN	$\text{HOOCH}_2\text{CHO} + \text{NO}_3 \rightarrow \text{OH} + \text{HCHO} + \text{CO} + \text{HNO}_3$	KN03AL	Rickard (2022)
G42060	TrGCN	$\text{HOOCH}_2\text{CO}_3 + \text{NO} \rightarrow \text{NO}_2 + \text{OH} + \text{HCHO} + \text{CO}_2$	KAPNO	Sander et al. (2019)
G42061	TrGCN	$\text{HOOCH}_2\text{CO}_3 + \text{NO}_3 \rightarrow \text{NO}_2 + \text{OH} + \text{HCHO} + \text{CO}_2$	$\text{KR02NO3} * 1.74$	Sander et al. (2019)
G42062a	TrGC	$\text{HOOCH}_2\text{CO}_3 + \text{HO}_2 \rightarrow 2 \text{OH} + \text{HCHO} + \text{CO}_2$	$\text{KAPH02} * r_{\text{CO3_OH}}$	Sander et al. (2019)
G42062b	TrGC	$\text{HOOCH}_2\text{CO}_3 + \text{HO}_2 \rightarrow \text{HOOCH}_2\text{CO3H}$	$\text{KAPH02} * r_{\text{CO3_00H}}$	Sander et al. (2019)
G42062c	TrGC	$\text{HOOCH}_2\text{CO}_3 + \text{HO}_2 \rightarrow \text{HOOCH}_2\text{CO2H} + \text{O}_3$	$\text{KAPH02} * r_{\text{CO3_03}}$	Sander et al. (2019)
G42063a	TrGC	$\text{HOOCH}_2\text{CO}_3 \rightarrow \text{OH} + \text{HCHO} + \text{CO}_2$	$k1_{\text{RO2RC03}} * 0.9$	Sander et al. (2019)
G42063b	TrGC	$\text{HOOCH}_2\text{CO}_3 \rightarrow \text{HOOCH}_2\text{CO2H}$	$k1_{\text{RO2RC03}} * 0.1$	Sander et al. (2019)
G42064a	TrGC	$\text{HOOCH}_2\text{CO3H} + \text{OH} \rightarrow \text{HOOCH}_2\text{CO}_3 + \text{H}_2\text{O}$	$2.*k_{\text{ROOHR}}_0$	Sander et al. (2019)
G42064b	TrGC	$\text{HOOCH}_2\text{CO3H} + \text{OH} \rightarrow \text{HCOCO}_3\text{H} + \text{OH} + \text{H}_2\text{O}$	$k_{\text{s}} * f_{\text{sOH}} * f_{\text{C02H}}$	Sander et al. (2019)
G42065	TrGC	$\text{HOOCH}_2\text{CO2H} + \text{OH} \rightarrow \text{HCOCO}_2\text{H} + \text{OH} + \text{H}_2\text{O}$	$k_{\text{s}} * f_{\text{sOH}} * f_{\text{C02H}} + k_{\text{C02H}}$	Sander et al. (2019)
G42066	TrGC	$\text{CH}_2\text{CO} + \text{OH} \rightarrow .6 \text{HCHO} + .6 \text{HO}_2 + .6 \text{CO} + .4 \text{HOOCH}_2\text{CO2H}$	$2.8E-12 * \exp(510./\text{temp})$	Baulch et al. (2005), Sander et al. (2019)
G42067a	TrGC	$\text{CH}_3\text{CHOHOOH} + \text{OH} \rightarrow \text{CH}_3\text{COOH} + \text{OH}$	$(k_t * f_{\text{t00H}} * f_{\text{tOH}} + k_{\text{ROHRO}})$	Sander et al. (2019)

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G42067b	TrGC	$\text{CH}_3\text{CHOHOOH} + \text{OH} \rightarrow \text{CH}_3\text{CHOHO}_2$	k_ROOHR0	Sander et al. (2019)
G42068	TrGC	$\text{CH}_3\text{CHOHO}_2 \rightarrow \text{CH}_3\text{CHO} + \text{HO}_2$	$3.46\text{E}12 * \text{EXP}(-12500. / (1.98 * \text{temp}))$	Hermans et al. (2005), Sander et al. (2019)
G42069	TrGC	$\text{CH}_3\text{CHO} + \text{HO}_2 \rightarrow \text{CH}_3\text{CHOHO}_2$	$3.46\text{E}12 * \text{EXP}(-12500. / (1.98 * \text{temp})) / (6.34\text{E}26 * \text{EXP}(-14700. / (1.98 * \text{temp})))$	Hermans et al. (2005), Sander et al. (2019)
G42070	TrGC	$\text{CH}_3\text{CHOHO}_2 + \text{HO}_2 \rightarrow .5 \text{CH}_3\text{CHOHOOH} + .3 \text{CH}_3\text{COOH} + .2 \text{CH}_3 + .2 \text{HCOOH} + .2 \text{OH}$	$5.6\text{E}-15 * \text{EXP}(2300. / \text{temp})$	Sander et al. (2019)
G42071	TrGC	$\text{CH}_3\text{CHOHO}_2 \rightarrow \text{CH}_3 + \text{HCOOH} + \text{OH}$	k1_R02s0R02	Sander et al. (2019)
G42072	TrGCN	$\text{CH}_3\text{CHOHO}_2 + \text{NO} \rightarrow \text{CH}_3 + \text{HCOOH} + \text{OH} + \text{NO}_2$	KR02NO	Sander et al. (2019)
G42073	TrGCN	$\text{C}_2\text{H}_5\text{ONO}_2 + \text{OH} \rightarrow \text{CH}_3\text{CHO} + \text{H}_2\text{O} + \text{NO}_2$	$6.7\text{E}-13 * \text{EXP}(-395. / \text{temp})$	Atkinson et al. (2006)
G42074a	TrGCN	$\text{NO}_3\text{CH}_2\text{CHO} + \text{OH} \rightarrow \text{GLYOX} + \text{NO}_2 + \text{H}_2\text{O}$	k_s*f_CH2ON02*f_CHO	Paulot et al. (2009a), Sander et al. (2019)*
G42074b	TrGCN	$\text{NO}_3\text{CH}_2\text{CHO} + \text{OH} \rightarrow \text{NO}_3\text{CH}_2\text{CO}_3 + \text{H}_2\text{O}$	k_t*f_0*f_CH2ON02*3.	Paulot et al. (2009a), Sander et al. (2019)*
G42075	TrGCN	$\text{NO}_3\text{CH}_2\text{CO}_3 + \text{HO}_2 \rightarrow \text{HCHO} + \text{NO}_2 + \text{CO}_2 + \text{OH}$	KAPH02	Rickard (2022)*
G42076	TrGCN	$\text{NO}_3\text{CH}_2\text{CO}_3 + \text{NO} \rightarrow \text{HCHO} + \text{NO}_2 + \text{CO}_2 + \text{NO}_2$	KAPNO	Rickard (2022)
G42077	TrGCN	$\text{NO}_3\text{CH}_2\text{CO}_3 + \text{NO}_2 \rightarrow \text{NO}_3\text{CH}_2\text{CHO}$	k_CH3C03_N02	Rickard (2022)
G42078	TrGCN	$\text{NO}_3\text{CH}_2\text{CO}_3 \rightarrow \text{HCHO} + \text{NO}_2 + \text{CO}_2$	k1_R02RC03	Rickard (2022)*
G42079	TrGCN	$\text{NO}_3\text{CH}_2\text{CHO} \rightarrow \text{NO}_3\text{CH}_2\text{CO}_3 + \text{NO}_2$	k_PAN_M	Rickard (2022)
G42080	StTrGCN	$\text{C}_2\text{H}_5\text{O}_2 + \text{NO}_2 \rightarrow \text{C}_2\text{H}_5\text{O}_2\text{NO}_2$	$\text{k_3rd_iupac}(\text{temp}, \text{cair}, 1.3\text{E}-29, 6.2, 8.8\text{E}-12, 0.0, 0.31)$	Atkinson et al. (2006)
G42081	StTrGCN	$\text{C}_2\text{H}_5\text{O}_2\text{NO}_2 \rightarrow \text{C}_2\text{H}_5\text{O}_2 + \text{NO}_2$	$\text{k_3rd_iupac}(\text{temp}, \text{cair}, \text{REAL}(4.8\text{E}-4 * \text{EXP}(-9285. / \text{temp}), \text{SP}), 0.0, \text{REAL}(8.8\text{E}15 * \text{EXP}(-10440. / \text{temp}), \text{SP}), 0.0, 0.31)$	Atkinson et al. (2006)
G42082	StTrGCN	$\text{C}_2\text{H}_5\text{O}_2\text{NO}_2 + \text{OH} \rightarrow \text{CH}_3\text{CHO} + \text{NO}_3 + \text{H}_2\text{O}$	$9.50\text{E}-13 * \text{EXP}(-650. / \text{temp})$	Sander et al. (2019)*
G42083a	TrGC	$\text{CH}_3\text{C(O)} + \text{O}_2 \rightarrow \text{CH}_3\text{C(O)OO}$	$5.1\text{E}-12 * (1. - 1. / (1. + 9.4\text{E}-18 * \text{cair}))$	Atkinson et al. (2006), Beyersdorf et al. (2010)*
G42083b	TrGC	$\text{CH}_3\text{C(O)} + \text{O}_2 \rightarrow \text{OH} + \text{HCHO} + \text{CO}$	$5.1\text{E}-12 * 1. / (1. + 9.4\text{E}-18 * \text{cair})$	Atkinson et al. (2006), Beyersdorf et al. (2010)*
G42084	TrGC	$\text{C}_2\text{H}_5\text{OH} + \text{OH} \rightarrow .95 \text{C}_2\text{H}_5\text{O}_2 + .95 \text{HO}_2 + .05 \text{HOCH}_2\text{CH}_2\text{O}_2 + \text{H}_2\text{O}$	$3.0\text{E}-12 * \text{EXP}(20. / \text{temp})$	Sander et al. (2019), Atkinson et al. (2006)
G42085a	TrGCN	$\text{CH}_3\text{CN} + \text{OH} \rightarrow \text{NCCH}_2\text{O}_2 + \text{H}_2\text{O}$	$8.1\text{E}-13 * \text{EXP}(-1080. / \text{temp}) * 0.40$	Atkinson et al. (2006), Tyndall et al. (2001b)*

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G42085b	TrGCN	$\text{CH}_3\text{CN} + \text{OH} \rightarrow \text{OH} + \text{CH}_3\text{C(O)} + \text{NO}$	$8.1\text{E-}13*\text{EXP}(-1080./\text{temp})*(1.-0.40)$	Atkinson et al. (2006), Tyndall et al. (2001b)*
G42086a	TrGCN	$\text{CH}_3\text{CN} + \text{O}^{\text{(1D)}} \rightarrow \text{O}^{\text{(3P)}} + \text{CH}_3\text{CN}$	$2.54\text{E-}10*\text{EXP}(-24./\text{temp})$ $*0.0269*\text{EXP}(137./\text{temp})$	Strekowski et al. (2010)
G42086b	TrGCN	$\text{CH}_3\text{CN} + \text{O}^{\text{(1D)}} \rightarrow 2 \text{H} + \text{CO} + \text{HCN}$	$2.54\text{E-}10*\text{EXP}(-24./\text{temp})*0.16$	Strekowski et al. (2010)*
G42086c	TrGCN	$\text{CH}_3\text{CN} + \text{O}^{\text{(1D)}} \rightarrow .5 \text{CH}_3 + .5 \text{NCO} + .5 \text{NCCH}_2\text{O}_2 + .5 \text{OH}$	$2.54\text{E-}10*\text{EXP}(-24./\text{temp})*(1.-(0.16+$ $0.0269*\text{EXP}(137./\text{temp}))$	Strekowski et al. (2010)*
G42087	TrGCN	$\text{NCCH}_2\text{O}_2 + \text{NO} \rightarrow \text{HCN} + \text{CO}_2 + \text{HO}_2 + \text{NO}_2$	KR02NO	see note*
G42088	TrGCN	$\text{NCCH}_2\text{O}_2 + \text{HO}_2 \rightarrow \text{HCN} + \text{CO}_2 + \text{HO}_2$	k_R02_HO2(temp, 2)	see note*
G42089a	TrGC	$\text{CH}_2\text{CHOH} + \text{OH} \rightarrow \text{HCOOH} + \text{OH} + \text{HCHO}$	k_CH2CHOH_OH_HCOOH	Sander et al. (2019), So et al. (2014)*
G42089b	TrGC	$\text{CH}_2\text{CHOH} + \text{OH} \rightarrow \text{HOCH}_2\text{CHO} + \text{HO}_2$	k_CH2CHOH_OH_ALD	Sander et al. (2019), So et al. (2014)
G42090	TrGC	$\text{CH}_2\text{CHOH} + \text{HCOOH} \rightarrow \text{CH}_3\text{CHO} + \text{HCOOH}$	k_CH2CHOH_HCOOH	Sander et al. (2019), da Silva (2010)*
G42091	TrGC	$\text{CH}_3\text{CHO} + \text{HCOOH} \rightarrow \text{CH}_2\text{CHOH} + \text{HCOOH}$	k_ALD_HCOOH	Sander et al. (2019), da Silva (2010)*
G42092	TrGC	$\text{HOOCOOH} + \text{OH} \rightarrow 2 \text{CO}_2 + \text{HO}_2 + \text{H}_2\text{O}$	$2.0 * k_{co2h}$	see note*
G42093a	TrGC	$\text{HOCH}_2\text{CHOHOH} + \text{OH} \rightarrow \text{HOCH}_2\text{CO}_2\text{H} + \text{HO}_2 + \text{H}_2\text{O}$	$k_t*f_{toh}*f_{toh}$	see note*
G42093b	TrGC	$\text{HOCH}_2\text{CHOHOH} + \text{OH} \rightarrow \text{CHOCHOHOH} + \text{HO}_2 + \text{H}_2\text{O}$	$k_s*f_{soh}*f_{pch2oh}$	see note*
G42093c	TrGC	$\text{HOCH}_2\text{CHOHOH} + \text{OH} \rightarrow \text{HCOOH} + \text{HOCH}_2\text{O}_2 + \text{H}_2\text{O}$	$2.0 * k_{rohro}$	see note*
G42093d	TrGC	$\text{HOCH}_2\text{CHOHOH} + \text{OH} \rightarrow \text{HCHO} + \text{HCOOH} + \text{HO}_2 + \text{H}_2\text{O}$	k_{rohro}	see note*
G42094a	TrGC	$\text{CH}_3\text{CHOHOH} + \text{OH} \rightarrow \text{CH}_3\text{COOH} + \text{HO}_2 + \text{H}_2\text{O}$	$k_t*f_{toh}*f_{toh}$	see note*
G42094b	TrGC	$\text{CH}_3\text{CHOHOH} + \text{OH} \rightarrow \text{CH}_3 + \text{HCOOH} + \text{H}_2\text{O}$	$2.0 * k_{rohro}$	see note*
G42095a	TrGC	$\text{CHOHOHCOOH} + \text{OH} \rightarrow \text{HOOCOOH} + \text{HO}_2 + \text{H}_2\text{O}$	$k_t*f_{toh}*f_{toh}*f_{co2h}$	see note*
G42095b	TrGC	$\text{CHOHOHCOOH} + \text{OH} \rightarrow \text{HCOOH} + \text{CO}_2 + \text{HO}_2 + \text{H}_2\text{O}$	$2.0 * k_{rohro} + k_{co2h}$	see note*
G42096a	TrGC	$\text{CHOHOHCHOHOH} + \text{OH} \rightarrow 2 \text{HCOOH} + \text{HO}_2 + \text{H}_2\text{O}$	$4.0 * k_{rohro}$	see note*
G42096b	TrGC	$\text{CHOHOHCHOHOH} + \text{OH} \rightarrow \text{CHOHOHCOOH} + \text{HO}_2 + \text{H}_2\text{O}$	$2.0 * k_t*f_{toh}*f_{toh}*f_{pch2oh}$	see note*
G42097a	TrGC	$\text{CHOHOHCHOHOH} + \text{OH} \rightarrow \text{HCOOH} + \text{CO} + \text{HO}_2 + \text{H}_2\text{O}$	$2.0 * k_{rohro} + k_t*f_{o*f_{pch2oh}}$	see note*
G42097b	TrGC	$\text{CHOHOHCHOHOH} + \text{OH} \rightarrow \text{HCOCO}_2\text{H} + \text{HO}_2 + \text{H}_2\text{O}$	$k_t*f_{toh}*f_{toh}*f_{cho}$	see note*
G42098a	TrGC	$\text{HOOCH}_2\text{CHOHOH} + \text{OH} \rightarrow \text{HOOCH}_2\text{CO}_2\text{H} + \text{HO}_2 + \text{H}_2\text{O}$	$k_t*f_{toh}*f_{toh}*f_{pch2oh}$	see note*

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G42098b	TrGC	HOOCH ₂ CHOHOH + OH → HCOOH + HCHO + OH + H ₂ O	2.0 * k_rohro	see note*
G42098c	TrGC	HOOCH ₂ CHOHOH + OH → CHOCHOHOH + OH + H ₂ O	k_s*f_pch2oh*f_sooh	see note*
G43000a	TrGC	C ₃ H ₈ + OH → iC ₃ H ₇ O ₂ + H ₂ O	k_s	Sander et al. (2019)
G43000b	TrGC	C ₃ H ₈ + OH → C ₃ H ₇ O ₂ + H ₂ O	2.*k_p	Sander et al. (2019)
G43001a	TrGC	C ₃ H ₆ + O ₃ → HCHO + .16 CH ₃ CHOHOH + .50 OH + .50 HCOCH ₂ O ₂ + .05 CH ₂ CO + .09 CH ₃ OH + .09 CO + .2 CH ₄ + .2 CO ₂	5.5E-15*EXP(-1880./temp)*.57	Atkinson et al. (2006)*
G43001b	TrGC	C ₃ H ₆ + O ₃ → CH ₃ CHO + CH ₂ OO*	5.5E-15*EXP(-1880./temp)*.43	Atkinson et al. (2006)*
G43002	TrGC	C ₃ H ₆ + OH → HYPROPO2	k_3rd_iupac(temp,cair,8.6E-27, 3.5, 3.E-11, 1., 0.5)	Atkinson et al. (2006), Rickard (2022)
G43003	TrGCN	C ₃ H ₆ + NO ₃ → PRONO3BO2	4.6E-13*EXP(-1155./temp)	Wallington et al. (2018)
G43004	TrGC	iC ₃ H ₇ O ₂ + HO ₂ → iC ₃ H ₇ OOH	1.9E-13*EXP(1300./temp)	Atkinson (1997)*
G43005a	TrGCN	iC ₃ H ₇ O ₂ + NO → CH ₃ COCH ₃ + HO ₂ + NO ₂	2.7E-12*EXP(360./temp)*(1.-alpha_AN(3,2,0,0,0,temp,cair))	Wallington et al. (2018)
G43005b	TrGCN	iC ₃ H ₇ O ₂ + NO → iC ₃ H ₇ ONO ₂	2.7E-12*EXP(360./temp)*alpha_AN(3,2,0,0,0,temp,cair)	Wallington et al. (2018)
G43006	TrGC	iC ₃ H ₇ O ₂ → .8 CH ₃ COCH ₃ + .2 IPROPOL + .6 HO ₂	2.*(1.6E-12*EXP(-2200./temp) *k_CH3O2)**(.5)*R02	Rickard (2022), Atkinson et al. (2006)
G43007a	TrGC	iC ₃ H ₇ OOH + OH → iC ₃ H ₇ O ₂ + H ₂ O	k_ROOHRO	Sander et al. (2019)
G43007b	TrGC	iC ₃ H ₇ OOH + OH → CH ₃ COCH ₃ + H ₂ O + OH	k_t*f_tOOH	Sander et al. (2019)
G43008	TrGC	C ₃ H ₇ O ₂ + HO ₂ → C ₃ H ₇ OOH	1.9E-13*EXP(1300./temp)	Atkinson (1997)*
G43009a	TrGCN	C ₃ H ₇ O ₂ + NO → C ₂ H ₅ CHO + HO ₂ + NO ₂	2.7E-12*EXP(360./temp)*(1.-alpha_AN(3,1,0,0,0,temp,cair))	Wallington et al. (2018)
G43009b	TrGCN	C ₃ H ₇ O ₂ + NO → C ₃ H ₇ ONO ₂	2.7E-12*EXP(360./temp)*alpha_AN(3,1,0,0,0,temp,cair)	Wallington et al. (2018)
G43010	TrGC	C ₃ H ₇ O ₂ → .8 CH ₃ COCH ₃ + .2 NPROPOL + .6 HO ₂	2.*(k_CH3O2*3.E-13)**(.5)*R02	Rickard (2022), Atkinson et al. (2006)
G43011	TrGC	CH ₃ COCH ₃ + OH → CH ₃ COCH ₂ O ₂ + H ₂ O	(8.8E-12*EXP(-1320./temp) + 1.7E-14*EXP(423./temp))	Atkinson et al. (2006)*
G43012a	TrGC	CH ₃ COCH ₂ O ₂ + HO ₂ → CH ₃ COCH ₂ O ₂ H	8.6E-13*EXP(700./temp)*r_COCH2O2_-OOH	Tyndall et al. (2001a), Sander et al. (2019)
G43012b	TrGC	CH ₃ COCH ₂ O ₂ + HO ₂ → OH + CH ₃ C(O) + HCHO	8.6E-13*EXP(700./temp)*r_COCH2O2_-OH	Tyndall et al. (2001a), Sander et al. (2019)

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G43013a	TrGCN	$\text{CH}_3\text{COCH}_2\text{O}_2 + \text{NO} \rightarrow \text{CH}_3\text{C(O)} + \text{HCHO} + \text{NO}_2$	$2.9\text{E-}12*\text{EXP}(300./\text{temp})*(1.-\text{alpha}_{\text{AN}}(4,1,1,0,0,\text{temp},\text{cair}))$	Burkholder et al. (2015)
G43013b	TrGCN	$\text{CH}_3\text{COCH}_2\text{O}_2 + \text{NO} \rightarrow \text{NOA}$	$2.9\text{E-}12*\text{EXP}(300./\text{temp})*\text{alpha}_{\text{AN}}(4,1,1,0,0,\text{temp},\text{cair})$	Burkholder et al. (2015)
G43014	TrGC	$\text{CH}_3\text{COCH}_2\text{O}_2 \rightarrow .3 \text{CH}_3\text{C(O)} + .3 \text{HCHO} + .5 \text{MGLYOX} + .2 \text{CH}_3\text{COCH}_2\text{OH}$	$k_1\text{_R02pOR02}$	Orlando and Tyndall (2012)
G43015a	TrGC	$\text{CH}_3\text{COCH}_2\text{O}_2\text{H} + \text{OH} \rightarrow \text{CH}_3\text{COCH}_2\text{O}_2 + \text{H}_2\text{O}$	k_{ROOHRO}	see note*
G43015b	TrGC	$\text{CH}_3\text{COCH}_2\text{O}_2\text{H} + \text{OH} \rightarrow \text{MGLYOX} + \text{OH} + \text{H}_2\text{O}$	$k_{\text{s*f_sOH*f_CO}}$	Sander et al. (2019)
G43016	TrGC	$\text{CH}_3\text{COCH}_2\text{OH} + \text{OH} \rightarrow \text{MGLYOX} + \text{HO}_2 + \text{H}_2\text{O}$	$1.6\text{E-}12*\text{EXP}(305./\text{temp})$	Atkinson et al. (2006)
G43017	TrGC	$\text{MGLYOX} + \text{OH} \rightarrow .4 \text{CH}_3 + .6 \text{CH}_3\text{C(O)} + 1.4 \text{CO} + \text{H}_2\text{O}$	$1.9\text{E-}12*\text{EXP}(575./\text{temp})$	Baeza-Romero et al. (2007), Atkinson et al. (2006)
G43020	TrGCN	$i\text{C}_3\text{H}_7\text{ONO}_2 + \text{OH} \rightarrow \text{CH}_3\text{COCH}_3 + \text{NO}_2$	$6.2\text{E-}13*\text{EXP}(-230./\text{temp})$	Wallington et al. (2018)
G43021	TrGCN	$\text{CH}_3\text{COCH}_2\text{O}_2 + \text{NO}_3 \rightarrow \text{CH}_3\text{C(O)} + \text{HCHO} + \text{NO}_2$	KR02NO3	Rickard (2022)
G43022	TrGC	$\text{HYPROPO2} \rightarrow \text{CH}_3\text{CHO} + \text{HCHO} + \text{HO}_2$	$k_1\text{_R02sOR02}$	Rickard (2022)
G43023a	TrGC	$\text{HYPROPO2} + \text{HO}_2 \rightarrow \text{HYPROPO2H}$	$k_{\text{R02_HO2(temp,3)*(1.-r}_\text{CHOHCH202_OH)}}$	Rickard (2022)
G43023b	TrGC	$\text{HYPROPO2} + \text{HO}_2 \rightarrow \text{CH}_3\text{CHO} + \text{HCHO} + \text{HO}_2 + \text{OH}$	$k_{\text{R02_HO2(temp,3)*r}_\text{CHOHCH202_OH}}$	Rickard (2022)
G43024a	TrGCN	$\text{HYPROPO2} + \text{NO} \rightarrow \text{CH}_3\text{CHO} + \text{HCHO} + \text{HO}_2 + \text{NO}_2$	$\text{KR02NO}*(1.-\text{alpha}_{\text{AN}}(4,1,0,0,0,\text{temp},\text{cair}))$	Rickard (2022)
G43024b	TrGCN	$\text{HYPROPO2} + \text{NO} \rightarrow \text{PROPOLNO3}$	$\text{KR02NO}*\text{alpha}_{\text{AN}}(4,1,0,0,0,\text{temp},\text{cair})$	Rickard (2022)
G43025	TrGCN	$\text{HYPROPO2} + \text{NO}_3 \rightarrow \text{CH}_3\text{CHO} + \text{HCHO} + \text{HO}_2 + \text{NO}_2$	KR02NO3	Rickard (2022)
G43026a	TrGC	$\text{HYPROPO2H} + \text{OH} \rightarrow \text{HYPROPO2}$	k_{ROOHRO}	Rickard (2022)
G43026b	TrGC	$\text{HYPROPO2H} + \text{OH} \rightarrow \text{CH}_3\text{COCH}_2\text{OH} + \text{OH}$	$(k_{\text{s*f_sOH*f_pCH2OH}}+k_{\text{t*f_tOOH*f_pCH2OH}})$	Sander et al. (2019)
G43027	TrGCN	$\text{PRONO3BO2} + \text{HO}_2 \rightarrow \text{PR2O2HNO3}$	$k_{\text{R02_HO2(temp,3)}}$	Rickard (2022)
G43028	TrGCN	$\text{PRONO3BO2} + \text{NO} \rightarrow \text{NOA} + \text{HO}_2 + \text{NO}_2$	KR02NO	Rickard (2022)*
G43029	TrGCN	$\text{PRONO3BO2} + \text{NO}_3 \rightarrow \text{NOA} + \text{HO}_2 + \text{NO}_2$	KR02NO3	Rickard (2022)
G43030a	TrGCN	$\text{PR2O2HNO3} + \text{OH} \rightarrow \text{PRONO3BO2}$	k_{ROOHRO}	Rickard (2022)
G43030b	TrGCN	$\text{PR2O2HNO3} + \text{OH} \rightarrow \text{NOA} + \text{OH}$	$k_{\text{t*f_tOOH*f_CH2ON02}}$	Sander et al. (2019)
G43031	TrGCN	$\text{MGLYOX} + \text{NO}_3 \rightarrow \text{CH}_3\text{C(O)} + \text{CO} + \text{HNO}_3$	KN03AL*2.4	Rickard (2022)
G43032	TrGCN	$\text{NOA} + \text{OH} \rightarrow \text{MGLYOX} + \text{NO}_2$	$(k_{\text{s*f_CO*f_ON02}}+k_{\text{p*f_CO}})$	Sander et al. (2019)
G43033	TrGC	$\text{HOCH2COCHO} + \text{OH} \rightarrow .8609 \text{HOCH2CO} + .8609 \text{CO} + .1391 \text{HCOCOCHO} + .1391 \text{HO}_2$	$(1.9\text{E-}12*\text{EXP}(575./\text{temp})+k_{\text{s*f_sOH*f_CO}})$	Sander et al. (2019)
G43034	TrGCN	$\text{HOCH2COCHO} + \text{NO}_3 \rightarrow \text{HOCH2CO} + \text{CO} + \text{HNO}_3$	KN03AL*2.4	Sander et al. (2019)

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G43035	TrGC	$\text{CH}_3\text{COCO}_2\text{H} + \text{OH} \rightarrow \text{CH}_3\text{C(O)} + \text{H}_2\text{O} + \text{CO}_2$	4.9E-14*EXP(276./temp)	Mellouki and Mu (2003), Sander et al. (2019)
G43036	TrGC	$\text{HCOCOCH}_2\text{O}_2 \rightarrow .6 \text{ HCOCO} + .6 \text{ HCHO} + .2 \text{ HOCH}_2\text{COCHO}$	k1_R02p0R02	Sander et al. (2019)
G43037	TrGCN	$\text{HCOCOCH}_2\text{O}_2 + \text{NO} \rightarrow \text{HCOCO} + \text{HCHO} + \text{NO}_2$	KR02NO	Sander et al. (2019)*
G43038a	TrGC	$\text{HCOCOCH}_2\text{O}_2 + \text{HO}_2 \rightarrow \text{HCOCOCH}_2\text{OOH}$	k_R02_H02(temp,3)*r_COCH202_00H	Sander et al. (2019)
G43038b	TrGC	$\text{HCOCOCH}_2\text{O}_2 + \text{HO}_2 \rightarrow \text{HCOCO} + \text{HCHO} + \text{OH}$	k_R02_H02(temp,3)*r_COCH202_OH	Sander et al. (2019)
G43039	TrGCN	$\text{HCOCOCH}_2\text{O}_2 + \text{NO}_3 \rightarrow \text{HCOCO} + \text{HCHO} + \text{NO}_2$	KR02NO3	Sander et al. (2019)
G43040a	TrGC	$\text{HCOCOCH}_2\text{OOH} + \text{OH} \rightarrow \text{HOOCH}_2\text{CO}_3 + \text{CO} + \text{H}_2\text{O}$	k_t*f_CO*f_O	Sander et al. (2019)*
G43040b	TrGC	$\text{HCOCOCH}_2\text{OOH} + \text{OH} \rightarrow \text{HCOCOCHO} + \text{H}_2\text{O} + \text{OH}$	k_s*f_SO0H*f_CO	Sander et al. (2019)*
G43040c	TrGC	$\text{HCOCOCH}_2\text{OOH} + \text{OH} \rightarrow \text{HCOCOCH}_2\text{O}_2 + \text{H}_2\text{O}$	k_ROOHRO	Sander et al. (2019)
G43041	TrGCN	$\text{HCOCOCH}_2\text{OOH} + \text{NO}_3 \rightarrow \text{HOOCH}_2\text{CO}_3 + \text{CO} + \text{HNO}_3$	KN03AL*2.4	Sander et al. (2019)
G43042	TrGC	$\text{HOCH}_2\text{COCH}_2\text{O}_2 \rightarrow \text{HCHO} + \text{HOCH}_2\text{CO}$	k1_R02p0R02	Sander et al. (2019)
G43043a	TrGC	$\text{HOCH}_2\text{COCH}_2\text{O}_2 + \text{HO}_2 \rightarrow \text{HOCH}_2\text{COCH}_2\text{OOH}$	k_R02_H02(temp,3)*r_COCH202_00H	Sander et al. (2019)
G43043b	TrGC	$\text{HOCH}_2\text{COCH}_2\text{O}_2 + \text{HO}_2 \rightarrow \text{HCHO} + \text{HOCH}_2\text{CO} + \text{OH}$	k_R02_H02(temp,3)*r_COCH202_OH	Sander et al. (2019)
G43044	TrGCN	$\text{HOCH}_2\text{COCH}_2\text{O}_2 + \text{NO} \rightarrow \text{HCHO} + \text{HOCH}_2\text{CO} + \text{NO}_2$	KR02NO	Sander et al. (2019)*
G43045a	TrGC	$\text{HOCH}_2\text{COCH}_2\text{OOH} + \text{OH} \rightarrow \text{HOCH}_2\text{COCHO} + \text{OH}$	k_s*f_SO0H*f_CO	Sander et al. (2019)
G43045b	TrGC	$\text{HOCH}_2\text{COCH}_2\text{OOH} + \text{OH} \rightarrow \text{HOCH}_2\text{COCH}_2\text{O}_2$	k_ROOHRO	Sander et al. (2019)
G43045c	TrGC	$\text{HOCH}_2\text{COCH}_2\text{OOH} + \text{OH} \rightarrow \text{HCOCOCH}_2\text{OOH} + \text{HO}_2$	1.60E-12*EXP(305./temp)	Sander et al. (2019)*
G43046	TrGC	$\text{CH}_3\text{CHCO} + \text{OH} \rightarrow .72 \text{ CO} + .72 \text{ CH}_3\text{CHO} + .72 \text{ HO}_2 + .21 \text{ CH}_3\text{COCO}_2\text{H} + .07 \text{ CH}_3\text{CHO} + .07 \text{ HO}_2 + .07 \text{ CO}_2$	7.6E-11	Hatakeyama et al. (1985), Sander et al. (2019)
G43047	TrGCN	$\text{PROPOLNO}_3 + \text{OH} \rightarrow \text{CH}_3\text{COCH}_2\text{OH} + \text{NO}_2$	k_t*f_ON02*f_pCH20H+k_s*f_SOH*f_CH20NO2	Sander et al. (2019)
G43048	TrGCN	$\text{CH}_3\text{COCH}_2\text{O}_2 + \text{NO}_2 \rightarrow \text{CH}_3\text{COCH}_2\text{OONO}_2$	2.3E-12*EXP(300./temp)	Tyndall et al. (2001a)*
G43049	TrGCN	$\text{CH}_3\text{COCH}_2\text{OONO}_2 \rightarrow \text{CH}_3\text{COCH}_2\text{O}_2 + \text{NO}_2$	1.9E16*EXP(-10830./temp)	Sehested et al. (1998)*
G43050	TrGCN	$\text{CH}_3\text{COCH}_2\text{OONO}_2 + \text{OH} \rightarrow \text{MGLYOX} + \text{NO}_3 + \text{H}_2\text{O}$	9.50E-13*EXP(-650./temp)*f_CO	Sander et al. (2019)*
G43051a	TrGC	$\text{C}_3\text{H}_7\text{OOH} + \text{OH} \rightarrow \text{C}_3\text{H}_7\text{O}_2 + \text{H}_2\text{O}$	k_ROOHRO	Sander et al. (2019)
G43051b	TrGC	$\text{C}_3\text{H}_7\text{OOH} + \text{OH} \rightarrow \text{C}_2\text{H}_5\text{CHO} + \text{H}_2\text{O} + \text{OH}$	k_s*f_SO0H	Sander et al. (2019)
G43051c	TrGC	$\text{C}_3\text{H}_7\text{OOH} + \text{OH} \rightarrow \text{C}_2\text{H}_5\text{CHO} + \text{HO}_2 + \text{H}_2\text{O}$	k_s*f_pCH20H	Sander et al. (2019)*
G43052	TrGC	$\text{C}_2\text{H}_5\text{CHO} + \text{OH} \rightarrow \text{C}_2\text{H}_5\text{CO}_3 + \text{H}_2\text{O}$	4.9E-12*EXP(405./temp)	Atkinson et al. (2006)*
G43053	TrGCN	$\text{C}_2\text{H}_5\text{CHO} + \text{NO}_3 \rightarrow \text{C}_2\text{H}_5\text{CO}_3 + \text{HNO}_3$	6.3E-15	Atkinson et al. (2006)
G43054a	TrGC	$\text{C}_2\text{H}_5\text{CO}_3 \rightarrow \text{C}_2\text{H}_5\text{O}_2 + \text{CO}_2$	k1_R02RC03*0.9	Sander et al. (2019)
G43054b	TrGC	$\text{C}_2\text{H}_5\text{CO}_3 \rightarrow \text{C}_2\text{H}_5\text{CO}_2\text{H}$	k1_R02RC03*0.1	Sander et al. (2019)
G43055a	TrGC	$\text{C}_2\text{H}_5\text{CO}_3 + \text{HO}_2 \rightarrow \text{C}_2\text{H}_5\text{O}_2 + \text{CO}_2 + \text{OH}$	KAPH02*r_CO3_OH	Sander et al. (2019), Groß et al. (2014)

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G43055b	TrGC	$C_2H_5CO_3 + HO_2 \rightarrow C_2H_5CO_3H$	KAPHO2*r_CO3_00H	Sander et al. (2019), Groß et al. (2014)
G43055c	TrGC	$C_2H_5CO_3 + HO_2 \rightarrow C_2H_5CO_2H + O_3$	KAPHO2*r_CO3_03	Sander et al. (2019), Groß et al. (2014)
G43056	TrGCN	$C_2H_5CO_3 + NO \rightarrow NO_2 + C_2H_5O_2 + CO_2$	KAPNO	Rickard (2022)
G43057	TrGCN	$C_2H_5CO_3 + NO_2 \rightarrow PPN$	k_CH3CO3_NO2	Rickard (2022)
G43058	TrGCN	$PPN \rightarrow C_2H_5CO_3 + NO_2$	k_PAN_M	Rickard (2022)
G43059	TrGC	$C_2H_5CO_2H + OH \rightarrow CH_3CHO + CO_2 + H_2O$	k_CO2H+k_p+k_s*f_CO2H	Sander et al. (2019)*
G43060a	TrGC	$C_2H_5CO_3H + OH \rightarrow C_2H_5CO_3 + H_2O$	k_ROOHRO	Sander et al. (2019)
G43060b	TrGC	$C_2H_5CO_3H + OH \rightarrow CH_3CHO + CO_2 + H_2O$	k_s*f_CO2H+k_p	Sander et al. (2019)*
G43061	TrGCN	$PPN + OH \rightarrow CH_3CHO + CO_2 + NO_2 + H_2O$	k_s*f_cpan+k_p	Sander et al. (2019)*
G43062	TrGC	$CH_3COCO_3H + OH \rightarrow CH_3COCO_3 + H_2O$	k_ROOHRO	Sander et al. (2019)
G43063a	TrGC	$CH_3COCO_3 + HO_2 \rightarrow CH_3C(O) + CO_2 + OH$	KAPHO2*r_CO3_OH	Sander et al. (2019)
G43063b	TrGC	$CH_3COCO_3 + HO_2 \rightarrow CH_3COCO_3H$	KAPHO2*(r_CO3_00H+r_CO3_03)	Sander et al. (2019)
G43064	TrGCN	$CH_3COCO_3 + NO \rightarrow CH_3C(O) + CO_2 + NO_2$	KAPNO	Sander et al. (2019)
G43065	TrGCN	$CH_3COCO_3 + NO_2 \rightarrow CH_3C(O) + CO_2 + NO_3$	k_CH3CO3_NO2	Sander et al. (2019)*
G43066	TrGCN	$CH_3COCO_3 + NO_3 \rightarrow CH_3C(O)OO + CO_2 + NO_2$	KR02NO3*1.74	Sander et al. (2019)
G43067	TrGC	$CH_3COCO_3 \rightarrow CH_3C(O)OO + CO_2$	k1_RO2RC03	Sander et al. (2019)
G43068	TrGC	$HCOCOCHO + OH \rightarrow 3 CO + HO_2$	2.*k_t*f_CO*f_0	Sander et al. (2019)
G43069	TrGC	$IPOPOL + OH \rightarrow CH_3COCH_3 + HO_2 + H_2O$	2.6E-12*EXP(200./temp)	Atkinson et al. (2006)
G43070a	TrGC	$NPOPOL + OH \rightarrow C_2H_5CHO + HO_2 + H_2O$	4.6E-12*EXP(70./temp)*(k_s*f_sOH/(k_p+k_s*f_pCH20H+k_s*f_sOH))	Atkinson et al. (2006), Sander et al. (2019)*
G43070b	TrGC	$NPOPOL + OH \rightarrow HYPROPO2 + H_2O$	4.6E-12*EXP(70./temp)*((k_p+k_s*f_pCH20H)/(k_p+k_s*f_pCH20H+k_s*f_sOH))	Atkinson et al. (2006), Sander et al. (2019)*
G43071a	TrGC	$CH_2CHCH_2OH + OH \rightarrow HCOOH + OH + CH_3CHO$	k_CH2CHOH_OH_HCOOH	Sander et al. (2019), So et al. (2014)*
G43072	TrGC	$CH_2CHCH_2OH + HCOOH \rightarrow C_2H_5CHO + HCOOH$	k_CH2CHOH_HCOOH	Sander et al. (2019), da Silva (2010)*
G43073	TrGC	$C_2H_5CHO + HCOOH \rightarrow CH_2CHCH_2OH + HCOOH$	k_ALD_HCOOH	Sander et al. (2019), da Silva (2010)*
G43074	TrGC	$HCOCOCH_2OOH + OH \rightarrow HCOCO + CO + HO_2 + OH$	k_s*f_sOOH*f_CO+k_ROOHRO	Sander et al. (2019)*
G43075a	TrGC	$CH_3COCHOHOH + OH \rightarrow CH_3C(O) + HCOOH + H_2O$	2.0 * k_rohro	see note*
G43075b	TrGC	$CH_3COCHOHOH + OH \rightarrow CH_3COCO_2H + H_2O$	k_t*f_toh*f_toh*f_co	see note*
G43202	TrGTerC	$HCOCH_2CHO + OH \rightarrow HCOCH_2CO_3$	4.29E-11	Rickard (2022)

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G43203	TrGTerCN	HCOCH ₂ CHO + NO ₃ → HCOCH ₂ CO ₃ + HNO ₃	2.*KN03AL*2.4	Rickard (2022)
G43204a	TrGTerC	HCOCH ₂ CO ₃ → HCOCH ₂ O ₂ + CO ₂	k1_RO2RC03*0.9	Sander et al. (2019)
G43204b	TrGTerC	HCOCH ₂ CO ₃ → HCOCH ₂ CO ₂ H	k1_RO2RC03*0.1	Sander et al. (2019)
G43205	TrGTerCN	HCOCH ₂ CO ₃ + NO → HCOCH ₂ O ₂ + CO ₂ + NO ₂	KAPNO	Rickard (2022)
G43206	TrGTerCN	HCOCH ₂ CO ₃ + NO ₂ → C ₃ PAN2	k_CH3C03_N02	Rickard (2022)
G43207a	TrGTerC	HCOCH ₂ CO ₃ + HO ₂ → HCOCH ₂ CO ₃ H	KAPH02*r_C03_00H	Rickard (2022)
G43207b	TrGTerC	HCOCH ₂ CO ₃ + HO ₂ → HCOCH ₂ CO ₂ H + O ₃	KAPH02*r_C03_03	Rickard (2022)
G43207c	TrGTerC	HCOCH ₂ CO ₃ + HO ₂ → HCOCH ₂ O ₂ + CO ₂ + OH	KAPH02*r_C03_OH	Rickard (2022)
G43210	TrGTerCN	C ₃ PAN2 → HCOCH ₂ CO ₃ + NO ₂	k_PAN_M	Rickard (2022)
G43211	TrGTerCN	C ₃ PAN2 + OH → GLYOX + CO + NO ₂	2.10E-11	Rickard (2022)
G43212	TrGTerC	HCOCH ₂ CO ₂ H + OH → HCOCH ₂ O ₂ + CO ₂	2.14E-11	Rickard (2022)
G43213a	TrGTerC	HOC ₂ H ₄ CO ₃ → HOCH ₂ CH ₂ O ₂ + CO ₂	k1_RO2RC03*0.9	Sander et al. (2019)
G43213b	TrGTerC	HOC ₂ H ₄ CO ₃ → HOC ₂ H ₄ CO ₂ H	k1_RO2RC03*0.1	Sander et al. (2019)
G43214	TrGTerCN	HOC ₂ H ₄ CO ₃ + NO → HOCH ₂ CH ₂ O ₂ + CO ₂ + NO ₂	KAPNO	Rickard (2022)
G43215a	TrGTerC	HOC ₂ H ₄ CO ₃ + HO ₂ → HOC ₂ H ₄ CO ₃ H	KAPH02*r_C03_00H	Rickard (2022)
G43215b	TrGTerC	HOC ₂ H ₄ CO ₃ + HO ₂ → HOCH ₂ CH ₂ O ₂ + CO ₂ + OH	KAPH02*r_C03_OH	Rickard (2022)
G43215c	TrGTerC	HOC ₂ H ₄ CO ₃ + HO ₂ → HOC ₂ H ₄ CO ₂ H + O ₃	KAPH02*r_C03_03	Rickard (2022)
G43218	TrGTerCN	HOC ₂ H ₄ CO ₃ + NO ₂ → C ₃ PAN1	k_CH3C03_N02	Rickard (2022)
G43219	TrGTerC	HOC ₂ H ₄ CO ₂ H + OH → HOCH ₂ CH ₂ O ₂ + CO ₂	1.39E-11	Rickard (2022)
G43220	TrGTerC	HOC ₂ H ₄ CO ₃ H + OH → HOC ₂ H ₄ CO ₃	1.73E-11	Rickard (2022)
G43221	TrGTerCN	C ₃ PAN1 → HOC ₂ H ₄ CO ₃ + NO ₂	k_PAN_M	Rickard (2022)
G43222	TrGTerCN	C ₃ PAN1 + OH → HOCH ₂ CHO + CO + NO ₂	4.51E-12	Rickard (2022)
G43223	TrGTerC	HCOCH ₂ CO ₃ H + OH → HCOCH ₂ O ₂ + CO ₂ + H ₂ O	2.49E-11	Rickard (2022)*
G43415	TrGAroC	C ₃ DIALOOH + OH → HCOCOCHO + OH	1.44E-10	Rickard (2022)
G43418a	TrGAroC	C ₃ DIALO ₂ + HO ₂ → C ₃ DIALOOH	k_RO2_H02(temp,3)*(r_C03_00H+r_C03_03)	Rickard (2022)
G43418b	TrGAroC	C ₃ DIALO ₂ + HO ₂ → GLYOX + CO + HO ₂ + OH	k_RO2_H02(temp,3)*r_C03_OH	Rickard (2022)
G43419	TrGAroCN	C ₃ DIALO ₂ + NO → GLYOX + CO + HO ₂ + NO ₂	KR02NO	Rickard (2022)*
G43420	TrGAroCN	C ₃ DIALO ₂ + NO ₃ → GLYOX + CO + HO ₂ + NO ₂	KR02N03	Rickard (2022)*
G43421	TrGAroC	C ₃ DIALO ₂ → GLYOX + CO + HO ₂	k1_RO2sOR02	Rickard (2022)*
G43422a	TrGAroC	HCOCOHCO ₃ + HO ₂ → GLYOX + CO ₂ + HO ₂ + OH	KAPH02*r_C03_OH	Rickard (2022)
G43422b	TrGAroC	HCOCOHCO ₃ + HO ₂ → HCOCOHCO ₃ H	KAPH02*(r_C03_00H+r_C03_03)	Rickard (2022)
G43424	TrGAroCN	HCOCOHCO ₃ + NO → GLYOX + CO ₂ + HO ₂ + NO ₂	KAPNO	Rickard (2022)
G43425	TrGAroCN	HCOCOHCO ₃ + NO ₂ → HCOCOHPAN	k_CH3C03_N02	Rickard (2022)
G43426	TrGAroCN	HCOCOHCO ₃ + NO ₃ → GLYOX + CO ₂ + HO ₂ + NO ₂	KR02N03*1.74	Rickard (2022)

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G43427	TrGAroC	$\text{HCOCOHC}3 \rightarrow \text{GLYOX} + \text{CO}_2 + \text{HO}_2$	k1_R02RC03	Rickard (2022)
G43428	TrGAroC	$\text{METACETHO} + \text{OH} \rightarrow \text{CH}_3\text{C(O)} + \text{CO}_2$	9.82E-11	Rickard (2022)
G43442	TrGAroCN	$\text{HCOCOHPAN} + \text{OH} \rightarrow \text{GLYOX} + \text{CO} + \text{NO}_2$	6.97E-11	Rickard (2022)
G43443	TrGAroCN	$\text{HCOCOHPAN} \rightarrow \text{HCOCOHC}3 + \text{NO}_2$	k_PAN_M	Rickard (2022)
G43444	TrGAroC	$\text{C32OH13CO} + \text{OH} \rightarrow \text{HCOCOHC}3$	1.36E-10	Rickard (2022)
G43446	TrGAroC	$\text{HCOCOHC}3\text{H} + \text{OH} \rightarrow \text{HCOCOHC}3$	7.33E-11	Rickard (2022)
G44000	TrGC	$\text{C}_4\text{H}_{10} + \text{OH} \rightarrow \text{LC}_4\text{H}_9\text{O}_2 + \text{H}_2\text{O}$	$2.03\text{E}-17 * \text{temp} * \text{temp} * \text{EXP}(78./\text{temp})$	Atkinson et al. (2006)*
G44001a	TrGC	$\text{LC}_4\text{H}_9\text{O}_2 \rightarrow \text{C}_3\text{H}_7\text{CHO} + \text{HO}_2$	$(k1_{_R02pR02}*0.1273+k1_{_R02sR02}*0.8727)*0.1273$	Rickard (2022), Sander et al. (2019)
G44001b	TrGC	$\text{LC}_4\text{H}_9\text{O}_2 \rightarrow .636 \text{ MEK} + .636 \text{ HO}_2 + .364 \text{ CH}_3\text{CHO} + .364 \text{ C}_2\text{H}_5\text{O}_2$	$(k1_{_R02pR02}*0.1273+k1_{_R02sR02}*0.8727)*0.8727$	Rickard (2022), Sander et al. (2019)*
G44002	TrGC	$\text{LC}_4\text{H}_9\text{O}_2 + \text{HO}_2 \rightarrow \text{LC}_4\text{H}_9\text{OOH}$	k_R02_H02(temp, 4)	Rickard (2022)
G44003a	TrGCN	$\text{LC}_4\text{H}_9\text{O}_2 + \text{NO} \rightarrow \text{NO}_2 + \text{C}_3\text{H}_7\text{CHO} + \text{HO}_2$	$\text{KR02NO}*(1.-(0.1273*\text{alpha_AN}(4,1,0,0,\text{temp},\text{cair})+0.8727*\text{alpha_AN}(4,2,0,0,0,\text{temp},\text{cair}))) * 0.1273$	Rickard (2022), Sander et al. (2019)
G44003b	TrGCN	$\text{LC}_4\text{H}_9\text{O}_2 + \text{NO} \rightarrow \text{NO}_2 + .636 \text{ MEK} + .636 \text{ HO}_2 + .364 \text{ CH}_3\text{CHO} + .364 \text{ C}_2\text{H}_5\text{O}_2$	$\text{KR02NO}*(1.-(0.1273*\text{alpha_AN}(4,1,0,0,\text{temp},\text{cair})+0.8727*\text{alpha_AN}(4,2,0,0,0,\text{temp},\text{cair}))) * 0.8727$	Rickard (2022), Sander et al. (2019)
G44003c	TrGCN	$\text{LC}_4\text{H}_9\text{O}_2 + \text{NO} \rightarrow \text{LC}_4\text{H}_9\text{NO}_3$	$\text{KR02NO}*(0.1273*\text{alpha_AN}(4,1,0,0,0,\text{temp},\text{cair})+0.8727*\text{alpha_AN}(4,2,0,0,0,\text{temp},\text{cair}))$	Rickard (2022)*
G44004a	TrGCN	$\text{LC}_4\text{H}_9\text{O}_2 + \text{NO}_3 \rightarrow \text{NO}_2 + \text{C}_3\text{H}_7\text{CHO} + \text{HO}_2$	KR02NO3*0.1273	Rickard (2022), Sander et al. (2019)
G44004b	TrGCN	$\text{LC}_4\text{H}_9\text{O}_2 + \text{NO}_3 \rightarrow \text{NO}_2 + .636 \text{ MEK} + .636 \text{ HO}_2 + .364 \text{ CH}_3\text{CHO} + .364 \text{ C}_2\text{H}_5\text{O}_2$	KR02NO3*0.8727	Rickard (2022), Sander et al. (2019)
G44005a	TrGC	$\text{LC}_4\text{H}_9\text{OOH} + \text{OH} \rightarrow \text{LC}_4\text{H}_9\text{O}_2 + \text{H}_2\text{O}$	k_ROOHRO	Sander et al. (2019)
G44005b	TrGC	$\text{LC}_4\text{H}_9\text{OOH} + \text{OH} \rightarrow \text{C}_3\text{H}_7\text{CHO} + \text{H}_2\text{O} + \text{OH}$	$k_s*f_{t00H}*f_{alk}*(k_p/(k_p+k_s))$	Sander et al. (2019)
G44005c	TrGC	$\text{LC}_4\text{H}_9\text{OOH} + \text{OH} \rightarrow \text{MEK} + \text{H}_2\text{O} + \text{OH}$	$k_t*f_{t00H}*f_{alk}*(k_s/(k_p+k_s))$	Sander et al. (2019)
G44006a	TrGC	$\text{iC}_4\text{H}_{10} + \text{OH} \rightarrow \text{TC}_4\text{H}_9\text{O}_2 + \text{H}_2\text{O}$	$1.17\text{E}-17 * \text{temp} * \text{temp} * \text{EXP}(213./\text{temp}) * k_t / (3.*k_p+k_t)$	Atkinson (2003)
G44006b	TrGC	$\text{iC}_4\text{H}_{10} + \text{OH} \rightarrow \text{IC}_4\text{H}_9\text{O}_2 + \text{H}_2\text{O}$	$1.17\text{E}-17 * \text{temp} * \text{temp} * \text{EXP}(213./\text{temp}) * 3.*k_p / (3.*k_p+k_t)$	Atkinson (2003)
G44007	TrGC	$\text{TC}_4\text{H}_9\text{O}_2 \rightarrow \text{CH}_3\text{COCH}_3 + \text{CH}_3$	k1_R02tR02	Rickard (2022), Sander et al. (2019)
G44008	TrGC	$\text{TC}_4\text{H}_9\text{O}_2 + \text{HO}_2 \rightarrow \text{TC}_4\text{H}_9\text{OOH}$	k_R02_H02(temp, 4)	Rickard (2022)

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G44009a	TrGCN	TC ₄ H ₉ O ₂ + NO → NO ₂ + CH ₃ COCH ₃ + CH ₃	KR02NO*(1.-alpha_AN(4,3,0,0,0, temp,cair))	Rickard (2022), Sander et al. (2019)
G44009b	TrGCN	TC ₄ H ₉ O ₂ + NO → TC4H9NO3	KR02NO*alpha_AN(4,3,0,0,0,temp, cair)	Rickard (2022)
G44010a	TrGC	TC ₄ H ₉ OOH + OH → TC ₄ H ₉ O ₂ + H ₂ O	k_ROOHRO	Sander et al. (2019)
G44010b	TrGC	TC ₄ H ₉ OOH + OH → CH ₃ COCH ₃ + HCHO + OH + H ₂ O	3.*k_p*f_tCH2OH	Sander et al. (2019)*
G44011	TrGCN	TC4H9NO3 + OH → CH ₃ COCH ₃ + HCHO + NO ₂ + H ₂ O	3.*k_p*f_CH2ON02	Sander et al. (2019)*
G44012	TrGC	IC ₄ H ₉ O ₂ → IPRCHO	k1_R02sR02	Rickard (2022), Sander et al. (2019)
G44013	TrGC	IC ₄ H ₉ O ₂ + HO ₂ → IC ₄ H ₉ OOH	k_R02_HO2(temp,4)	Rickard (2022)
G44014a	TrGCN	IC ₄ H ₉ O ₂ + NO → NO ₂ + IPRCHO	KR02NO*(1.-alpha_AN(4,2,0,0,0, temp,cair))	Rickard (2022), Sander et al. (2019)
G44014b	TrGCN	IC ₄ H ₉ O ₂ + NO → IC4H9NO3	KR02NO*alpha_AN(4,2,0,0,0,temp, cair)	Rickard (2022)
G44015a	TrGC	IC ₄ H ₉ OOH + OH → IC ₄ H ₉ O ₂ + H ₂ O	k_ROOHRO	Sander et al. (2019)
G44015b	TrGC	IC ₄ H ₉ OOH + OH → IPRCHO + OH + H ₂ O	k_s*f_SO0H+2.*k_s+k_t*f_pCH2OH	Sander et al. (2019)*
G44016	TrGCN	IC4H9NO3 + OH → IPRCHO + NO ₂ + H ₂ O	k_s*f_ON02+2.*k_p+k_t*f_CH2ON02	Sander et al. (2019)*
G44017	TrGC	MVK + O ₃ → .87 MGLYOX + .5481 CO + .1392 HO ₂ + .1392 OH + .3219 CH ₂ OO + .13 HCHO + .04680 OH + .04680 CO + .07280 CH ₃ C(O) + .026 CH ₃ CHO + .026 CO ₂ + .026 HCHO + .026 HO ₂ + .02402 MGLYOX + .02402 H ₂ O ₂ + .00718 CH ₃ COCO ₂ H	8.5E-16*EXP(-1520./temp)	Sander et al. (2019)
G44018	TrGC	MVK + OH → LHMVKABO2	2.6E-12*EXP(610./temp)	Sander et al. (2019), Atkinson et al. (2006)*
G44019	TrGC	MEK + OH → LMEKO2 + H ₂ O	1.5E-12*EXP(-90./temp)	Atkinson et al. (2006), Sander et al. (2019)*
G44020	TrGC	LMEKO2 + HO ₂ → LMEKOOH	k_R02_HO2(temp,4)	Sander et al. (2019)
G44021a	TrGCN	LMEKO2 + NO → .62 CH ₃ CHO + .62 CH ₃ C(O) + .38 HCHO + .38 CO ₂ + .38 HOCH ₂ CH ₂ O ₂ + NO ₂	KR02NO*(1.-(.62*alpha_AN(4,2,1, 0,0,temp,cair)+.38*alpha_AN(4,1, 0,1,0,temp,cair)))	Sander et al. (2019)*
G44021b	TrGCN	LMEKO2 + NO → LMEKNO3	KR02NO*(.62*alpha_AN(4,2,1,0,0, temp,cair)+.38*alpha_AN(4,1,0,1, 0,temp,cair))	Sander et al. (2019)
G44022a	TrGC	LMEKOOH + OH → LMEKO2 + H ₂ O	k_ROOHRO	Sander et al. (2019)

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G44022b	TrGC	LMEKOOH + OH → .62 BIACET + .38 HCHO + .38 CO ₂ + .38 HOCH ₂ CH ₂ O ₂ + H ₂ O + OH	(.62*k_t*f_t0OH*f_CO+.38*k_s*f_sOH)	Sander et al. (2019)
G44023a	TrGCN	LC4H9NO ₃ + OH → MEK + NO ₂ + H ₂ O	(k_t*f_ONO2*f_alk+k_p*f_alk+k_s*f_CH2ONO2+k_p)*(k_s/(k_p+k_s))	Sander et al. (2019)*
G44023b	TrGCN	LC4H9NO ₃ + OH → C ₃ H ₇ CHO + NO ₂ + H ₂ O	(k_p+k_s*(1+f_CH2ONO2+f_ONO2)*f_alk)*(k_p/(k_p+k_s))	Sander et al. (2019)*
G44024	TrGCN	MPAN + OH → CH ₃ COCH ₂ OH + CO + NO ₂	3.2E-11	Orlando et al. (2002)
G44025	TrGCN	MPAN → MACO ₃ + NO ₂	k_PAN_M	see note*
G44026	TrGC	LMEKO ₂ → .538 HCHO + .538 CO ₂ + .459 HOCH ₂ CH ₂ O ₂ + .079 C ₂ H ₅ O ₂ + .462 CH ₃ C(O) + .462 CH ₃ CHO	(.62*k1_R02sOR02+.38*k1_R02pOR02)	Rickard (2022)*
G44027	TrGC	MACR + OH → .45 MACO ₃ + .55 MACRO ₂	8.E-12*EXP(380./temp)	Orlando et al. (1999b), Sander et al. (2019)
G44028	TrGC	MACR + O ₃ → .5481 CO + .1392 HO ₂ + .1392 OH + .3219 CH ₂ OO + .87 MGLYOX + .13 HCHO + .13 OH + .065 HCOCOCH ₂ O ₂ + .065 CO + .065 CH ₃ C(O)	1.36E-15*EXP(-2112./temp)	Sander et al. (2019)
G44029	TrGCN	MACR + NO ₃ → MACO ₃ + HNO ₃	KN03AL*2.0	Rickard (2022)
G44030a	TrGC	MACO ₃ → CH ₃ C(O) + HCHO + CO ₂	k1_R02RC03*0.9	Sander et al. (2019)
G44030b	TrGC	MACO ₃ → MACO ₂ H	k1_R02RC03*0.1	Sander et al. (2019)
G44031a	TrGC	MACO ₃ + HO ₂ → MACO ₂ + OH	KAPH02*r_CO3_OH	Sander et al. (2019)
G44031b	TrGC	MACO ₃ + HO ₂ → MACO ₃ H	KAPH02*r_CO3_OOH	Sander et al. (2019)
G44031c	TrGC	MACO ₃ + HO ₂ → MACO ₂ H + O ₃	KAPH02*r_CO3_03	Sander et al. (2019)
G44032	TrGCN	MACO ₃ + NO → MACO ₂ + NO ₂	8.70E-12*EXP(290./temp)	Sander et al. (2019)
G44033	TrGCN	MACO ₃ + NO ₂ → MPAN	k_CH3C03_N02	Rickard (2022)
G44034	TrGCN	MACO ₃ + NO ₃ → MACO ₂ + NO ₂	KR02N03*1.74	Sander et al. (2019)
G44035	TrGC	MACRO ₂ → .7 CH ₃ COCH ₂ OH + .7 HCHO + .7 HO ₂ + .3 MACROH	k1_R02tOR02	Rickard (2022)*
G44036a	TrGC	MACRO ₂ + HO ₂ → MACRO + OH	k_R02_H02(temp,4)*r_COCH202_OH	Sander et al. (2019)
G44036b	TrGC	MACRO ₂ + HO ₂ → MACROOH	k_R02_H02(temp,4)*r_COCH202_OOH	Sander et al. (2019)
G44037a	TrGCN	MACRO ₂ + NO → MACRO + NO ₂	KR02NO*(1.-alpha_AN(6,3,1,0,0, temp, cair))	Sander et al. (2019)
G44037b	TrGCN	MACRO ₂ + NO → MACRNO ₃	KR02NO*alpha_AN(6,3,1,0,0,temp, cair)	Sander et al. (2019)
G44038	TrGCN	MACRO ₂ + NO ₃ → MACRO + NO ₂	KR02N03	Sander et al. (2019)
G44039a	TrGC	MACROOH + OH → MACRO ₂	k_ROOHRO	Sander et al. (2019)

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G44039b	TrGC	MACROOH + OH → CO + CH ₃ COCH ₂ OH + OH	k_t*f_0*f_tCH2OH*f_alk	Sander et al. (2019)
G44039c	TrGC	MACROOH + OH → CO + MGLYOX + HO ₂	(k_s*f_sOH*f_pCH2OH + k_ROHRO)	Sander et al. (2019)
G44040	TrGC	MACROH + OH → CH ₃ COCH ₂ OH + CO + HO ₂	k_t*f_0*f_tCH2OH*f_alk	Sander et al. (2019)
G44041	TrGC	MACRO → .885 CH ₃ COCH ₂ OH + .885 CO + .115 MGLYOX + .115 HCHO + HO ₂	KDEC	Sander et al. (2019)
G44042	TrGC	MACO2H + OH → CH ₃ COCH ₂ OH + HO ₂ + CO ₂	((k_adt+k_adp)*a_CO2H+k_CO2H)	Sander et al. (2019)
G44043a	TrGC	MACO3H + OH → CH ₃ COCH ₂ OH + CO ₂ + OH	(k_adt+k_adp)*a_CO2H	Sander et al. (2019)
G44043b	TrGC	MACO3H + OH → MACO3	k_ROOHRO	Sander et al. (2019)
G44044	TrGC	LHMVKABO2 → .024 CO2H3CHO + .072 MGLYOX + .072 HO ₂ + .072 HCHO + .5280 CH ₃ C(O) + .5280 HOCH ₂ CHO + .176 BIACETOH + .2 HO12CO3C4	(.12*k1_R02p0R02+.88*k1_R02s0R02)	Sander et al. (2019)
G44045a	TrGC	LHMVKABO2 + HO ₂ → OH + HOCH ₂ CHO + CH ₃ C(O)	k_R02_H02(temp,4)*.88*r_COCH202_OH	Sander et al. (2019)
G44045b	TrGC	LHMVKABO2 + HO ₂ → LHMVKABOOH	k_R02_H02(temp,4)*(.12+.88*r_COCH202_OOH)	Sander et al. (2019)
G44046a	TrGCN	LHMVKABO2 + NO → .12 MGLYOX + .12 HO ₂ + .88 HOCH ₂ CHO + .88 CH ₃ C(O) + .12 HCHO + NO ₂	KR02NO*(1.-(.12*alpha_AN(6,1,0,1,0,temp,cair)+.88*alpha_AN(6,2,1,0,0,temp,cair)))	Sander et al. (2019)
G44046b	TrGCN	LHMVKABO2 + NO → MVKNO3	KR02NO*(.12*alpha_AN(6,1,0,1,0,temp,cair)+.88*alpha_AN(6,2,1,0,0,temp,cair))	Sander et al. (2019)*
G44047	TrGCN	LHMVKABO2 + NO ₃ → .12 MGLYOX + .12 HO ₂ + .88 HOCH ₂ CHO + .88 CH ₃ C(O) + .12 HCHO + .12 HO ₂ + NO ₂	KR02NO3	Sander et al. (2019)
G44048a	TrGC	LHMVKABOOH + OH → LHMVKABO2	k_ROOHRO	Sander et al. (2019)
G44048b	TrGC	LHMVKABOOH + OH → .12 CO2H3CHO + .88 BIACETOH + OH	(.12*k_s*f_sOOH*f_pCH2OH+.88*k_t*f_tOOH*f_pCH2OH*f_CO)	Sander et al. (2019)
G44049a	TrGC	CO2H3CHO + OH → CO2H3CO3	k_t*f_0*f_alk	Sander et al. (2019)
G44049b	TrGC	CO2H3CHO + OH → CH ₃ COCOCHO + HO ₂ + H ₂ O	k_t*f_CO*f_tOH*f_CHO	Sander et al. (2019)
G44050	TrGCN	CO2H3CHO + NO ₃ → CO2H3CO3 + HNO ₃	KNO3AL*4.0	Rickard (2022)
G44051	TrGC	CO2H3CO3 → MGLYOX + HO ₂ + CO ₂	k1_R02RC03	Sander et al. (2019)
G44052a	TrGC	CO2H3CO3 + HO ₂ → OH + MGLYOX + HO ₂ + CO ₂	KAPH02*r_CO3_OH	Sander et al. (2019)
G44052b	TrGC	CO2H3CO3 + HO ₂ → CO2H3CO2H + O ₃	KAPH02*r_CO3_03	Sander et al. (2019)
G44052c	TrGC	CO2H3CO3 + HO ₂ → CO2H3CO3H	KAPH02*r_CO3_OOH	Sander et al. (2019)
G44053	TrGCN	CO2H3CO3 + NO → MGLYOX + HO ₂ + NO ₂ + CO ₂	KAPNO	Sander et al. (2019)

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G44054	TrGCN	$\text{CO}_2\text{H}_3\text{CO}_3 + \text{NO}_3 \rightarrow \text{MGLYOX} + \text{HO}_2 + \text{NO}_2 + \text{CO}_2$	KR02N03*1.74	Sander et al. (2019)
G44055a	TrGC	$\text{CO}_2\text{H}_3\text{CO}_3\text{H} + \text{OH} \rightarrow \text{CO}_2\text{H}_3\text{CO}_3$	k_{ROOHRO}	Sander et al. (2019)
G44055b	TrGC	$\text{CO}_2\text{H}_3\text{CO}_3\text{H} + \text{OH} \rightarrow \text{CH}_3\text{C(O)} + \text{CO} + \text{CO}_2 + \text{OH}$	$(k_{\text{t*f_CO2H*f_CO*f_tOH}})$	Sander et al. (2019)
G44056	TrGC	$\text{CO}_2\text{H}_3\text{CO}_2\text{H} + \text{OH} \rightarrow \text{CH}_3\text{COCOCO}_2\text{H} + \text{HO}_2$	$k_{\text{t*f_CO2H*f_CO*f_tOH+k_CO2H}}$	Sander et al. (2019)
G44057a	TrGC	$\text{HO}_12\text{CO}_3\text{C}_4 + \text{OH} \rightarrow \text{BIACETOH} + \text{HO}_2$	$k_{\text{t*f_tOH*f_alk*f_CO}}$	Sander et al. (2019)
G44057b	TrGC	$\text{HO}_12\text{CO}_3\text{C}_4 + \text{OH} \rightarrow \text{CO}_2\text{H}_3\text{CHO} + \text{HO}_2$	$k_{\text{s*f_sOH*f_alk}}$	Sander et al. (2019)
G44058	TrGC	$\text{MACO}_2 \rightarrow .65 \text{ CH}_3 + .65 \text{ CO} + .65 \text{ HCHO} + .35 \text{ OH} + .35 \text{ CH}_3\text{COCH}_2\text{O}_2 + \text{CO}_2$	KDEC	Sander et al. (2019)
G44059	TrGC	$\text{LHMVKABO}_2 \rightarrow .88 \text{ MGLYOX} + .88 \text{ HCHO} + .12 \text{ HOOCH}_2\text{CHO} + .12 \text{ CH}_3\text{C(O)} + \text{OH}$	k_{hsd}	Sander et al. (2019)
G44060	TrGC	$\text{MACRO}_2 \rightarrow \text{MGLYOX} + \text{HCHO} + \text{OH}$	k_{hsb}	Sander et al. (2019)
G44061a	TrGCN	$\text{MVKN}_3 + \text{OH} \rightarrow \text{MGLYOX} + \text{CO}_2 + \text{HO}_2 + \text{NO}_2 + \text{H}_2\text{O}$	$k_{\text{s*f_sOH*f_CH2ON02+k_ROHRO}}$	Sander et al. (2019)*
G44061b	TrGCN	$\text{MVKN}_3 + \text{OH} \rightarrow \text{BIACETOH} + \text{NO}_2 + \text{H}_2\text{O}$	$k_{\text{t*f_ON02*f_CO*f_pCH2OH}}$	Sander et al. (2019)*
G44062a	TrGCN	$\text{MACRNO}_3 + \text{OH} \rightarrow \text{CH}_3\text{COCH}_2\text{OH} + \text{CO}_2 + \text{NO}_2 + \text{H}_2\text{O}$	$k_{\text{t*f_O*f_CH2ON02}}$	Sander et al. (2019)*
G44062b	TrGCN	$\text{MACRNO}_3 + \text{OH} \rightarrow \text{MGLYOX} + \text{CO} + \text{NO}_2 + \text{H}_2\text{O}$	$k_{\text{ROHRO+k_s*f_sOH*f_CH2ON02}}$	Sander et al. (2019)*
G44063	TrGC	$\text{MACRO}_2 \rightarrow \text{CH}_3\text{COCH}_2\text{OH} + \text{OH} + \text{CO}$	k_{14hsal}	Sander et al. (2019)
G44064	TrGC	$\text{EZCH}_3\text{CO}_2\text{CHCHO} \rightarrow .9 \text{ CH}_3\text{COCHCO} + .1 \text{ CH}_3\text{C(O)} + .01 \text{ GLYOX} + .18 \text{ CO} + .09 \text{ HO}_2 + \text{OH}$	$k_{\text{15hs24vynal}}$	Sander et al. (2019)
G44065	TrGC	$\text{EZCH}_3\text{CO}_2\text{CHCHO} + \text{HO}_2 \rightarrow \text{CH}_3\text{COOHCHCHO}$	$k_{\text{R02_HO2(temp,4)}}$	Sander et al. (2019)
G44066	TrGCN	$\text{EZCH}_3\text{CO}_2\text{CHCHO} + \text{NO} \rightarrow \text{CH}_3\text{COCHO}_2\text{CHO} + \text{NO}_2$	KR02NO	Sander et al. (2019)*
G44067	TrGCN	$\text{EZCH}_3\text{CO}_2\text{CHCHO} + \text{NO}_3 \rightarrow \text{CH}_3\text{COCHO}_2\text{CHO} + \text{NO}_2$	KR02N03	Sander et al. (2019)
G44068	TrGC	$\text{EZCH}_3\text{CO}_2\text{CHCHO} \rightarrow \text{CH}_3\text{COCHO}_2\text{CHO}$	$k_{\text{1_R02sOR02}}$	Sander et al. (2019)
G44069	TrGC	$\text{EZCHOCCH}_3\text{CHO}_2 \rightarrow \text{HCOCCH}_3\text{CO} + \text{OH}$	$k_{\text{15hs24vynal}}$	Sander et al. (2019)
G44070	TrGCN	$\text{EZCHOCCH}_3\text{CHO}_2 + \text{NO} \rightarrow \text{HCOCO}_2\text{CH}_3\text{CHO} + \text{NO}_2$	KR02NO	Sander et al. (2019)*
G44071	TrGC	$\text{EZCHOCCH}_3\text{CHO}_2 + \text{HO}_2 \rightarrow \text{HCOCCH}_3\text{CHOOH}$	$k_{\text{R02_HO2(temp,4)}}$	Sander et al. (2019)
G44072	TrGCN	$\text{EZCHOCCH}_3\text{CHO}_2 + \text{NO}_3 \rightarrow \text{HCOCO}_2\text{CH}_3\text{CHO} + \text{NO}_2$	KR02N03	Sander et al. (2019)
G44073	TrGC	$\text{EZCHOCCH}_3\text{CHO}_2 \rightarrow \text{HCOCO}_2\text{CH}_3\text{CHO}$	$k_{\text{1_R02pOR02}}$	Sander et al. (2019)
G44074	TrGC	$\text{CH}_3\text{COOHCHCHO} \rightarrow \text{CH}_3\text{COCHO}_2\text{CHO} + \text{OH}$	k_{hydec}	Sander et al. (2019)
G44075	TrGC	$\text{HCOCCH}_3\text{CHOOH} \rightarrow \text{HCOCO}_2\text{CH}_3\text{CHO} + \text{OH}$	k_{hydec}	Sander et al. (2019)
G44076	TrGCN	$\text{CH}_3\text{COCHO}_2\text{CHO} + \text{NO} \rightarrow \text{CH}_3\text{C(O)} + \text{GLYOX} + \text{NO}_2$	KR02NO	Sander et al. (2019)*
G44077	TrGCN	$\text{CH}_3\text{COCHO}_2\text{CHO} + \text{NO}_3 \rightarrow \text{CH}_3\text{C(O)} + \text{GLYOX} + \text{NO}_2$	KR02N03	Sander et al. (2019)
G44078	TrGC	$\text{CH}_3\text{COCHO}_2\text{CHO} + \text{HO}_2 \rightarrow \text{CH}_3\text{C(O)} + \text{GLYOX} + \text{OH}$	$k_{\text{R02_HO2(temp,4)}}$	Sander et al. (2019)*
G44079	TrGC	$\text{CH}_3\text{COCHO}_2\text{CHO} \rightarrow \text{CH}_3\text{C(O)} + \text{GLYOX}$	$k_{\text{1_R02sOR02}}$	Sander et al. (2019)

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G44080	TrGC	$\text{HCOCO}_2\text{CH}_3\text{CHO} \rightarrow \text{MGLYOX} + \text{CO} + \text{HO}_2$	$k_{1_R02t0R02}$	Sander et al. (2019)
G44081	TrGCN	$\text{HCOCO}_2\text{CH}_3\text{CHO} + \text{NO} \rightarrow \text{MGLYOX} + \text{CO} + \text{HO}_2 + \text{NO}_2$	k_{R02NO}	Sander et al. (2019)*
G44082	TrGC	$\text{HCOCO}_2\text{CH}_3\text{CHO} + \text{HO}_2 \rightarrow \text{MGLYOX} + \text{CO} + \text{HO}_2 + \text{OH}$	$k_{R02_HO2(temp,4)}$	Sander et al. (2019)*
G44083	TrGCN	$\text{HCOCO}_2\text{CH}_3\text{CHO} + \text{NO}_3 \rightarrow \text{MGLYOX} + \text{CO} + \text{HO}_2 + \text{NO}_2$	k_{R02NO3}	Sander et al. (2019)
G44084	TrGC	$\text{HCOCCH}_3\text{CO} + \text{OH} \rightarrow \text{CO} + \text{MGLYOX} + \text{HO}_2$	$1E-10*a_{\text{CHO}}$	Hatakeyama et al. (1985), Sander et al. (2019)
G44085	TrGC	$\text{CH}_3\text{COCHCO} + \text{OH} \rightarrow \text{CO} + \text{MGLYOX} + \text{HO}_2$	$7.6E-11*a_{\text{COCH3}}$	Hatakeyama et al. (1985), Sander et al. (2019)*
G44086	TrGCN	$\text{LMEKNO}_3 + \text{OH} \rightarrow .62 \text{ MGLYOX} + .62 \text{ HCHO} + .62 \text{ HO}_2 + .62 \text{ NO}_2 + .38 \text{ CH}_3\text{C(O)} + .38 \text{ NO}_3\text{CH}_2\text{CHO}$	$.62*(k_p*(f_{\text{CO}}+f_{\text{CH2ON02}})) + .38*(k_s*f_{\text{CH2ON02}}*f_{\text{CO}})$	Sander et al. (2019)*
G44087	TrGC	$\text{MEPROPENE} + \text{OH} \rightarrow \text{IBUTOLBO2}$	$9.4E-12*\text{EXP}(505./\text{temp})$	Atkinson et al. (2006)
G44088a	TrGC	$\text{MEPROPENE} + \text{O}_3 \rightarrow \text{CH}_3\text{COCH}_3 + \text{CH}_2\text{OO}^*$	$2.7E-15*\text{EXP}(-1630./\text{temp})*0.33$	Atkinson et al. (2006), Sander et al. (2019)
G44088b	TrGC	$\text{MEPROPENE} + \text{O}_3 \rightarrow \text{CH}_3\text{COCH}_2\text{O}_2 + \text{OH} + \text{HCHO}$	$2.7E-15*\text{EXP}(-1630./\text{temp})*0.67$	Atkinson et al. (2006), Sander et al. (2019)
G44089	TrGCN	$\text{MEPROPENE} + \text{NO}_3 \rightarrow \text{CH}_3\text{COCH}_3 + \text{HCHO} + \text{NO}_2$	$3.4E-13$	Atkinson et al. (2006), Sander et al. (2019)*
G44090	TrGC	$\text{IBUTOLBO2} \rightarrow \text{CH}_3\text{COCH}_3 + \text{HCHO} + \text{HO}_2$	$k_{1_R02t0R02}$	Sander et al. (2019)
G44091a	TrGC	$\text{IBUTOLBO2} + \text{HO}_2 \rightarrow \text{IBUTOLBOOH}$	$k_{R02_HO2(temp,4)*r_{\text{COCH2O2_OOH}}}$	Sander et al. (2019)
G44091b	TrGC	$\text{IBUTOLBO2} + \text{HO}_2 \rightarrow \text{CH}_3\text{COCH}_3 + \text{HCHO} + \text{HO}_2 + \text{OH}$	$k_{R02_HO2(temp,4)*r_{\text{COCH2O2_OH}}}$	Sander et al. (2019)
G44092a	TrGCN	$\text{IBUTOLBO2} + \text{NO} \rightarrow \text{CH}_3\text{COCH}_3 + \text{HCHO} + \text{HO}_2 + \text{NO}_2$	$k_{R02NO}*(1.-\alpha_{\text{AN}}(5,3,0,0,0,\text{temp},\text{cair}))$	Sander et al. (2019)
G44092b	TrGCN	$\text{IBUTOLBO2} + \text{NO} \rightarrow \text{IBUTOLBNO3}$	$k_{R02NO}*\alpha_{\text{AN}}(5,3,0,0,0,\text{temp},\text{cair})$	Sander et al. (2019)
G44093	TrGCN	$\text{IBUTOLBO2} + \text{NO}_3 \rightarrow \text{CH}_3\text{COCH}_3 + \text{HCHO} + \text{HO}_2 + \text{NO}_2$	k_{R02NO3}	Sander et al. (2019)
G44094a	TrGC	$\text{IBUTOLBOOH} + \text{OH} \rightarrow \text{IBUTOLBO2}$	k_{ROOHRO}	Sander et al. (2019)
G44094b	TrGC	$\text{IBUTOLBOOH} + \text{OH} \rightarrow \text{CH}_3\text{COCH}_3 + \text{HCHO} + \text{HO}_2$	$k_s*f_{\text{SOOH}}*f_{\text{pCH2OH}}$	Sander et al. (2019)
G44095	TrGCN	$\text{IBUTOLBNO3} + \text{OH} \rightarrow \text{CH}_3\text{COCH}_3 + \text{HCHO} + \text{HO}_2 + \text{NO}_2$	$3.*k_p$	Sander et al. (2019)
G44096	TrGC	$\text{BUT1ENE} + \text{OH} \rightarrow \text{LBUT1ENO2}$	$6.6E-12*\text{EXP}(465./\text{temp})$	Atkinson et al. (2006)*

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G44097a	TrGC	BUT1ENE + O ₃ → HCHO + .5 C ₂ H ₅ CHO + .5 H ₂ O ₂ + .5 CH ₃ CHO + .5 CO + .5 HO ₂	3.35E-15*EXP(-1745./temp)*.57	Atkinson et al. (2006), Sander et al. (2019)*
G44097b	TrGC	BUT1ENE + O ₃ → C ₂ H ₅ CHO + CH ₂ OO*	3.35E-15*EXP(-1745./temp)*.43	Atkinson et al. (2006), Sander et al. (2019)*
G44098	TrGCN	BUT1ENE + NO ₃ → C ₂ H ₅ CHO + HCHO + NO ₂	3.2E-13*EXP(-950./temp)	Atkinson et al. (2006), Sander et al. (2019)*
G44099	TrGC	LBUT1ENO2 → C ₂ H ₅ CHO + HCHO + HO ₂	k1_R02sOR02	Sander et al. (2019)
G44100a	TrGC	LBUT1ENO2 + HO ₂ → LBUT1ENOOH	k_R02_HO2(temp,4)*r_COCH202_00H	Sander et al. (2019)
G44100b	TrGC	LBUT1ENO2 + HO ₂ → C ₂ H ₅ CHO + HCHO + HO ₂ + OH	k_R02_HO2(temp,4)*r_COCH202_OH	Sander et al. (2019)
G44101a	TrGCN	LBUT1ENO2 + NO → C ₂ H ₅ CHO + HCHO + HO ₂ + NO ₂	KR02NO*(1.-alpha_AN(5,2,0,0,0,temp,cair))	Sander et al. (2019)
G44101b	TrGCN	LBUT1ENO2 + NO → LBUT1ENNO3	KR02NO*alpha_AN(5,2,0,0,0,temp,cair)	Sander et al. (2019)
G44102	TrGCN	LBUT1ENO2 + NO ₃ → C ₂ H ₅ CHO + HCHO + HO ₂ + NO ₂	KR02NO3	Sander et al. (2019)
G44103a	TrGC	LBUT1ENOOH + OH → LBUT1ENO2	k_ROOHRO	Sander et al. (2019)
G44103b	TrGC	LBUT1ENOOH + OH → C ₂ H ₅ CO ₃ + HCHO + HO ₂	k_t*f_t00H*f_pCH20H	Sander et al. (2019)*
G44104	TrGCN	LBUT1ENNO3 + OH → C ₂ H ₅ CHO + CO + HO ₂ + NO ₂	k_s*f_SOH*f_CH2ON02	Sander et al. (2019)*
G44105	TrGC	CBUT2ENE + OH → BUT2OLO2	1.1E-11*EXP(485./temp)	Atkinson et al. (2006)
G44106	TrGC	CBUT2ENE + O ₃ → CH ₃ CHO + .16 CH ₃ CHOHOOH + .50 OH + .50 HCOCH ₂ O ₂ + .05 CH ₂ CO + .09 CH ₃ OH + .09 CO + .2 CH ₄ + .2 CO ₂	3.2E-15*EXP(-965./temp)	Atkinson et al. (2006), Sander et al. (2019)*
G44107	TrGCN	CBUT2ENE + NO ₃ → 2 CH ₃ CHO + NO ₂	3.5E-13	Atkinson et al. (2006), Sander et al. (2019)*
G44108	TrGC	TBUT2ENE + OH → BUT2OLO2	1.0E-11*EXP(553./temp)	Atkinson et al. (2006)
G44109	TrGC	TBUT2ENE + O ₃ → CH ₃ CHO + .16 CH ₃ CHOHOOH + .50 OH + .50 HCOCH ₂ O ₂ + .05 CH ₂ CO + .09 CH ₃ OH + .09 CO + .2 CH ₄ + .2 CO ₂	6.6E-15*EXP(-1060./temp)	Atkinson et al. (2006), Sander et al. (2019)
G44110	TrGCN	TBUT2ENE + NO ₃ → 2 CH ₃ CHO + NO ₂	1.78E-12*EXP(-530./temp) +1.28E-14*EXP(570./temp)	Atkinson et al. (2006), Sander et al. (2019)*
G44111	TrGC	BUT2OLO2 → C ₂ H ₅ CHO + HCHO + HO ₂	k1_R02sOR02	Sander et al. (2019)
G44112a	TrGC	BUT2OLO2 + HO ₂ → BUT2OLOOH	k_R02_HO2(temp,4)*r_COCH202_00H	Sander et al. (2019)
G44112b	TrGC	BUT2OLO2 + HO ₂ → 2 CH ₃ CHO + HO ₂ + OH	k_R02_HO2(temp,4)*r_COCH202_OH	Sander et al. (2019)

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G44113a	TrGCN	BUT2OLO2 + NO → 2 CH ₃ CHO + HO ₂ + NO ₂	KR02NO*(1.-alpha_AN(5,2,0,0,0, temp,cair))	Sander et al. (2019)
G44113b	TrGCN	BUT2OLO2 + NO → BUT2OLNO3	KR02NO*alpha_AN(5,2,0,0,0,temp, cair)	Sander et al. (2019)
G44114	TrGCN	BUT2OLO2 + NO ₃ → 2 CH ₃ CHO + HO ₂ + NO ₂	KR02NO3	Sander et al. (2019)
G44115a	TrGC	BUT2OLOOH + OH → BUT2OLO2	k_ROOHRO	Sander et al. (2019)
G44115b	TrGC	BUT2OLOOH + OH → LMEKOOH + HO ₂	k_t*f_tOH*f_pCH2OH	Sander et al. (2019)
G44115c	TrGC	BUT2OLOOH + OH → BUT2OLO + OH	k_t*f_tOH*f_pCH2OH	Sander et al. (2019)
G44116	TrGCN	BUT2OLNO3 + OH → LMEKNO3 + HO ₂	k_t*f_tOH*f_CH2ON02	Sander et al. (2019)
G44117	TrGC	BUT2OLO + OH → BIACET + HO ₂	k_t*f_tOH*f_CO	Sander et al. (2019)
G44118	TrGC	IPRCHO + OH → IPRCO3 + H ₂ O	6.8E-12*EXP(410./temp)	Atkinson et al. (2006)
G44119	TrGCN	IPRCHO + NO ₃ → IPRCO3 + HNO ₃	1.67E-12*EXP(-1460./temp)	Atkinson et al. (2006)
G44120	TrGC	IPRCO3 → iC ₃ H ₇ O ₂ + CO ₂	k1_R02RC03	Rickard (2022)
G44121a	TrGC	IPRCO3 + HO ₂ → PERIBUACID	KAPH02*r_CO3_OOH	Rickard (2022), Sander et al. (2019)
G44121b	TrGC	IPRCO3 + HO ₂ → iC ₃ H ₇ O ₂ + CO ₂ + OH	KAPH02*(1.-r_CO3_OOH)	Rickard (2022), Sander et al. (2019)
G44122	TrGCN	IPRCO3 + NO ₂ → PIPN	k_CH3CO3_N02	Rickard (2022)
G44123	TrGCN	IPRCO3 + NO → iC ₃ H ₇ O ₂ + CO ₂ + NO ₂	KAPNO	Rickard (2022)
G44124a	TrGC	PERIBUACID + OH → IPRCO3 + H ₂ O	k_ROOHRO	Rickard (2022)
G44124b	TrGC	PERIBUACID + OH → CH ₃ COCH ₃ + H ₂ O + CO ₂	k_s*f_CO2H	Sander et al. (2019)*
G44125	TrGCN	PIPN → IPRCO3 + NO ₂	k_PAN_M	Rickard (2022)
G44126	TrGCN	PIPN + OH → CH ₃ COCH ₃ + CO ₂ + NO ₂	k_s*f_cpan	Sander et al. (2019)*
G44127	TrGC	MROPENOL + OH → HCOOH + OH + CH ₃ COCH ₃	k_CH2CHOH_OH_HCOOH	Sander et al. (2019), So et al. (2014)*
G44128	TrGC	MROPENOL + HCOOH → IPRCHO + HCOOH	k_CH2CHOH_HCOOH	Sander et al. (2019), da Silva (2010)*
G44129	TrGC	IPRCHO + HCOOH → MROPENOL + HCOOH	k_ALD_HCOOH	Sander et al. (2019), da Silva (2010)*
G44130	TrGC	BUTENOL + OH → HCOOH + OH + C ₂ H ₅ CHO	k_CH2CHOH_OH_HCOOH	Sander et al. (2019), So et al. (2014)*
G44131	TrGC	BUTENOL + HCOOH → C ₃ H ₇ CHO + HCOOH	k_CH2CHOH_HCOOH	Sander et al. (2019), da Silva (2010)*
G44132	TrGC	C ₃ H ₇ CHO + HCOOH → BUTENOL + HCOOH	k_ALD_HCOOH	Sander et al. (2019), da Silva (2010)*

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G44133	TrGC	HVMK + OH → HCOOH + OH + MGLYOX	8.8E-11	Sander et al. (2019), So et al. (2014), Messaadia et al. (2015)*
G44134	TrGC	HVMK + HCOOH → CO2C3CHO + HCOOH	k_CH2CHOH_HCOOH	Sander et al. (2019), da Silva (2010)*
G44135	TrGC	CO2C3CHO + HCOOH → HVMK + HCOOH	k_ALD_HCOOH	Sander et al. (2019), da Silva (2010)*
G44136	TrGC	HMAC + OH → HCOOH + OH + MGLYOX	8.8E-11	Sander et al. (2019), So et al. (2014), Messaadia et al. (2015)*
G44137	TrGC	HMAC + HCOOH → IBUTDIAL + HCOOH	k_CH2CHOH_HCOOH	Sander et al. (2019), da Silva (2010)*
G44138	TrGC	IBUTDIAL + HCOOH → HMAC + HCOOH	k_ALD_HCOOH	Sander et al. (2019), da Silva (2010)*
G44139	TrGC	CO2C3CHO + OH → CH ₃ COCH ₂ O ₂ + CO ₂ + H ₂ O	k_t*f_0*f_alk+k_s*f_CHO*f_CO	Sander et al. (2019)*
G44140	TrGCN	CO2C3CHO + NO ₃ → CH ₃ COCH ₂ O ₂ + CO ₂ + HNO ₃	KN03AL*4.0	Sander et al. (2019)*
G44141	TrGC	IBUTDIAL + OH → CH ₃ CHO + CO + HO ₂ + CO ₂ + H ₂ O	2.*k_t*f_0*f_alk+k_t*f_CHO*f_CHO	Sander et al. (2019)*
G44142	TrGCN	IBUTDIAL + NO ₃ → CH ₃ CHO + CO + HO ₂ + CO ₂ + HNO ₃	2.*KN03AL*4.0	Sander et al. (2019)*
G44200	TrGTerC	CH ₃ COCOCH ₂ O ₂ → CH ₃ C(O) + HCHO + CO	k1_R02p0R02	Rickard (2022)
G44201	TrGTerC	CH ₃ COCOCH ₂ O ₂ + HO ₂ → CH ₃ COCOCH ₂ OOH	k_R02_H02(temp, 4)	Rickard (2022)
G44202	TrGTerCN	CH ₃ COCOCH ₂ O ₂ + NO → CH ₃ C(O) + HCHO + CO + NO ₂	KR02NO	Rickard (2022)*
G44203a	TrGTerC	CH ₃ COCOCH ₂ OOH + OH → CH ₃ COCOCHO + OH	k_s*f_CO*f_sOOH	Rickard (2022)*
G44203b	TrGTerC	CH ₃ COCOCH ₂ OOH + OH → CH ₃ COCOCH ₂ O ₂	k_ROOHRO	Rickard (2022)
G44204	TrGTerC	C44O2 + HO ₂ → C44OOH	k_R02_H02(temp, 4)	Rickard (2022)
G44205	TrGTerCN	C44O2 + NO → HCOCH ₂ CHO + CO ₂ + HO ₂ + NO ₂	KR02NO	Rickard (2022)*
G44206	TrGTerC	C44O2 → HCOCH ₂ CHO + CO ₂ + HO ₂	k1_R02s0R02	Rickard (2022)
G44207	TrGTerC	C44OOH + OH → C44O2	7.46E-11	Rickard (2022)
G44208	TrGTerC	CHOC3COO ₂ → HCOCH ₂ CO ₃ + HCHO	k1_R02p0R02	Rickard (2022)
G44209	TrGTerC	CHOC3COO ₂ + HO ₂ → C413COOOH	k_R02_H02(temp, 4)	Rickard (2022)
G44210	TrGTerCN	CHOC3COO ₂ + NO → HCOCH ₂ CO ₃ + HCHO + NO ₂	KR02NO	Rickard (2022)*
G44211	TrGTerC	C413COOOH + OH → CHOC3COO ₂	8.33E-11	Rickard (2022)
G44212	TrGTerC	C4CODIAL + OH → C312COCO ₃	3.39E-11	Rickard (2022)
G44213	TrGTerCN	C4CODIAL + NO ₃ → C312COCO ₃ + HNO ₃	2.*KN03AL*4.0	Rickard (2022)
G44214	TrGTerC	C312COCO ₃ → HCOCOCH ₂ O ₂ + CO ₂	k1_R02RC03	Rickard (2022)

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G44215a	TrGTerC	C312COCO3 + HO ₂ → C312COCO3H	KAPH02*r_CO3_00H	Rickard (2022)
G44215b	TrGTerC	C312COCO3 + HO ₂ → HCOCOCH ₂ O ₂ + CO ₂ + OH	KAPH02*(1.-r_CO3_00H)	Rickard (2022)
G44216	TrGTerCN	C312COCO3 + NO ₂ → C312COPAN	k_CH3CO3_NO2	Rickard (2022)
G44217	TrGTerCN	C312COCO3 + NO → HCOCOCH ₂ O ₂ + CO ₂ + NO ₂	KAPNO	Rickard (2022)
G44218	TrGTerC	C312COCO3H + OH → C312COCO3	1.63E-11	Rickard (2022)
G44219	TrGTerCN	C312COPAN → C312COCO3 + NO ₂	k_PAN_M	Rickard (2022)
G44220	TrGTerCN	C312COPAN + OH → HCOCOCHO + CO + NO ₂	1.27E-11	Rickard (2022)
G44221	TrGTerC	CH ₃ COCOCHO + OH → CH ₃ C(O) + 2 CO	8.4E-13*EXP(830./temp)	Sander et al. (2019)*
G44222	TrGTerCN	CH ₃ COCOCHO + NO ₃ → CH ₃ C(O) + 2 CO + HNO ₃	KN03AL*4.0	Rickard (2022)
G44223	TrGTerC	IBUTALOH + OH → IPRHOCO3	1.4E-11	Rickard (2022)
G44224a	TrGTerC	IPRHOCO3 + HO ₂ → CH ₃ COCH ₃ + CO ₂ + HO ₂ + OH	KAPH02*r_CO3_OH	Rickard (2022), Sander et al. (2019)
G44224b	TrGTerC	IPRHOCO3 + HO ₂ → IPRHOCO2H + O ₃	KAPH02*r_CO3_03	Rickard (2022), Sander et al. (2019)
G44224c	TrGTerC	IPRHOCO3 + HO ₂ → IPRHOCO3H	KAPH02*r_CO3_00H	Rickard (2022), Sander et al. (2019)
G44225	TrGTerCN	IPRHOCO3 + NO → CH ₃ COCH ₃ + CO ₂ + HO ₂ + NO ₂	KAPNO	Rickard (2022)
G44226	TrGTerCN	IPRHOCO3 + NO ₂ → C4PAN5	k_CH3CO3_NO2	Rickard (2022)
G44227	TrGTerCN	IPRHOCO3 + NO ₃ → CH ₃ COCH ₃ + CO ₂ + HO ₂ + NO ₂	KR02NO3*1.74	Rickard (2022)
G44228a	TrGTerC	IPRHOCO3 → CH ₃ COCH ₃ + CO ₂ + HO ₂	k1_R02RC03*0.7	Rickard (2022)
G44228b	TrGTerC	IPRHOCO3 → IPRHOCO2H	k1_R02RC03*0.3	Rickard (2022)
G44229	TrGTerC	IPRHOCO2H + OH → CH ₃ COCH ₃ + CO ₂ + HO ₂ + H ₂ O	1.72E-12	Rickard (2022)
G44230	TrGTerC	OH + IPRHOCO3H → IPRHOCO3	4.80E-12	Rickard (2022)
G44231	TrGTerCN	C4PAN5 → IPRHOCO3 + NO ₂	k_PAN_M	Rickard (2022)
G44232	TrGTerCN	C4PAN5 + OH → CH ₃ COCH ₃ + CO + NO ₂	4.75E-13	Rickard (2022)
G44233a	TrGTerC	MBOOO → IPRHOCO2H	1.60E-17*C(ind_H2O)*(0.08+0.15)	Rickard (2022), Sander et al. (2019)
G44233b	TrGTerC	MBOOO → IBUTALOH + H ₂ O ₂	1.60E-17*C(ind_H2O)*0.77	Rickard (2022), Sander et al. (2019)
G44234	TrGTerC	MBOOO + CO → IBUTALOH + CO ₂	1.20E-15	Rickard (2022)
G44235	TrGTerCN	MBOOO + NO → IBUTALOH + NO ₂	1.00E-14	Rickard (2022)
G44236	TrGTerCN	MBOOO + NO ₂ → IBUTALOH + NO ₃	1.00E-15	Rickard (2022)
G44400	TrGAroC	MALANHY + OH → MALANHYO2	1.4E-12	Rickard (2022)
G44401a	TrGAroC	MALDIALOOH + OH → HOCOC4DIAL + OH	1.22E-10	Rickard (2022)
G44401b	TrGAroC	MALDIALOOH + OH → MALDIALO2	k_ROOHRO	Rickard (2022)

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G44402	TrGAroCN	NC4DCO2H + OH → MALANHY + NO ₂	k_ROOHRO	Rickard (2022)*
G44403	TrGAroC	CO14O3CO2H + OH → HCOCH ₂ O ₂ + 2 CO ₂	2.19E-11	Rickard (2022)
G44404	TrGAroC	BZFUOOH + OH → BZFUO ₂	3.68E-11	Rickard (2022)
G44405	TrGAroC	HOCOC4DIAL + OH → CO2C4DIAL + HO ₂	3.67E-11	Rickard (2022)
G44406a	TrGAroC	MALDIALCO ₃ + HO ₂ → MALDALCO2H + O ₃	KAPH02*r_CO3_03	Rickard (2022)
G44406b	TrGAroC	MALDIALCO ₃ + HO ₂ → MALDALCO3H	KAPH02*r_CO3_OOH	Rickard (2022)
G44406c	TrGAroC	MALDIALCO ₃ + HO ₂ → .6 MALANHY + HO ₂ + .4 GLYOX + .4 CO + .4 CO ₂ + OH	KAPH02*r_CO3_OH	Rickard (2022)*
G44407	TrGAroCN	MALDIALCO ₃ + NO → .6 MALANHY + HO ₂ + .4 GLYOX + .4 CO + .4 CO ₂ + NO ₂	KAPNO	Rickard (2022)*
G44408	TrGAroCN	MALDIALCO ₃ + NO ₂ → MALDIALPAN	k_CH3CO3_N02	Rickard (2022)
G44409	TrGAroCN	MALDIALCO ₃ + NO ₃ → .6 MALANHY + HO ₂ + .4 GLYOX + .4 CO + .4 CO ₂ + NO ₂	KR02N03*1.74	Rickard (2022)*
G44410	TrGAroC	MALDIALCO ₃ → .6 MALANHY + HO ₂ + .4 GLYOX + .4 CO + .4 CO ₂	k1_R02RC03	Rickard (2022)*
G44411	TrGAroCN	BZFUONE + NO ₃ → NBZFUO ₂	3.00E-13	Rickard (2022)
G44412	TrGAroC	BZFUONE + O ₃ → .3125 CO14O3CO2H + .1875 CO14O3CHO + .1875 H ₂ O ₂ + .5 CO + .5 CO ₂ + .5 HCOCH ₂ O ₂ + .5 OH	2.20E-19	see note*
G44413	TrGAroC	BZFUONE + OH → BZFUO ₂	4.45E-11	Rickard (2022)
G44414	TrGAroCN	NBZFUOOH + OH → NBZFUO ₂	6.18E-12	Rickard (2022)
G44415	TrGAroC	MALDALCO3H + OH → MALDIALCO ₃	4.00E-11	Rickard (2022)
G44416	TrGAroC	EPXDLCO2H + OH → C3DIALO ₂ + CO ₂	2.31E-11	Rickard (2022)
G44417a	TrGAroC	EPXDLCO ₃ + HO ₂ → C3DIALO ₂ + CO ₂ + OH	KAPH02*r_CO3_OH	Rickard (2022)
G44417b	TrGAroC	EPXDLCO ₃ + HO ₂ → EPXDLCO2H + O ₃	KAPH02*r_CO3_03	Rickard (2022)
G44417c	TrGAroC	EPXDLCO ₃ + HO ₂ → EPXDLCO3H	KAPH02*r_CO3_OOH	Rickard (2022)
G44418	TrGAroCN	EPXDLCO ₃ + NO → C3DIALO ₂ + CO ₂ + NO ₂	KAPNO	Rickard (2022)
G44419	TrGAroCN	EPXDLCO ₃ + NO ₂ → EPXDLPAN	k_CH3CO3_N02	Rickard (2022)
G44420	TrGAroCN	EPXDLCO ₃ + NO ₃ → C3DIALO ₂ + CO ₂ + NO ₂	KR02N03*1.74	Rickard (2022)
G44421	TrGAroC	EPXDLCO ₃ → C3DIALO ₂ + CO ₂	k1_R02RC03	Rickard (2022)*
G44422	TrGAroC	MALNHYOHCO + OH → CO + CO + CO + CO ₂ + HO ₂	5.68E-12	Rickard (2022)
G44423	TrGAroCN	MALDIAL + NO ₃ → MALDIALCO ₃ + HNO ₃	2.*KN03AL*2.0	Rickard (2022)
G44424	TrGAroC	MALDIAL + O ₃ → 1.0675 GLYOX + .125 HCHO + .1125 HCOCO ₂ H + .0675 H ₂ O ₂ + .82 HO ₂ + .57 OH + 1.265 CO + .25 CO ₂	2.00E-18	Rickard (2022)*

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G44425	TrGAroC	MALDIAL + OH → .83 MALDIALCO3 + .17 MALDIALO2	5.20E-11	Rickard (2022)*
G44426	TrGAroC	MALANHYOOH + OH → MALNHYOHCO + OH	4.66E-11	Rickard (2022)
G44427	TrGAroCN	MALDIALPAN + OH → GLYOX + CO + CO + NO ₂	3.70E-11	Rickard (2022)
G44428	TrGAroCN	MALDIALPAN → MALDIALCO3 + NO ₂	k_PAN_M	Rickard (2022)
G44429a	TrGAroC	MALANHYO2 + HO ₂ → MALANHYOOH	k_RO2_HO2(temp,4)*(1.-r_COCH202_OH-r_CHOHCH202_OH)	Rickard (2022), Sander et al. (2019)
G44429b	TrGAroC	MALANHYO2 + HO ₂ → HCOCOHC03 + CO ₂ + OH	k_RO2_HO2(temp,4)*(r_COCH202_OH+r_CHOHCH202_OH)	Rickard (2022), Sander et al. (2019)
G44430	TrGAroCN	MALANHYO2 + NO → HCOCOHC03 + CO ₂ + NO ₂	KR02NO	Rickard (2022)*
G44431	TrGAroCN	MALANHYO2 + NO ₃ → HCOCOHC03 + CO ₂ + NO ₂	KR02NO3	Rickard (2022)*
G44432	TrGAroC	MALANHYO2 → HCOCOHC03 + CO ₂	k1_RO2sOR02	Rickard (2022)*
G44433	TrGAroC	EPXDLCO3H + OH → EPXDLCO3	2.62E-11	Rickard (2022)
G44434	TrGAroC	CO2C4DIAL + OH → CO + CO + CO + CO + HO ₂	2.45E-11	Rickard (2022)
G44435a	TrGAroCN	NBZFUO2 + HO ₂ → NBZFUOOH	k_RO2_HO2(temp,4)*(1.-r_COCH202_OH)	Rickard (2022), Sander et al. (2019)
G44435b	TrGAroCN	NBZFUO2 + HO ₂ → .5 CO14O3CHO + .5 NO ₂ + .5 NBZFUONE + .5 HO ₂ + OH	k_RO2_HO2(temp,4)*r_COCH202_OH	Rickard (2022), Sander et al. (2019)
G44436	TrGAroCN	NBZFUO2 + NO → .5 CO14O3CHO + .5 NO ₂ + .5 NBZFUONE + .5 HO ₂ + NO ₂	KR02NO	Rickard (2022)*
G44437	TrGAroCN	NBZFUO2 + NO ₃ → .5 CO14O3CHO + .5 NO ₂ + .5 NBZFUONE + .5 HO ₂ + NO ₂	KR02NO3	Rickard (2022)*
G44438	TrGAroCN	NBZFUO2 → .5 CO14O3CHO + .5 NO ₂ + .5 NBZFUONE + .5 HO ₂	k1_RO2sOR02	Rickard (2022)*
G44439	TrGAroC	MALDALCO2H + OH → .6 MALANHY + HO ₂ + .4 GLYOX + .4 CO + .4 CO ₂	3.70E-11	Rickard (2022)*
G44440	TrGAroCN	EPXC4DIAL + NO ₃ → EPXDLCO3 + HNO ₃	2.*KN03AL*4.0	Rickard (2022)
G44441	TrGAroC	EPXC4DIAL + OH → EPXDLCO3	4.32E-11	Rickard (2022)
G44442a	TrGAroC	MECOACETO2 + HO ₂ → MECOACEOOH	k_RO2_HO2(temp,4)*(1.-r_COCH202_OH)	Rickard (2022), Sander et al. (2019)
G44442b	TrGAroC	MECOACETO2 + HO ₂ → CH ₃ C(O)OO + HCHO + CO ₂ + OH	k_RO2_HO2(temp,4)*r_COCH202_OH	Rickard (2022), Sander et al. (2019)
G44443	TrGAroCN	MECOACETO2 + NO → CH ₃ C(O)OO + HCHO + CO ₂ + NO ₂	KR02NO	Rickard (2022)*

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G44444	TrGAroCN	MECOACETO2 + NO ₃ → CH ₃ C(O)OO + HCHO + CO ₂ + NO ₂	KR02N03	Rickard (2022)*
G44445	TrGAroC	MECOACETO2 → CH ₃ C(O)OO + HCHO + CO ₂	k1_R02p0R02	Rickard (2022)*
G44446	TrGAroCN	CO14O3CHO + NO ₃ → CO + HCOCH ₂ O ₂ + CO ₂ + HNO ₃	KN03AL*8.0	Rickard (2022)
G44447	TrGAroC	CO14O3CHO + OH → CO + HCOCH ₂ O ₂ + CO ₂	3.44E-11	Rickard (2022)
G44448	TrGAroCN	NBZFUONE + OH → BZFUCO + NO ₂	1.16E-12	Rickard (2022)
G44449a	TrGAroC	BZFUO2 + HO ₂ → BZFUOOH	k_R02_H02(temp,4)*(1.-r_COCH202_OH-r_CHOHCH202_OH)	Rickard (2022), Sander et al. (2019)
G44449b	TrGAroC	BZFUO2 + HO ₂ → CO14O3CHO + HO ₂ + OH	k_R02_H02(temp,4)*(r_COCH202_OH+r_CHOHCH202_OH)	Rickard (2022), Sander et al. (2019)
G44450	TrGAroCN	BZFUO2 + NO → CO14O3CHO + HO ₂ + NO ₂	KR02NO	Rickard (2022)*
G44451	TrGAroCN	BZFUO2 + NO ₃ → CO14O3CHO + HO ₂ + NO ₂	KR02N03	Rickard (2022)*
G44452	TrGAroC	BZFUO2 → CO14O3CHO + HO ₂	k1_R02s0R02	Rickard (2022)*
G44453	TrGAroC	BZFUCO + OH → CO14O3CHO + HO ₂	1.78E-11	Rickard (2022)
G44456a	TrGAroC	MALDIALO2 + HO ₂ → MALDIALOOH	k_R02_H02(temp,4)*(1.-r_COCH202_OH-r_CHOHCH202_OH)	Rickard (2022)
G44456b	TrGAroC	MALDIALO2 + HO ₂ → GLYOX + GLYOX + HO ₂ + OH	k_R02_H02(temp,4)*(r_COCH202_OH+r_CHOHCH202_OH)	Rickard (2022)
G44457	TrGAroCN	MALDIALO2 + NO → GLYOX + GLYOX + HO ₂ + NO ₂	KR02NO	Rickard (2022)*
G44458	TrGAroCN	MALDIALO2 + NO ₃ → GLYOX + GLYOX + HO ₂ + NO ₂	KR02N03	Rickard (2022)*
G44459	TrGAroC	MALDIALO2 → GLYOX + GLYOX + HO ₂	k1_R02s0R02	Rickard (2022)*
G44460	TrGAroCN	EPXDL PAN + OH → HCOCOCHO + CO + NO ₂	2.29E-11	Rickard (2022)
G44461	TrGAroCN	EPXDL PAN → EPXDL CO ₃ + NO ₂	k_PAN_M	Rickard (2022)*
G44462	TrGAroC	MECOACEOOH + OH → MECOACETO2	3.59E-12	Rickard (2022)
G45000	TrGC	C ₅ H ₈ + O ₃ → .3508 MACR + .01518 MACO2H + .2440 MVK + .7085 HCHO + .11 CH ₂ OO + .1275 C ₃ H ₆ + .1575 CH ₃ C(O) + .0510 CH ₃ + .2625 HO ₂ + .27 OH + .09482 H ₂ O ₂ + .255 CO ₂ + .522 CO + .07182 HCHO + .03618 HCOCH ₂ O ₂ + .01782 CO + 0.05408 LCARBON	1.03E-14*EXP(-1995./temp)	Atkinson et al. (2006), Sander et al. (2019)
G45001	TrGC	C ₅ H ₈ + OH → .63 LISOPAB + .30 LISOPCD + .07 LISOPEFO2	2.7E-11*EXP(390./temp)	Atkinson et al. (2006), Sander et al. (2019)
G45002	TrGCN	C ₅ H ₈ + NO ₃ → NISOPO2	3.0E-12*EXP(-450./temp)	Atkinson et al. (2006)
G45003a	TrGC	LISOPAB + O ₂ → LISOPACO2	5.530E-13	Sander et al. (2019)

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G45003b	TrGC	LISOPAB + O ₂ → ISOPBO2	3.E-12	Sander et al. (2019)
G45004a	TrGC	LISOPCD + O ₂ → LDISOPACO2	6.780E-13	Sander et al. (2019)
G45004b	TrGC	LISOPCD + O ₂ → ISOPDO2	3.E-12	Sander et al. (2019)
G45005	TrGC	LISOPACO2 → LISOPAB + O ₂	3.1E12*exp(-7900./temp)*.6+ 7.8E13*exp(-8600./temp)*.4	Sander et al. (2019)
G45006	TrGC	ISOPBO2 → LISOPAB + O ₂	3.7E14*exp(-9570./temp) +4.2E14*exp(-9970./temp)	Sander et al. (2019)
G45007	TrGC	LDISOPACO2 → LISOPCD + O ₂	5.65E12*exp(-8410./temp) *.42+1.4E14*exp(-9110./temp)*.58	Sander et al. (2019)
G45008	TrGC	ISOPDO2 → LISOPCD + O ₂	5.0E14*exp(-10120./temp) +8.25E14*exp(-10220./temp)	Sander et al. (2019)
G45009a	TrGC	LISOPACO2 → C1ODC2O2C4OOH	k_16hsz14 * 2./3.*(1.-f_HPAL)	Sander et al. (2019)
G45009b	TrGC	LISOPACO2 → LZCDOC23DBCOOH + HO ₂	k_16hsz14 * (2./3.*f_HPAL + 1./3.)	Sander et al. (2019)
G45010a	TrGC	LDISOPACO2 → C1OOHC3O2C4OD	k_16hsz41 * 2./3.*(1.-f_HPAL)	Sander et al. (2019)
G45010b	TrGC	LDISOPACO2 → LZCDOC23DBCOOH + HO ₂	k_16hsz41 * (2./3.*f_HPAL + 1./3.)	Sander et al. (2019)
G45011	TrGC	LISOPACO2 → .9 LISOPACO + .1 ISOPAOH	k1_R02LISOPACO2	Rickard (2022), Sander et al. (2019)
G45012	TrGC	LISOPACO2 + HO ₂ → LISOPACOOH	k_R02_HO2(temp,5)	Rickard (2022)
G45013a	TrGCN	LISOPACO2 + NO → LISOPACO + NO ₂	KR02NO*(1.-alpha_AN(6,1,0,0,0, temp,cair))	Lockwood et al. (2010), Paulot et al. (2009a), Sander et al. (2019)
G45013b	TrGCN	LISOPACO2 + NO → LISOPACNO3	KR02NO*alpha_AN(6,1,0,0,0,temp, cair)	Lockwood et al. (2010), Paulot et al. (2009a), Sander et al. (2019)
G45014	TrGCN	LISOPACO2 + NO ₃ → LISOPACO + NO ₂	KR02NO3	Rickard (2022)
G45015	TrGC	LDISOPACO2 → .9 LISOPACO + .1 ISOPAOH	k1_R02LISOPACO2	Rickard (2022), Sander et al. (2019)
G45016	TrGC	LDISOPACO2 + HO ₂ → LISOPACOOH	k_R02_HO2(temp,5)	Rickard (2022)
G45017a	TrGCN	LDISOPACO2 + NO → LISOPACO + NO ₂	KR02NO*(1.-alpha_AN(6,1,0,0,0, temp,cair))	Lockwood et al. (2010), Paulot et al. (2009a), Sander et al. (2019)
G45017b	TrGCN	LDISOPACO2 + NO → LISOPACNO3	KR02NO*alpha_AN(6,1,0,0,0,temp, cair)	Lockwood et al. (2010), Paulot et al. (2009a), Sander et al. (2019)
G45018	TrGCN	LDISOPACO2 + NO ₃ → LISOPACO + NO ₂	KR02NO3	Rickard (2022)

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G45019a	TrGC	LISOPACOOH + OH → LISOPACO2	k_ROOHRO	Sander et al. (2019)
G45019b	TrGC	LISOPACOOH + OH → LZCDC23DBCOOH + HO ₂	k_s*f_allyl*f_sOH	Sander et al. (2019)
G45019c	TrGC	LISOPACOOH + OH → LHC4ACCHO + OH	(k_s*f_sOH*f_allyl+ k_ROHRO)	Sander et al. (2019)
G45019d	TrGC	LISOPACOOH + OH → LIEPOX + OH	(k_adt+k_ads)*a_CH2OH*a_CH200H	Sander et al. (2019)*
G45020	TrGC	ISOPAOH + OH → LHC4ACCHO + HO ₂	(k_adt+k_ads)*a_CH2OH*a_CH20H+k_s*f_sOH*f_allyl+k_ROHRO	Sander et al. (2019)
G45021	TrGCN	LISOPACNO3 + OH → LISOPACNO3O2	(k_adt+k_ads)*a_CH2ON02*a_CH2OH	Sander et al. (2019)*
G45022	TrGC	ISOPBO2 → .8 MVK + .8 HCHO + .8 HO ₂ + .2 ISOPBOH	k1_RO2ISOPBO2	Rickard (2022)
G45023a	TrGC	ISOPBO2 + HO ₂ → ISOPBOOH	k_RO2_HO2(temp,5)*(1.-r_CHOHCH202_OH)	Sander et al. (2019)
G45023b	TrGC	ISOPBO2 + HO ₂ → MVK + HCHO + HO ₂ + OH	k_RO2_HO2(temp,5)*r_CHOHCH202_OH	Sander et al. (2019)
G45024a	TrGCN	ISOPBO2 + NO → MVK + HCHO + HO ₂ + NO ₂	KR02NO*(1.-alpha_AN(6,3,0,0,0, temp, cair))	Lockwood et al. (2010), Sander et al. (2019)
G45024b	TrGCN	ISOPBO2 + NO → ISOPBNO3	KR02NO*alpha_AN(6,3,0,0,0,temp, cair)	Lockwood et al. (2010), Sander et al. (2019)
G45025	TrGCN	ISOPBO2 + NO ₃ → MVK + .75 HCHO + .75 HO ₂ + .25 CH ₃ + NO ₂	KR02N03	Rickard (2022)
G45026a	TrGC	ISOPBOOH + OH → LIEPOX + OH	(k_ads+k_adp)*a_CH200H	Paulot et al. (2009b), Sander et al. (2019)
G45026b	TrGC	ISOPBOOH + OH → ISOPBO2	k_ROOHRO	Sander et al. (2019)
G45026c	TrGC	ISOPBOOH + OH → MGLYOX + HOCH ₂ CHO	k_ROHRO+k_s*f_alk*f_sOH	Sander et al. (2019)
G45027	TrGC	ISOPBOOH + O ₃ → .1368 MACROOH + .1368 H ₂ O ₂ + .2280 HO ₂ + .4332 CH ₃ COCH ₂ OH + .2280 CO ₂ + .6384 OH + .2052 CO + .57 HCHO + .43 MACROOH + .06880 HO ₂ + .06880 OH + .2709 CO + .1591 CH ₂ OO	1.E-17	Sander et al. (2019)
G45028	TrGC	ISOPBOH + OH → MVK + .75 HCHO + .75 HO ₂ + .25 CH ₃	k_s*f_alk*f_sOH+(k_adp+k_ads)*a_CH2OH	Sander et al. (2019)
G45029	TrGCN	ISOPBNO3 + OH → ISOPBDNO3O2	(k_adt+k_adp)*f_CH2ON02	Sander et al. (2019)
G45030	TrGC	ISOPDO2 → .8 MACR + .8 HCHO + .8 HO ₂ + .1 HCOC5 + .1 ISOPDOH	k1_RO2ISOPD02	Rickard (2022)
G45031a	TrGC	ISOPDO2 + HO ₂ → ISOPDOOH	k_RO2_HO2(temp,5)*(1.-r_CHOHCH202_OH)	Sander et al. (2019)
G45031b	TrGC	ISOPDO2 + HO ₂ → MACR + HCHO + HO ₂ + OH	k_RO2_HO2(temp,5)*r_CHOHCH202_OH	Sander et al. (2019)
G45032a	TrGCN	ISOPDO2 + NO → MACR + HCHO + HO ₂ + NO ₂	KR02NO*(1.-alpha_AN(6,2,0,0,0, temp, cair))	Lockwood et al. (2010), Sander et al. (2019)

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G45032b	TrGCN	ISOPDO2 + NO → ISOPDNO3	KR02NO*alpha_AN(6, 2, 0, 0, 0, temp, cair)	Lockwood et al. (2010), Sander et al. (2019)
G45033	TrGCN	ISOPDO2 + NO ₃ → MACR + HCHO + HO ₂ + NO ₂	KR02NO3	Rickard (2022)
G45034a	TrGC	ISOPDOOH + OH → LIEPOX + OH	(k_adt+k_adp)*a_CH20OH	Paulot et al. (2009b), Sander et al. (2019)
G45034b	TrGC	ISOPDOOH + OH → ISOPDO2	k_ROOHRO	Sander et al. (2019)
G45034c	TrGC	ISOPDOOH + OH → HCOC5 + OH	k_t*f_t0OH*f_allyl*f_pCH2OH	Sander et al. (2019)
G45034d	TrGC	ISOPDOOH + OH → CH ₃ COCH ₂ OH + GLYOX + OH	k_s*f_pCH2OH*f_sOH	Sander et al. (2019)
G45035	TrGC	ISOPDOOH + O ₃ → 1.393 OH + BIACETOH + .67 HCHO + .05280 HO ₂ + .2079 CO + .1221 CH ₂ OO	1.E-17	Sander et al. (2019)
G45036	TrGC	ISOPDOH + OH → HCOC5 + HO ₂	2.*k_ROHRO+(k_t*f_t0H*f_allyl+k_s*f_sOH)*f_pCH2OH+(k_adt+k_adp)*a_CH20H	Sander et al. (2019)
G45037	TrGCN	ISOPDNO3 + OH → ISOPBDNO3O2	(k_adp+k_ads)*a_CH20N02	Sander et al. (2019)*
G45038	TrGCN	NISOPO2 → .8 NC4CHO + .6 HO ₂ + .2 LISOPACNO3	k1_R02LISOPACO2	Rickard (2022)
G45039	TrGCN	NISOPO2 + HO ₂ → NISOPOOH	k_R02_H02(temp,5)	Rickard (2022)
G45040	TrGCN	NISOPO2 + NO → NC4CHO + HO ₂ + NO ₂	KR02NO	Rickard (2022)*
G45041	TrGCN	NISOPO2 + NO ₃ → NC4CHO + HO ₂ + NO ₂	KR02NO3	Rickard (2022)
G45042	TrGCN	NISOPOOH + OH → NC4CHO + OH	1.03E-10	Rickard (2022)
G45043	TrGCN	NC4CHO + OH → LNISO3	(k_adt+k_ads)*a_CHO*a_CH20N02	Sander et al. (2019)*
G45044	TrGCN	NC4CHO + O ₃ → .27 NOA + .027 HCOCO ₂ H + .0162 GLYOX + .0162 H ₂ O ₂ + .1458 HCOCO + .0405 HCOOH + .0405 CO + .8758 OH + .365 MGLYOX + .73 NO ₂ + 0.7705 HCHO + .4055 CO ₂ + .73 GLYOX	2.40E-17	Sander et al. (2019)
G45045	TrGCN	NC4CHO + NO ₃ → LNISO3 + HNO ₃	KN03AL*4.25	Rickard (2022)
G45046	TrGCN	LNISO3 + HO ₂ → LNISOOH	0.5*k_R02_H02(temp,5)+0.5*KAPH02	Rickard (2022)
G45047	TrGCN	LNISO3 + NO → NOA + .5 HOCHCHO + .5 CO + .5 HO ₂ + NO ₂ + .5 CO ₂	0.5*KAPNO+0.5*KR02NO	Rickard (2022)*
G45048	TrGCN	LNISO3 + NO ₃ → NOA + .5 HOCHCHO + .5 CO + .5 HO ₂ + NO ₂ + .5 CO ₂	KR02NO3*1.37	Rickard (2022)
G45049	TrGCN	LNISOOH + OH → LNISO3	2.65E-11	Rickard (2022)
G45050a	TrGC	LHC4ACCHO + OH → LC578O2	(k_adtertpim+k_ads)*a_CHO*a_CH20H	Sander et al. (2019)
G45050b	TrGC	LHC4ACCHO + OH → LHC4ACCO3	k_t*f_0	Sander et al. (2019)
G45050c	TrGC	LHC4ACCHO + OH → C4MDIAL + HO ₂	k_s*f_sOH*f_allyl	Sander et al. (2019)

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G45051	TrGC	LHC4ACCHO + O ₃ → .2225 CH ₃ C(O) + .89 CO + .0171875 HOCH ₂ CO ₂ H + .075625 H ₂ O ₂ + .0171875 HCOCO ₂ H + .2775 CH ₃ COCH ₂ OH + .6675 HO ₂ + .2603125 GLYOX + .2225 HCHO + .89 OH + .2603125 HOCH ₂ CHO + .5 MGLYOX	2.40E-17	Rickard (2022)
G45052	TrGCN	LHC4ACCHO + NO ₃ → LHC4ACCO3 + HNO ₃	KNO3AL*4.25	Rickard (2022)
G45053	TrGC	LC578O2 → .25 CH ₃ COCH ₂ OH + .75 MGLYOX + .25 HOCHCHO + .75 HOCH ₂ CHO + .75 HO ₂	k1_R02t0R02	Rickard (2022)
G45054a	TrGC	LC578O2 + HO ₂ → MGLYOX + HOCH ₂ CHO + OH	k_R02_HO2(temp,5)*r_COCH2O2_OH	Rickard (2022)
G45054b	TrGC	LC578O2 + HO ₂ → LC578OOH	k_R02_HO2(temp,5)*r_COCH2O2_OOH	Rickard (2022)
G45055	TrGCN	LC578O2 + NO → .25 CH ₃ COCH ₂ OH + .75 MGLYOX + .25 HOCHCHO + .75 HOCH ₂ CHO + .75 HO ₂ + NO ₂	KR02NO	Rickard (2022)*
G45056	TrGCN	LC578O2 + NO ₃ → .25 CH ₃ COCH ₂ OH + .75 MGLYOX + .25 HOCHCHO + .75 HOCH ₂ CHO + .75 HO ₂ + NO ₂	KR02NO3	Rickard (2022)
G45057	TrGC	LC578O2 → .25 CH ₃ COCH ₂ OH + .75 MGLYOX + .25 HOCH ₂ CHO + .75 HO ₂ + OH	k_hsb	Sander et al. (2019)
G45058a	TrGC	LC578OOH + OH → LC578O2	k_ROOHRO	Sander et al. (2019)
G45058b	TrGC	LC578OOH + OH → C1ODC2OOHC4OD + HO ₂	k_t*f_0*f_tCH2OH*f_alk+k_t*f_tOH*f_pCH2OH*f_pCH2OH+k_s*f_sOH*f_pCH2OH	Sander et al. (2019)
G45059a	TrGC	LHC4ACCO3 → OH + .5 MACRO2 + .5 LHMVKABO2 + CO ₂	k1_R02RC03*0.9	Sander et al. (2019)
G45059b	TrGC	LHC4ACCO3 → LHC4ACCO2H	k1_R02RC03*0.1	Sander et al. (2019)
G45060a	TrGC	LHC4ACCO3 + HO ₂ → 2 OH + .5 MACRO2 + .5 LHMVKABO2 + CO ₂	KAPH02*r_C03_OH	Sander et al. (2019)
G45060b	TrGC	LHC4ACCO3 + HO ₂ → LHC4ACCO3H	KAPH02*r_C03_OOH	Sander et al. (2019)
G45060c	TrGC	LHC4ACCO3 + HO ₂ → LHC4ACCO2H + O ₃	KAPH02*r_C03_03	Sander et al. (2019)
G45061	TrGCN	LHC4ACCO3 + NO → .5 MACRO2 + .5 LHMVKABO2 + NO ₂ + CO ₂	KAPNO	Sander et al. (2019)
G45062	TrGCN	LHC4ACCO3 + NO ₂ → LC5PAN1719	k_CH3C03_N02	Rickard (2022)
G45063	TrGCN	LHC4ACCO3 + NO ₃ → .5 MACRO2 + .5 LHMVKABO2 + NO ₂ + CO ₂	KR02NO3*1.74	Sander et al. (2019)
G45064a	TrGC	LHC4ACCO2H + OH → OH + .5 MACRO2 + .5 LHMVKABO2 + CO ₂	2.52E-11	Sander et al. (2019)
G45064b	TrGC	LHC4ACCO3H + OH → LHC4ACCO3	2.88E-11	Rickard (2022)

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G45065	TrGCN	$\text{LC5PAN1719} \rightarrow \text{LHC4ACCO3} + \text{NO}_2$	k_PAN_M	Rickard (2022)
G45066	TrGCN	$\text{LC5PAN1719} + \text{OH} \rightarrow .5 \text{ MACROH} + .5 \text{ HO12CO3C4} + \text{CO} + \text{NO}_2$	2.52E-11	Rickard (2022)
G45067	TrGC	$\text{HCOC5} + \text{OH} \rightarrow \text{C59O2}$	3.81E-11	Rickard (2022)
G45068	TrGC	$\text{HCOC5} + \text{O}_3 \rightarrow \text{BIACETOH} + .335 \text{ H}_2\text{O}_2 + .67 \text{ HCHO} + .2079 \text{ CO} + .1221 \text{ CH}_2\text{OO} + .05280 \text{ OH}$	$7.51\text{E}-16 * \text{EXP}(-1521./\text{temp})$	Sander et al. (2019)
G45069	TrGC	$\text{C59O2} \rightarrow \text{CH}_3\text{COCH}_2\text{OH} + \text{HOCH}_2\text{CO}$	k1_R02t0R02	Sander et al. (2019)
G45070a	TrGC	$\text{C59O2} + \text{HO}_2 \rightarrow \text{OH} + \text{CH}_3\text{COCH}_2\text{OH} + \text{HOCH}_2\text{CO}$	$k_{\text{R02}_*\text{HO2}}(\text{temp}, 5) * r_{\text{COCH202}_*\text{OH}}$	Sander et al. (2019)
G45070b	TrGC	$\text{C59O2} + \text{HO}_2 \rightarrow \text{C59OOH}$	$k_{\text{R02}_*\text{HO2}}(\text{temp}, 5) * r_{\text{COCH202}_*\text{OOH}}$	Sander et al. (2019)
G45071	TrGCN	$\text{C59O2} + \text{NO} \rightarrow \text{CH}_3\text{COCH}_2\text{OH} + \text{HOCH}_2\text{CO} + \text{NO}_2$	KR02NO	Sander et al. (2019)*
G45072	TrGCN	$\text{C59O2} + \text{NO}_3 \rightarrow \text{CH}_3\text{COCH}_2\text{OH} + \text{HOCH}_2\text{CO} + \text{NO}_2$	KR02NO3	Sander et al. (2019)
G45073	TrGC	$\text{C59OOH} + \text{OH} \rightarrow \text{C59O2}$	9.7E-12	Rickard (2022)
G45074	TrGC	$\text{LIEPOX} + \text{OH} \rightarrow \text{DB1O2} + \text{H}_2\text{O}$	$5.78\text{E}-11 * \text{EXP}(-400./\text{temp}) * (1.52/3.+0.98*2./3.)/1.51$	Paulot et al. (2009b), Bates et al. (2014), Sander et al. (2019)*
G45075	TrGC	$\text{ISOPBO2} \rightarrow \text{MVK} + \text{HCHO} + \text{OH}$	k_hsb	Sander et al. (2019)
G45076	TrGC	$\text{ISOPDO2} \rightarrow \text{MACR} + \text{HCHO} + \text{OH}$	k_hsd	Sander et al. (2019)
G45077a	TrGC	$\text{LZC0DC23DBC0OH} + \text{OH} \rightarrow .6 \text{ C10DC2O2C4OOH} + .4 \text{ C1OOHC2O2C4OD}$	$k_{\text{adt}} * a_{\text{CHO}} * a_{\text{CH200H}}$	Sander et al. (2019)
G45077b	TrGC	$\text{LZC0DC23DBC0OH} + \text{OH} \rightarrow .6 \text{ C10DC3O2C4OOH} + .4 \text{ C1OOHC3O2C4OD}$	$k_{\text{ads}} * a_{\text{CHO}} * a_{\text{CH200H}}$	Sander et al. (2019)
G45077c	TrGC	$\text{LZC0DC23DBC0OH} + \text{OH} \rightarrow \text{LZCO3HC23DBC0D}$	$k_t * f_0 * f_{\text{alk}} + k_{\text{ROOHRO}}$	Sander et al. (2019)
G45077d	TrGC	$\text{LZC0DC23DBC0OH} + \text{OH} \rightarrow \text{C4MDIAL} + \text{OH}$	$k_s * f_{\text{SOOH}} * f_{\text{allyl}}$	Sander et al. (2019)
G45078	TrGC	$\text{LZC0DC23DBC0OH} + \text{O}_3 \rightarrow .4672 \text{ OH} + .2336 \text{ HCOCOCH}_2\text{O}_2 + .2336 \text{ CO} + .2336 \text{ CH}_3\text{C(O)} + .4672 \text{ HOOCH}_2\text{CHO} + .1728 \text{ MGLYOX} + .1901 \text{ OH} + .0864 \text{ GLYOX} + .02765 \text{ HOOCH}_2\text{CHO} + .02765 \text{ H}_2\text{O}_2 + .02592 \text{ CH}_3\text{OOH} + .02592 \text{ CO}_2 + .01037 \text{ HCOCO} + .01555 \text{ CH}_2\text{OO} + .01555 \text{ CO} + .006908 \text{ HOOCH}_2\text{CO}_3 + .2628 \text{ OH} + .1314 \text{ MGLYOX} + .1314 \text{ OH} + .1314 \text{ HCOCOCH}_2\text{OOH} + .2628 \text{ GLYOX} + .0972 \text{ CH}_3\text{COCH}_2\text{O}_2\text{H} + .00972 \text{ HCOCO}_2\text{H} + .005832 \text{ GLYOX} + .005832 \text{ H}_2\text{O}_2 + .05249 \text{ OH} + .05249 \text{ HCOCO} + .01458 \text{ HCHO} + .01458 \text{ CO}_2 + .01458 \text{ HCOOH} + .01458 \text{ CO}$	2.4E-17	Sander et al. (2019)
G45079	TrGC	$\text{C1OOHC2O2C4OD} \rightarrow .78 \text{ CH}_3\text{COCH}_2\text{O}_2\text{H} + .78 \text{ HOCHCHO} + .22 \text{ CO}_2\text{H}_3\text{CHO} + .22 \text{ HCHO} + .22 \text{ OH}$	k1_R02t0R02	Sander et al. (2019)

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G45080	TrGCN	C1OOHC2O2C4OD + NO → .78 CH ₃ COCH ₂ O ₂ H + .78 HOCHCHO + .22 CO ₂ H3CHO + .22 HCHO + .22 OH + NO ₂	KR02NO	Sander et al. (2019)*
G45081a	TrGC	C1OOHC2O2C4OD + HO ₂ → C1OOHC2OOHC4OD	k_R02_H02(temp,5)*r_COCH202_00H	Sander et al. (2019)
G45081b	TrGC	C1OOHC2O2C4OD + HO ₂ → .78 CH ₃ COCH ₂ O ₂ H + .78 HOCHCHO + .22 CO ₂ H3CHO + .22 HCHO + 1.22 OH	k_R02_H02(temp,5)*r_COCH202_OH	Sander et al. (2019)
G45082	TrGC	C1OOHC2O2C4OD → CH ₃ COCH ₂ O ₂ H + GLYOX + OH	k_hsb	Sander et al. (2019)
G45083	TrGC	C1ODC2O2C4OOH → OH + C1ODC2OOHC4OD	k_15hsdhb	Sander et al. (2019)
G45084a	TrGC	C1OOHC2OOHC4OD + OH → C1ODC2OOHC4OD + OH	2.*k_s*f_s00H*f_tCH20H	Sander et al. (2019)
G45084b	TrGC	C1OOHC2OOHC4OD + OH → CH ₃ COCH ₂ O ₂ H + 2 CO + 2 HO ₂ + OH	k_t*f_tOH*f_pCH20H*f_pCH20H	Sander et al. (2019)
G45084c	TrGC	C1OOHC2OOHC4OD + OH → C1OOHC2O2C4OD	k_ROOHR0	Sander et al. (2019)
G45085	TrGC	C1ODC2OOHC4OD + OH → CO ₂ H3CHO + CO + H ₂ O + OH	k_t*f_0*f_tCH20H+k_t*f_tOH*f_tOH*f_CHO	Sander et al. (2019)
G45086	TrGC	C1ODC3O2C4OOH → MGLYOX + HOOCH ₂ CHO + HO ₂	k1_R02s0R02	Sander et al. (2019)
G45087	TrGCN	C1ODC3O2C4OOH + NO → MGLYOX + HOOCH ₂ CHO + HO ₂ + NO ₂	KR02NO	Sander et al. (2019)
G45088	TrGC	C1ODC3O2C4OOH + HO ₂ → .5 CH ₃ C(O) + .5 CO + .5 MGLYOX + .5 HO ₂ + HOOCH ₂ CO ₃	k_R02_H02(temp,5)	Sander et al. (2019)
G45089	TrGC	C1ODC3O2C4OOH → MGLYOX + OH + HOOCH ₂ CHO	k_hsd	Sander et al. (2019)
G45090	TrGC	C1OOHC3O2C4OD → .625 MGLYOX + 2 CO + 1.625 HO ₂ + .375 CH ₃ C(O) + .375 CO ₂ + OH	k_15hsdhb	Sander et al. (2019)
G45091	TrGC	LHC4ACCO3 → LZCO3HC23DBCOD + HO ₂	k_16hs	Sander et al. (2019)
G45092a	TrGC	C4MDIAL + OH → C1ODC2O2C4OD	(k_adt+k_ads)*a_CHO*a_CHO	Sander et al. (2019)*
G45092b	TrGC	C4MDIAL + OH → LZCO3C23DBCOD	2.*k_t*f_0*f_alk	Sander et al. (2019)*
G45093	TrGCN	C4MDIAL + NO ₃ → LZCO3C23DBCOD + HNO ₃	KN03AL*4.25*2.	Sander et al. (2019)*
G45094a	TrGC	C1ODC2O2C4OD + HO ₂ → OH + MGLYOX + HOCHCHO	k_R02_H02(temp,5)*r_COCH202_OH	Sander et al. (2019)
G45094b	TrGC	C1ODC2O2C4OD + HO ₂ → C1ODC2OOHC4OD	k_R02_H02(temp,5)*r_COCH202_00H	Sander et al. (2019)
G45095	TrGCN	C1ODC2O2C4OD + NO → NO ₂ + MGLYOX + HOCHCHO	KR02NO	Sander et al. (2019)*
G45096	TrGC	C1ODC2O2C4OD → MGLYOX + HOCHCHO	k1_R02t0R02	Sander et al. (2019)

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G45097a	TrGC	C1ODC2OOHC4OD + OH → MGLYOX + 2 CO	(2.*k_t*f_0*f_tCH2OH*f_alk+k_t*f_tOH*f_CHO*f_pCH2OH)*.5	Sander et al. (2019)
G45097b	TrGC	C1ODC2OOHC4OD + OH → MGLYOX + 2 CO + OH	(2.*k_t*f_0*f_tCH2OH*f_alk+k_t*f_tOH*f_CHO*f_pCH2OH)*.5	Sander et al. (2019)
G45098	TrGCN	LISOPACNO3O2 + NO → .21 NOA + .21 HOCH ₂ CHO + .21 HO ₂ + .49 HO12CO3C4 + .49 HCHO + .49 NO ₂ + .045 MVKNO3 + .045 HCHO + .255 CH ₃ COCH ₂ OH + .255 NO ₃ CH ₂ CHO + .225 H ₂ O ₂ + NO ₂	KR02NO	Sander et al. (2019)*
G45099	TrGCN	LISOPACNO3O2 → .21 NOA + .21 HOCH ₂ CHO + .21 HO ₂ + .49 HO12CO3C4 + .49 HCHO + .49 NO ₂ + .045 MVKNO3 + .045 HCHO + .255 CH ₃ COCH ₂ OH + .255 NO ₃ CH ₂ CHO + .225 H ₂ O ₂	k1_R02t0R02+k_R02_H02(temp,5)*c(ind_H02)	Sander et al. (2019)
G45100	TrGCN	ISOPBDNO3O2 + NO → .6 CH ₃ COCH ₂ OH + .6 HOCH ₂ CHO + .26 MACRNO3 + .14 MVKNO3 + .4 HCHO + .4 HO ₂ + 1.6 NO ₂	KR02NO	Sander et al. (2019)*
G45101	TrGCN	ISOPBDNO3O2 → .6 CH ₃ COCH ₂ OH + .6 HOCH ₂ CHO + .26 MACRNO3 + .14 MVKNO3 + .4 HCHO + .4 HO ₂ + .6 NO ₂	k1_R02s0R02+k_R02_H02(temp,5)*c(ind_H02)	Sander et al. (2019)
G45102	TrGCN	LISOPACNO3 + O ₃ → .8704 OH + .365 HO ₂ + .73 MGLYOX + .4325 NO ₃ CH ₂ CHO + .135 CH ₃ COCH ₂ OH + .0675 GLYOX + .4325 NO ₂ + .0891 H ₂ O ₂ + .135 NOA + .0675 HOCHCHO + .3866 HOCH ₂ CHO + .0405 CH ₃ OH + .0405 CO + .0054 HOCH ₂ CO	2.8E-17	Feierabend et al. (2008), Sander et al. (2019)
G45103	TrGC	DB1O2 → DB1O2	k1_R02s0R02	Sander et al. (2019)
G45104a	TrGC	DB1O2 + HO ₂ → DB1OOH	k_R02_H02(temp,5)*(1.-r_CHOCH2O2_OH)	Sander et al. (2019)*
G45104b	TrGC	DB1O2 + HO ₂ → DB1O2 + OH	k_R02_H02(temp,5)*r_CHOCH2O2_OH	Sander et al. (2019)
G45105a	TrGCN	DB1O2 + NO → DB1O2 + NO ₂	KR02NO*(1.-alpha_AN(7,2,0,0,0,temp,cair))	Sander et al. (2019)
G45105b	TrGCN	DB1O2 + NO → DB1NO3	KR02NO*alpha_AN(7,2,0,0,0,temp,cair)	Sander et al. (2019)
G45106	TrGCN	DB1O2 + NO ₃ → DB1O2 + NO ₂	KR02NO3	Sander et al. (2019)
G45107	TrGC	DB1O2 → DB1O2 + OH	1.E4	Peeters and Nguyen (2012)*
G45108a	TrGC	DB1O2 → DB1O2	KDEC*0.72	see note*
G45108b	TrGC	DB1O2 → .5 HVMK + .5 HMAC + HCHO + HO ₂	KDEC*0.28	see note*

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G45109	TrGC	DB1O2 → .48 CH ₃ COCH ₂ OH + .52 HOCH ₂ CHO + .52 MGLYOX + .48 GLYOX + HO ₂	k1_R02sOR02	Sander et al. (2019)
G45110a	TrGC	DB1O2 + HO ₂ → DB2OOH	k_R02_HO2(temp,5)*(1.-r_CHOCH2O2_OH)	Sander et al. (2019)
G45110b	TrGC	DB1O2 + HO ₂ → .48 CH ₃ COCH ₂ OH + .52 HOCH ₂ CHO + .52 MGLYOX + .48 GLYOX + HO ₂ + OH	k_R02_HO2(temp,5)*r_CHOCH2O2_OH	Sander et al. (2019)
G45111	TrGCN	DB1O2 + NO → .48 CH ₃ COCH ₂ OH + .52 HOCH ₂ CHO + .52 MGLYOX + .48 GLYOX + HO ₂ + NO ₂	KR02NO	see note*
G45112	TrGCN	DB1O2 + NO ₃ → .48 CH ₃ COCH ₂ OH + .52 HOCH ₂ CHO + .52 MGLYOX + .48 GLYOX + HO ₂ + NO ₂	KR02N03	Sander et al. (2019)
G45113	TrGC	DB1O2 → .48 MACROOH + .52 LHMVKABOOH + CO + OH	k_14hsal	Sander et al. (2019)
G45114a	TrGC	DB1OOH + OH → DB1O2	k_ROOHRO	Sander et al. (2019)
G45114b	TrGC	DB1OOH + OH → HCOOH + HO ₂ + CH ₃ COCHO ₂ CHO	k_adt	Sander et al. (2019)*
G45115	TrGC	DB1OOH + HCOOH → C1ODC2OOHC4OD + HCOOH	4.67E-26*(temp)**(3.286)*EXP(4509./(1.987*temp))	Sander et al. (2019), da Silva (2010)*
G45116	TrGCN	DB1NO3 + OH → HCOOH + NO ₂ + CH ₃ COCHO ₂ CHO	k_adt	Sander et al. (2019)*
G45117	TrGC	DB2OOH + OH → DB1O2	k_ROOHRO	Sander et al. (2019)*
G45118	TrGC	LISOPACOOH + O ₃ → 1.3272 OH + .36986 HO ₂ + .0432 H ₂ O ₂ + .08422 CO + .2025 CH ₃ OOH + .01215 CH ₂ OO + .3704 HCHO + .00405 CH ₃ OH + .0405 CO ₂ + .1825 HOCH ₂ COCH ₂ O ₂ + .365 MGLYOX + .3866 HOOCH ₂ CHO + .135 CH ₃ COCH ₂ OH + .0675 GLYOX + .00324 HCOCO + .3866 HOCH ₂ CHO + .135 CH ₃ COCH ₂ O ₂ H + .0675 HOCHCHO + .0054 HOCH ₂ CO	4.829E-16	Sander et al. (2019)
G45119a	TrGC	LZCO3HC23DBCOD + OH → .62 CO2H3CHO + .62 OH + .62 CO ₂ + .38 MGLYOX + .38 HCOCO ₃ H + .38 HO ₂	k_adt*a_CHO*a_CO2H	Sander et al. (2019)
G45119b	TrGC	LZCO3HC23DBCOD + OH → .62 CH ₃ COCO ₃ H + 1.24 CO + 1.24 HO ₂ + .38 MGLYOX + .38 HO ₂ + .38 CO + .38 HO ₂ + .38 OH + .38 CO ₂	k_ads*a_CHO*a_CO2H	Sander et al. (2019)
G45120	TrGC	LISOPEFO2 → LISOPEFO	k1_R02pOR02	Sander et al. (2019)
G45121a	TrGCN	LISOPEFO2 + NO → LISOPEFO + NO ₂	KR02NO*(1.-alpha_AN(6,1,0,0,0,temp,cair))	Sander et al. (2019)
G45121b	TrGCN	LISOPEFO2 + NO → ISOPDNO3	KR02NO*alpha_AN(6,1,0,0,0,temp,cair)	Sander et al. (2019)*

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G45122a	TrGC	LISOPEFO2 + HO ₂ → .7143 ISOPDOOH + .2857 ISOPBOOH	k_R02_HO2(temp,5)*(1.-r_CHOHCH2O2_OH)	Sander et al. (2019)
G45122b	TrGC	LISOPEFO2 + HO ₂ → LISOPEFO + OH	k_R02_HO2(temp,5)*r_CHOHCH2O2_OH	Sander et al. (2019)
G45123	TrGCN	LISOPEFO2 + NO ₃ → LISOPEFO + NO ₂	KR02NO3	Sander et al. (2019)
G45124	TrGC	LISOPEFO2 → .7143 MACR + .2857 MVK + HCHO + OH	0.7143*k_hsd+.2857*k_hsb	Sander et al. (2019)
G45125	TrGC	LISOPEFO → .7143 MACR + .2857 MVK + HCHO + HO ₂	KDEC	Sander et al. (2019)
G45126a	TrGC	LISOPACO → 3METHYLFURAN + HO ₂	KDEC*0.37	Sander et al. (2019), Paulot et al. (2009a), Francisco-Marquez et al. (2003)
G45126b	TrGC	LISOPACO → .65 LHC4ACCHO + .65 HO ₂ + .35 DB1O2	KDEC*(1.-0.37)	Sander et al. (2019), Paulot et al. (2009a), Francisco-Marquez et al. (2003)
G45127a	TrGC	LISOPACO → 3METHYLFURAN + HO ₂	KDEC*0.37	Sander et al. (2019), Paulot et al. (2009a), Francisco-Marquez et al. (2003)
G45127b	TrGC	LISOPACO → .65 LHC4ACCHO + .65 HO ₂ + .35 DB1O2	KDEC*(1.-0.37)	Sander et al. (2019), Paulot et al. (2009a), Francisco-Marquez et al. (2003)
G45128	TrGC	3METHYLFURAN + OH → L3METHYLFURANO2	3.2E-11*EXP(310./temp)	Sander et al. (2019)*
G45129	TrGCN	3METHYLFURAN + NO ₃ → L3METHYLFURANO2 + NO ₂	1.9E-11	Sander et al. (2019), Atkinson et al. (2006)*
G45130	TrGC	L3METHYLFURANO2 → C4MDIAL + HO ₂	k1_R02sOR02	Sander et al. (2019)
G45131	TrGCN	L3METHYLFURANO2 + NO → C4MDIAL + HO ₂ + NO ₂	KR02NO	Sander et al. (2019)*
G45132	TrGC	L3METHYLFURANO2 + HO ₂ → C4MDIAL + HO ₂	k_R02_HO2(temp,5)	Sander et al. (2019)*
G45133	TrGC	LZCO3C23DBCOD → .62 EZCH3CO2CHCHO + .38 EZCHOCCH3CHO2 + CO ₂	k1_R02RC03	Sander et al. (2019)
G45134a	TrGC	LZCO3C23DBCOD + HO ₂ → .62 EZCH3CO2CHCHO + .38 EZCHOCCH3CHO2 + CO ₂ + OH	KAPH02*r_CO3_OH	Sander et al. (2019)
G45134b	TrGC	LZCO3C23DBCOD + HO ₂ → LZCO3HC23DBCOD	KAPH02*(r_CO3_00H+r_CO3_03)	Sander et al. (2019)*
G45135	TrGCN	LZCO3C23DBCOD + NO → .62 EZCH3CO2CHCHO + .38 EZCHOCCH3CHO2 + CO ₂ + NO ₂	KAPNO	Sander et al. (2019)
G45136	TrGCN	LZCO3C23DBCOD + NO ₂ → LZCPANC23DBCOD	k_CH3C03_N02	Rickard (2022)

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G45137	TrGCN	LZCO3C23DBCOD + NO ₃ → .62 EZCH3CO2CHCHO + .38 EZCHOCCH3CHO ₂ + CO ₂ + NO ₂	KR02N03*1.74	Sander et al. (2019)
G45138	TrGCN	LZCPANC23DBCOD → LZCO3C23DBCOD + NO ₂	k_PAN_M	Rickard (2022)
G45139	TrGCN	LZCPANC23DBCOD + OH → .62 EZCH3CO2CHCHO + .38 EZCHOCCH3CHO ₂ + CO ₂ + NO ₂	2.52E-11	Sander et al. (2019)*
G45200	TrGTerC	C511O2 → CH ₃ C(O) + HCOCH ₂ CHO	k1_R02sOR02	Rickard (2022)
G45201	TrGTerCN	C511O2 + NO → CH ₃ C(O) + HCOCH ₂ CHO + NO ₂	KR02NO	Rickard (2022)*
G45202a	TrGTerC	C511O2 + HO ₂ → C511OOH	k_R02_H02(temp,5)*r_COCH202_00H	Rickard (2022), Sander et al. (2019)
G45202b	TrGTerC	C511O2 + HO ₂ → CH ₃ C(O) + HCOCH ₂ CHO + OH	k_R02_H02(temp,5)*r_COCH202_OH	Rickard (2022), Sander et al. (2019)
G45203	TrGTerC	C511OOH + OH → C511O2	7.49E-11	Rickard (2022)
G45204	TrGTerC	CO23C4CHO + OH → CO23C4CO3	6.65E-11	Rickard (2022)
G45205	TrGTerCN	CO23C4CHO + NO ₃ → CO23C4CO3 + HNO ₃	KN03AL*5.5	Rickard (2022)
G45206	TrGTerC	CO23C4CO3 → CH ₃ COCOCH ₂ O ₂ + CO ₂	k1_R02RC03	Rickard (2022)
G45207	TrGTerCN	CO23C4CO3 + NO → CH ₃ COCOCH ₂ O ₂ + CO ₂ + NO ₂	KAPNO	Rickard (2022)*
G45208	TrGTerCN	CO23C4CO3 + NO ₂ → C5PAN9	k_CH3C03_N02	Rickard (2022)
G45209a	TrGTerC	CO23C4CO3 + HO ₂ → CO23C4CO3H	KAPH02*(r_C03_00H+r_C03_03)	Rickard (2022)
G45209b	TrGTerC	CO23C4CO3 + HO ₂ → CH ₃ COCOCH ₂ O ₂ + CO ₂ + OH	KAPH02*r_C03_OH	Rickard (2022)
G45210	TrGTerCN	C5PAN9 → CO23C4CO3 + NO ₂	k_PAN_M	Rickard (2022)
G45211	TrGTerCN	C5PAN9 + OH → CH ₃ COCOCHO + CO + NO ₂	3.12E-13	Rickard (2022)
G45212	TrGTerC	C512O2 → C513O2	k1_R02pR02	Rickard (2022)
G45213	TrGTerC	C512O2 + HO ₂ → C512OOH	k_R02_H02(temp,5)	Rickard (2022)
G45214	TrGTerCN	C512O2 + NO → C513O2 + NO ₂	KR02NO	Rickard (2022)*
G45215	TrGTerC	C512OOH + OH → CO13C4CHO + OH	1.01E-10	Rickard (2022)
G45216	TrGTerC	C513O2 → GLYOX + HOC ₂ H ₄ CO ₃	k1_R02sOR02	Rickard (2022)
G45217	TrGTerCN	C513O2 + NO → GLYOX + HOC ₂ H ₄ CO ₃ + NO ₂	KR02NO	Rickard (2022)*
G45218a	TrGTerC	C513O2 + HO ₂ → C513OOH	k_R02_H02(temp,5)*r_COCH202_00H	Rickard (2022), Sander et al. (2019)
G45218b	TrGTerC	C513O2 + HO ₂ → GLYOX + HOC ₂ H ₄ CO ₃ + OH	k_R02_H02(temp,5)*r_COCH202_OH	Rickard (2022), Sander et al. (2019)
G45219	TrGTerC	CO13C4CHO + OH → CHOC3COCO3	1.33E-10	Rickard (2022)
G45220	TrGTerCN	CO13C4CHO + NO ₃ → CHOC3COCO3 + HNO ₃	2.*KN03AL*5.5	Rickard (2022)
G45221	TrGTerC	C513OOH + OH → C513CO + OH	9.23E-11	Rickard (2022)
G45222	TrGTerC	CHOC3COCO3 → CHOC3COO2 + CO ₂	k1_R02RC03	Rickard (2022)

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G45223	TrGTerC	$\text{CHOC3COCO}_3 + \text{HO}_2 \rightarrow \text{CHOC3COOOH}$	KAPH02	Rickard (2022)
G45224	TrGTerCN	$\text{CHOC3COCO}_3 + \text{NO}_2 \rightarrow \text{CHOC3COPAN}$	k_CH3C03_N02	Rickard (2022)
G45225	TrGTerCN	$\text{CHOC3COCO}_3 + \text{NO} \rightarrow \text{CHOC3COO}_2 + \text{CO}_2 + \text{NO}_2$	KAPNO	Rickard (2022)*
G45226	TrGTerC	$\text{C513CO} + \text{OH} \rightarrow \text{HOC}_2\text{H}_4\text{CO}_3 + \text{CO} + \text{CO}$	2.64E-11	Rickard (2022)
G45227	TrGTerC	$\text{C514O}_2 + \text{HO}_2 \rightarrow \text{C514OOH}$	k_RO2_H02(temp,5)	Rickard (2022)
G45228a	TrGTerCN	$\text{C514O}_2 + \text{NO} \rightarrow \text{CO13C4CHO} + \text{HO}_2 + \text{NO}_2$	KRO2NO*(1.-alpha_AN(7,2,0,1,0, temp,cair))	Rickard (2022), Sander et al. (2019)
G45228b	TrGTerCN	$\text{C514O}_2 + \text{NO} \rightarrow \text{C514NO}_3$	KRO2NO*alpha_AN(7,2,0,1,0,temp, cair)	Rickard (2022), Sander et al. (2019)
G45229	TrGTerCN	$\text{C514O}_2 + \text{NO}_3 \rightarrow \text{CO13C4CHO} + \text{HO}_2 + \text{NO}_2$	KRO2N03	Rickard (2022)
G45230	TrGTerC	$\text{C514O}_2 \rightarrow \text{CO13C4CHO} + \text{HO}_2$	k1_RO2sR02	Rickard (2022)
G45231	TrGTerC	$\text{C514OOH} + \text{OH} \rightarrow \text{CO13C4CHO} + \text{OH}$	1.10E-10	Rickard (2022)
G45232	TrGTerCN	$\text{C514NO}_3 + \text{OH} \rightarrow \text{CO13C4CHO} + \text{NO}_2$	4.33E-11	Rickard (2022)
G45233	TrGTerC	$\text{CHOC3COOOH} + \text{OH} \rightarrow \text{CHOC3COCO}_3$	7.55E-11	Rickard (2022)
G45234	TrGTerCN	$\text{CHOC3COPAN} \rightarrow \text{CHOC3COCO}_3 + \text{NO}_2$	k_PAN_M	Rickard (2022)
G45235	TrGTerCN	$\text{CHOC3COPAN} + \text{OH} \rightarrow \text{C4CODIAL} + \text{CO} + \text{NO}_2$	7.19E-11	Rickard (2022)
G45236	TrGTerC	$\text{MBO} + \text{OH} \rightarrow \text{LMBOABO}_2$	8.1E-12*EXP(610./temp)	Rickard (2022), Sander et al. (2019)*
G45237a	TrGTerC	$\text{MBO} + \text{O}_3 \rightarrow \text{HCHO} + .16 \text{CH}_3\text{COCH}_3 + .16 \text{HO}_2 + .16 \text{CO} + .16 \text{OH} + .84 \text{MBOOO}$	1.0E-17*0.57	Rickard (2022), Sander et al. (2019)
G45237b	TrGTerC	$\text{MBO} + \text{O}_3 \rightarrow \text{IBUTALOH} + .63 \text{CO} + .37 \text{HOCH}_2\text{OOH} + .16 \text{OH} + .16 \text{HO}_2$	1.0E-17*0.43	Rickard (2022), Sander et al. (2019)
G45238	TrGTerCN	$\text{MBO} + \text{NO}_3 \rightarrow \text{LNMBABO}_2$	4.6E-14*EXP(-400./temp)	Rickard (2022), Sander et al. (2019)
G45239	TrGTerC	$\text{LMBOABO}_2 + \text{HO}_2 \rightarrow \text{LMBOABOOH}$	k_RO2_H02(temp,5)	Rickard (2022), Sander et al. (2019)
G45240a	TrGTerCN	$\text{LMBOABO}_2 + \text{NO} \rightarrow \text{LMBOABNO}_3$	KRO2NO*(.67*alpha_AN(7,2,0,0,0, temp,cair)+.33*alpha_AN(7,1,0,0, 0,temp,cair))	Rickard (2022), Sander et al. (2019)
G45240b	TrGTerCN	$\text{LMBOABO}_2 + \text{NO} \rightarrow \text{HOCH}_2\text{CHO} + \text{CH}_3\text{COCH}_3 + \text{HO}_2 + \text{NO}_2$	KRO2NO*(1.-(.67*alpha_AN(7,2,0, 0,0,temp,cair)+.33*alpha_AN(7,1, 0,0,0,temp,cair)))*.67	Rickard (2022), Sander et al. (2019)
G45240c	TrGTerCN	$\text{LMBOABO}_2 + \text{NO} \rightarrow \text{IBUTALOH} + \text{HCHO} + \text{HO}_2 + \text{NO}_2$	KRO2NO*(1.-(.67*alpha_AN(7,2,0, 0,0,temp,cair)+.33*alpha_AN(7,1, 0,0,0,temp,cair)))*.33	Rickard (2022), Sander et al. (2019)

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G45241a	TrGTerC	$\text{LMBOABO}_2 \rightarrow \text{HOCH}_2\text{CHO} + \text{CH}_3\text{COCH}_3 + \text{HO}_2$	$k_{1_R02s0R02*}.67$	Rickard (2022), Sander et al. (2019)
G45241b	TrGTerC	$\text{LMBOABO}_2 \rightarrow \text{IBUTALOH} + \text{HCHO} + \text{HO}_2$	$k_{1_R02p0R02*}.33$	Rickard (2022), Sander et al. (2019)
G45242a	TrGTerC	$\text{LMBOABOOH} + \text{OH} \rightarrow \text{MBOACO}$	$0.67*2.93\text{E}-11+.33*2.05\text{E}-12$	Rickard (2022), Sander et al. (2019)
G45242b	TrGTerC	$\text{LMBOABOOH} + \text{OH} \rightarrow \text{LMBOABO}_2$	$k_{_ROOHRO}$	Rickard (2022), Sander et al. (2019)
G45243	TrGTerCN	$\text{LMBOABNO}_3 + \text{OH} \rightarrow \text{MBOACO} + \text{NO}_2$	$0.67*1.75\text{E}-12+.33*2.69\text{E}-12$	Rickard (2022), Sander et al. (2019)
G45244	TrGTerC	$\text{MBOACO} + \text{OH} \rightarrow \text{MBOCOCO} + \text{HO}_2$	$3.79\text{E}-12$	Rickard (2022)
G45245	TrGTerC	$\text{MBOCOCO} + \text{OH} \rightarrow \text{CO} + \text{IPRHOCO}_3$	$1.38\text{E}-11$	Rickard (2022)
G45246	TrGTerCN	$\text{LNMBABO}_2 + \text{HO}_2 \rightarrow \text{LNMBABOOH}$	$k_{_R02_H02(temp,5)}$	Rickard (2022), Sander et al. (2019)
G45247	TrGTerCN	$\text{LNMBABO}_2 + \text{NO} \rightarrow .65 \text{NO}_3\text{CH}_2\text{CHO} + .65 \text{CH}_3\text{COCH}_3 + .65 \text{HO}_2 + .35 \text{IBUTALOH} + .35 \text{HCHO} + .35 \text{NO}_2 + \text{NO}_2$	K_{R02NO}	Rickard (2022), Sander et al. (2019)*
G45248	TrGTerCN	$\text{LNMBABO}_2 + \text{NO}_3 \rightarrow .65 \text{NO}_3\text{CH}_2\text{CHO} + .65 \text{CH}_3\text{COCH}_3 + .65 \text{HO}_2 + .35 \text{IBUTALOH} + .35 \text{HCHO} + .35 \text{NO}_2 + \text{NO}_2$	K_{R02N03}	Rickard (2022), Sander et al. (2019)
G45249	TrGTerCN	$\text{LNMBABO}_2 \rightarrow .65 \text{NO}_3\text{CH}_2\text{CHO} + .65 \text{CH}_3\text{COCH}_3 + .65 \text{HO}_2 + .35 \text{IBUTALOH} + .35 \text{HCHO} + .35 \text{NO}_2$	$k_{1_R02s0R02}$	Rickard (2022), Sander et al. (2019)
G45250a	TrGTerCN	$\text{LNMBABOOH} + \text{OH} \rightarrow .65 \text{C4MCONO}_3\text{OH} + .35 \text{NMBOBCO}$	$0.65*4.89\text{E}-12+.35*2.52\text{E}-12$	Rickard (2022), Sander et al. (2019)
G45250b	TrGTerCN	$\text{LNMBABOOH} + \text{OH} \rightarrow \text{LNMBABO}_2$	$k_{_ROOHRO}$	Rickard (2022), Sander et al. (2019)
G45251	TrGTerCN	$\text{NMBOBCO} + \text{OH} \rightarrow \text{NC4OHCO}_3$	$4.26\text{E}-12$	Rickard (2022)
G45252a	TrGTerCN	$\text{NC4OHCO}_3 + \text{HO}_2 \rightarrow \text{IBUTALOH} + \text{CO}_2 + \text{NO}_2 + \text{OH}$	$K_{AHO2*r_C03_OH}$	Rickard (2022), Sander et al. (2019)
G45252b	TrGTerCN	$\text{NC4OHCO}_3 + \text{HO}_2 \rightarrow \text{NC4OHCO}_3\text{H}$	$K_{AHO2*(r_C03_03+r_C03_00H)}$	Rickard (2022), Sander et al. (2019)
G45253	TrGTerCN	$\text{NC4OHCO}_3 + \text{NO} \rightarrow \text{IBUTALOH} + \text{CO}_2 + \text{NO}_2 + \text{NO}_2$	K_{APNO}	Rickard (2022)
G45254	TrGTerCN	$\text{NC4OHCO}_3 + \text{NO}_2 \rightarrow \text{NC4OHCPAN}$	$k_{_CH3C03_N02}$	Rickard (2022)
G45255	TrGTerCN	$\text{NC4OHCO}_3 + \text{NO}_3 \rightarrow \text{IBUTALOH} + \text{CO}_2 + \text{NO}_2 + \text{NO}_2$	$K_{R02N03*1.74}$	Rickard (2022)
G45256	TrGTerCN	$\text{NC4OHCO}_3 \rightarrow \text{IBUTALOH} + \text{CO}_2 + \text{NO}_2$	$k_{1_R02RC03}$	Rickard (2022)

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G45257	TrGTerCN	NC4OHCO3H + OH → NC4OHCO3	4.50E-12	Rickard (2022)
G45258	TrGTerCN	NC4OHCPAN + OH → IBUTALOH + CO + NO ₂ + NO ₂	1.27E-12	Rickard (2022)
G45259	TrGTerCN	NC4OHCPAN → NC4OHCO3 + NO ₂	k_PAN_M	Rickard (2022)
G45260	TrGTerCN	C4MCONO3OH + OH → CH ₃ COCH ₃ + HCHO + CO ₂ + NO ₂	1.23E-12	Rickard (2022), Sander et al. (2019)
G45400	TrGAroCN	NC4MDCO2HN + OH → MMALANHY + NO ₂	k_ROOHRO	Rickard (2022)*
G45401	TrGAroCN	C54CO + NO ₃ → 3 CO + CH ₃ C(O)OO + HNO ₃	KNO3AL*5.5	Rickard (2022)
G45402	TrGAroC	C54CO + OH → 3 CO + CH ₃ C(O)OO	1.72E-11	Rickard (2022)
G45403a	TrGAroCN	NTLFUO2 + HO ₂ → NTLFUOOH	k_R02_H02(temp,5)*(1.-r_COCH202_OH)	Rickard (2022)
G45403b	TrGAroCN	NTLFUO2 + HO ₂ → ACCOMECHO + NO ₂ + OH	k_R02_H02(temp,5)*r_COCH202_OH	Rickard (2022)
G45404	TrGAroCN	NTLFUO2 + NO → ACCOMECHO + NO ₂ + NO ₂	KR02NO	Rickard (2022)*
G45405	TrGAroCN	NTLFUO2 + NO ₃ → ACCOMECHO + NO ₂ + NO ₂	KR02NO3	Rickard (2022)*
G45406	TrGAroCN	NTLFUO2 → ACCOMECHO + NO ₂	k1_R02tOR02	Rickard (2022)*
G45407	TrGAroC	C5134CO2OH + OH → C54CO + HO ₂	7.48E-11	Rickard (2022)
G45408	TrGAroCN	C5COO2NO2 + OH → MGLYOX + CO + CO + NO ₂	5.43E-11	Rickard (2022)
G45409	TrGAroCN	C5COO2NO2 → C5CO14O2 + NO ₂	k_PAN_M	Rickard (2022)*
G45410	TrGAroC	C5DIALOOH + OH → C5DIALCO + OH	7.52E-11	Rickard (2022)
G45411a	TrGAroC	C4CO2DBCO3 + HO ₂ → C4CO2DCO3H	KAPHO2*(r_C03_00H+r_C03_03)	Rickard (2022)
G45411b	TrGAroC	C4CO2DBCO3 + HO ₂ → HO ₂ + CO + HCOCOCHO + CO ₂ + OH	KAPHO2*r_C03_OH	Rickard (2022), Sander et al. (2019)
G45412	TrGAroCN	C4CO2DBCO3 + NO → HO ₂ + CO + HCOCOCHO + CO ₂ + NO ₂	KAPNO	Rickard (2022)
G45413	TrGAroCN	C4CO2DBCO3 + NO ₂ → C4CO2DBPAN	k_CH3C03_N02	Rickard (2022)*
G45414	TrGAroCN	C4CO2DBCO3 + NO ₃ → HO ₂ + CO + HCOCOCHO + CO ₂ + NO ₂	KR02NO3*1.74	Rickard (2022)
G45415	TrGAroC	C4CO2DBCO3 → HO ₂ + CO + HCOCOCHO + CO ₂	k1_R02RC03	Rickard (2022)
G45416	TrGAroC	MMALANHY + OH → MMALANHYO2	1.50E-12	Rickard (2022)
G45421a	TrGAroC	MMALANHYO2 + HO ₂ → MMALNHYOOH	k_R02_H02(temp,5)*(1.-r_COCH202_OH-r_CHOHCH202_OH)	Rickard (2022), Sander et al. (2019)
G45421b	TrGAroC	MMALANHYO2 + HO ₂ → CO2H3CO3 + CO ₂ + OH	k_R02_H02(temp,5)*(r_COCH202_OH+r_CHOHCH202_OH)	Rickard (2022), Sander et al. (2019)
G45422	TrGAroCN	MMALANHYO2 + NO → CO2H3CO3 + CO ₂ + NO ₂	KR02NO	Rickard (2022)*
G45423	TrGAroCN	MMALANHYO2 + NO ₃ → CO2H3CO3 + CO ₂ + NO ₂	KR02NO3	Rickard (2022)*
G45424	TrGAroC	MMALANHYO2 → CO2H3CO3 + CO ₂	k1_R02tOR02	Rickard (2022)*

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G45428	TrGAroCN	C4CO2DBPAN + OH → HCOCOCHO + CO ₂ + CO + NO ₂	2.74E-11	Rickard (2022)
G45429	TrGAroCN	C4CO2DBPAN → C4CO2DBCO3 + NO ₂	k_PAN_M	Rickard (2022)*
G45430a	TrGAroC	C5CO14O2 + HO ₂ → .83 MALANHY + .83 CH ₃ + .17 MGLYOX + .17 HO ₂ + .17 CO + .17 CO ₂ + OH	KAPH02*r_CO3_OH	Rickard (2022)*
G45430b	TrGAroC	C5CO14O2 + HO ₂ → C5CO14OH + O ₃	KAPH02*r_CO3_03	Rickard (2022)
G45430c	TrGAroC	C5CO14O2 + HO ₂ → C5CO14OOH	KAPH02*r_CO3_OOH	Rickard (2022)
G45431	TrGAroCN	C5CO14O2 + NO → .83 MALANHY + .83 CH ₃ + .17 MGLYOX + .17 HO ₂ + .17 CO + .17 CO ₂ + NO ₂	KAPNO	Rickard (2022)*
G45432	TrGAroCN	C5CO14O2 + NO ₂ → C5COO2NO ₂	k_CH3CO3_NO2	Rickard (2022)*
G45433	TrGAroCN	C5CO14O2 + NO ₃ → .83 MALANHY + .83 CH ₃ + .17 MGLYOX + .17 HO ₂ + .17 CO + .17 CO ₂ + NO ₂	KR02NO3*1.74	Rickard (2022)*
G45434	TrGAroC	C5CO14O2 → .83 MALANHY + .83 CH ₃ + .17 MGLYOX + .17 HO ₂ + .17 CO + .17 CO ₂	k1_R02RC03	Rickard (2022)*
G45436	TrGAroC	C5CO14OH + OH → .83 MALANHY + .83 CH ₃ + .17 MGLYOX + .17 HO ₂ + .17 CO + .17 CO ₂	5.44E-11	Rickard (2022)*
G45441	TrGAroCN	C5DICARB + NO ₃ → C5CO14O2 + HNO ₃	KN03AL*2.75	Rickard (2022)
G45442	TrGAroC	C5DICARB + O ₃ → .5338 GLYOX + .063 CH ₃ CHO + .348 CH ₃ C(O)OO + .918 CO + .57 OH + .473 HO ₂ + .0563 CH ₃ COCO ₂ H + .5338 MGLYOX + .676 H ₂ O ₂ + .063 HCHO + .0563 HCOCO ₂ H + .2465 CO ₂	2.00E-18	Rickard (2022)
G45443	TrGAroC	C5DICARB + OH → .48 C5CO14O2 + .52 C5DICARBO2	6.2E-11	Rickard (2022)
G45444	TrGAroC	MC3ODBCO2H + OH → .35 GLYOX + .35 CH ₃ + .35 CO + .35 CO ₂ + .65 MMALANHY + .65 HO ₂	4.38E-11	Rickard (2022)*
G45451	TrGAroCN	TLFUONE + NO ₃ → NTLFUO2	1.00E-12	Rickard (2022)
G45452	TrGAroC	TLFUONE + O ₃ → .5 CO + .5 OH + .5 MECOACETO2 + .3125 C24O3CCO2H + .1875 ACCOMECHO + .1875 H ₂ O ₂	8.00E-19	see note*
G45453	TrGAroC	TLFUONE + OH → TLFUO2	6.90E-11	Rickard (2022)
G45454a	TrGAroC	ACCOMEKO3 + HO ₂ → ACCOMEKO3H	KAPH02*(r_CO3_OOH+r_CO3_03)	Rickard (2022)
G45454b	TrGAroC	ACCOMEKO3 + HO ₂ → MECOACETO2 + CO ₂ + OH	KAPH02*r_CO3_OH	Rickard (2022)
G45455	TrGAroCN	ACCOMEKO3 + NO → MECOACETO2 + CO ₂ + NO ₂	KAPNO	Rickard (2022)
G45456	TrGAroCN	ACCOMEKO3 + NO ₂ → ACCOME PAN	k_CH3CO3_NO2	Rickard (2022)*
G45457	TrGAroCN	ACCOMEKO3 + NO ₃ → MECOACETO2 + CO ₂ + NO ₂	KR02NO3*1.74	Rickard (2022)
G45458	TrGAroC	ACCOMEKO3 → MECOACETO2 + CO ₂	k1_R02RC03	Rickard (2022)

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G45459	TrGAroC	C4CO2DCO3H + OH → C4CO2DBCO3	3.06E-11	Rickard (2022)
G45464	TrGAroCN	ACCOMECHO + NO ₃ → ACCOMEKO3 + HNO ₃	KNO3AL*5.5	Rickard (2022)
G45465	TrGAroC	ACCOMECHO + OH → ACCOMEKO3	7.09E-11	Rickard (2022)
G45466	TrGAroC	MMALNHYOOH + OH → MMALANHYO2	1.69E-11	Rickard (2022)
G45467a	TrGAroC	C5DICAROOH + OH → C5134CO2OH + OH	1.21E-10	Rickard (2022)
G45467b	TrGAroC	C5DICAROOH + OH → C5DICARBO2	k_ROOHR0	Rickard (2022)
G45468	TrGAroC	C24O3CCO2H + OH → MECOACETO2 + CO ₂	8.76E-13	Rickard (2022)
G45469	TrGAroCN	NTLFUOOH + OH → NTLFUO2	4.44E-12	Rickard (2022)
G45470	TrGAroCN	ACCOMEPAH + OH → METACETHO + CO + CO + NO ₂	1.00E-14	Rickard (2022)
G45471	TrGAroCN	ACCOMEPAH → ACCOMEKO3 + NO ₂	k_PAN_M	Rickard (2022)
G45476a	TrGAroC	TLFUO2 + HO ₂ → TLFUOOH	k_RO2_HO2(temp,5)*(1.-r_COCH202_OH-r_CHOHCH202_OH)	Rickard (2022)
G45476b	TrGAroC	TLFUO2 + HO ₂ → ACCOMECHO + HO ₂ + OH	k_RO2_HO2(temp,5)*(r_COCH202_OH+r_CHOHCH202_OH)	Rickard (2022)*
G45477	TrGAroCN	TLFUO2 + NO → ACCOMECHO + HO ₂ + NO ₂	KRO2NO	Rickard (2022)*
G45478	TrGAroCN	TLFUO2 + NO ₃ → ACCOMECHO + HO ₂ + NO ₂	KRO2NO3	Rickard (2022)*
G45479	TrGAroC	TLFUO2 → ACCOMECHO + HO ₂	k1_R02t0R02	Rickard (2022)*
G45480	TrGAroC	C5CO14OOH + OH → C5CO14O2	3.59E-12	Rickard (2022)
G45483	TrGAroC	TLFUOOH + OH → TLFUO2	2.53E-11	Rickard (2022)
G45485	TrGAroC	ACCOMEKO3H + OH → ACCOMEKO3	3.59E-12	Rickard (2022)
G45486a	TrGAroC	C5DIALO2 + HO ₂ → C5DIALOOH	k_RO2_HO2(temp,5)*(1.-r_COCH202_OH)	Rickard (2022)
G45486b	TrGAroC	C5DIALO2 + HO ₂ → MALDIAL + CO + HO ₂ + OH	k_RO2_HO2(temp,5)*r_COCH202_OH	Rickard (2022)*
G45487	TrGAroCN	C5DIALO2 + NO → MALDIAL + CO + HO ₂ + NO ₂	KRO2NO	Rickard (2022)*
G45488	TrGAroCN	C5DIALO2 + NO ₃ → MALDIAL + CO + HO ₂ + NO ₂	KRO2NO3	Rickard (2022)*
G45489	TrGAroC	C5DIALO2 → MALDIAL + CO + HO ₂	k1_R02s0R02	Rickard (2022)*
G45490a	TrGAroC	C5DICARBO2 + HO ₂ → C5DICAROOH	k_RO2_HO2(temp,5)*(r_CO3_00H+r_CO3_03)	Rickard (2022)
G45491b	TrGAroC	C5DICARBO2 + HO ₂ → MGLYOX + GLYOX + HO ₂ + OH	k_RO2_HO2(temp,5)*r_CO3_OH	Rickard (2022)*
G45492	TrGAroCN	C5DICARBO2 + NO → MGLYOX + GLYOX + HO ₂ + NO ₂	KRO2NO	Rickard (2022)*
G45493	TrGAroCN	C5DICARBO2 + NO ₃ → MGLYOX + GLYOX + HO ₂ + NO ₂	KRO2NO3	Rickard (2022)*

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G45494	TrGAroC	C5DICARBO2 → MGLYOX + GLYOX + HO ₂	k1_R02sOR02	Rickard (2022)*
G46200a	TrGTerC	CO235C6O2 + HO ₂ → CO235C6OOH	k_R02_H02(temp,6)*r_COCH202_OOH	Rickard (2022), Sander et al. (2019)
G46200b	TrGTerC	CO235C6O2 + HO ₂ → CO23C4CO3 + HCHO + OH	k_R02_H02(temp,6)*r_COCH202_OH	Rickard (2022), Sander et al. (2019)
G46201	TrGTerCN	CO235C6O2 + NO → CO23C4CO3 + HCHO + NO ₂	KR02NO	Rickard (2022)*
G46202	TrGTerC	CO235C6O2 → CO23C4CO3 + HCHO	k1_R02pOR02	Rickard (2022)
G46203	TrGTerC	CO235C6OOH + OH → CO235C6O2	1.01E-11	Rickard (2022)
G46204	TrGTerC	C614O2 → CO23C4CHO + HCHO + HO ₂	k1_R02sOR02	Rickard (2022)
G46205a	TrGTerCN	C614O2 + NO → CO23C4CHO + HCHO + HO ₂ + NO ₂	KR02NO*(1.-alpha_AN(9,2,0,1,0, temp, cair))	Rickard (2022)
G46205b	TrGTerCN	C614O2 + NO → C614NO3	KR02NO*alpha_AN(9,2,0,1,0, temp, cair)	Rickard (2022)
G46206a	TrGTerC	C614O2 + HO ₂ → C614OOH	k_R02_H02(temp,6)*(1.-r_CHOCH202_OH)	Rickard (2022), Sander et al. (2019)
G46206b	TrGTerC	C614O2 + HO ₂ → CO23C4CHO + HCHO + HO ₂ + OH	k_R02_H02(temp,6)*r_CHOCH202_OH	Rickard (2022), Sander et al. (2019)
G46207	TrGTerCN	C614NO3 + OH → C614CO + NO ₂	7.11E-12	Rickard (2022)
G46208	TrGTerC	C614OOH + OH → C614CO + OH	8.69E-11	Rickard (2022)
G46209	TrGTerC	C614CO + OH → CO235C5CHO + HO ₂	3.22E-12	Rickard (2022)
G46210	TrGTerC	CO235C5CHO + OH → CO23C4CO3 + CO	1.33E-11	Rickard (2022)
G46211	TrGTerCN	CO235C5CHO + NO ₃ → CO23C4CO3 + CO + HNO ₃	KN03AL*5.5	Rickard (2022)
G46400	TrGAroC	PHENOOH + OH → PHENO2	1.16E-10	Rickard (2022)
G46401	TrGAroC	C6CO4DB + OH → CO + CO + HO ₂ + CO + HCOCOCHO	7.70E-11	Rickard (2022)
G46402	TrGAroC	C5CO2DCO3H + OH → C5CO2DBC03	3.60E-11	Rickard (2022)
G46403	TrGAroCN	NDNPHENOOH + OH → NDNPHENO2	k_ROOHRO	Rickard (2022)
G46404a	TrGAroC	C615CO2O2 + HO ₂ → C615CO2OOH	k_R02_H02(temp,6)*(1.-r_COCH202_OH)	Rickard (2022)
G46404b	TrGAroC	C615CO2O2 + HO ₂ → C5DICARB + CO + HO ₂ + OH	k_R02_H02(temp,6)*r_COCH202_OH	Rickard (2022)*
G46405	TrGAroCN	C615CO2O2 + NO → C5DICARB + CO + HO ₂ + NO ₂	KR02NO	Rickard (2022)*
G46406	TrGAroCN	C615CO2O2 + NO ₃ → C5DICARB + CO + HO ₂ + NO ₂	KR02NO3	Rickard (2022)*
G46407	TrGAroC	C615CO2O2 → C5DICARB + CO + HO ₂	k1_R02sOR02	Rickard (2022)*
G46408	TrGAroCN	BZEMUCPAN + OH → MALDIAL + CO + CO ₂ + NO ₂	4.05E-11	Rickard (2022)
G46409	TrGAroCN	BZEMUCPAN → BZEMUCCO3 + NO ₂	k_PAN_M	Rickard (2022)

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G46410	TrGAroCN	BZBIPERNO3 + OH → BZOBIPEROH + NO ₂	7.30E-11	Rickard (2022)
G46411	TrGAroCN	HOC6H4NO ₂ + NO ₃ → NPHEN1O + HNO ₃	9.00E-14	Rickard (2022)
G46412	TrGAroCN	HOC6H4NO ₂ + OH → NPHEN1O	9.00E-13	Rickard (2022)
G46413a	TrGAroCN	NDNPHENO ₂ + HO ₂ → NDNPHENOOH	k_R02_H02(temp,6)*(1.-r_ CHOHCH2O2_OH)	Rickard (2022)
G46413b	TrGAroCN	NDNPHENO ₂ + HO ₂ → NC4DCO2H + HNO ₃ + CO + CO + NO ₂ + OH	k_R02_H02(temp,6)*r_CHOHCH2O2_OH	Rickard (2022)*
G46414	TrGAroCN	NDNPHENO ₂ + NO → NC4DCO2H + HNO ₃ + CO + CO + NO ₂ + NO ₂	KR02NO	Rickard (2022)*
G46415	TrGAroCN	NDNPHENO ₂ + NO ₃ → NC4DCO2H + HNO ₃ + CO + CO + NO ₂ + NO ₂	KR02N03	Rickard (2022)*
G46416	TrGAroCN	NDNPHENO ₂ → NC4DCO2H + HNO ₃ + CO + CO + NO ₂	k1_R02ISOPD02	Rickard (2022)*
G46417	TrGAroC	PBZQCO + OH → C5CO2OHCO3	6.07E-11	Rickard (2022)
G46418	TrGAroCN	CATECHOL + NO ₃ → CATEC1O + HNO ₃	9.9E-11	Rickard (2022)*
G46419	TrGAroC	CATECHOL + O ₃ → MALDALCO2H + HCOCO ₂ H + HO ₂ + OH	9.2E-18	Rickard (2022)
G46420	TrGAroC	CATECHOL + OH → CATEC1O	1.0E-10	Rickard (2022)
G46421	TrGAroC	C5COOHCO3H + OH → C5CO2OHCO3	8.01E-11	Rickard (2022)
G46422	TrGAroCN	NCATECHOL + NO ₃ → NNCATECO2	2.60E-12	Rickard (2022)
G46423	TrGAroCN	NCATECHOL + OH → NCATECO2	3.47E-12	Rickard (2022)
G46424a	TrGAroC	C5CO2OHCO3 + HO ₂ → C5COOHCO3H	KAPH02*(r_C03_00H+r_C03_03)	Rickard (2022)
G46424b	TrGAroC	C5CO2OHCO3 + HO ₂ → HOCOC4DIAL + HO ₂ + CO + CO ₂ + OH	KAPH02*r_C03_OH	Rickard (2022)
G46425	TrGAroCN	C5CO2OHCO3 + NO → HOCOC4DIAL + HO ₂ + CO + CO ₂ + NO ₂	KAPNO	Rickard (2022)
G46426	TrGAroCN	C5CO2OHCO3 + NO ₂ → C5CO2OHPAN	k_CH3C03_N02	Rickard (2022)*
G46427	TrGAroCN	C5CO2OHCO3 + NO ₃ → HOCOC4DIAL + HO ₂ + CO + CO ₂ + NO ₂	KR02N03*1.74	Rickard (2022)
G46428	TrGAroC	C5CO2OHCO3 → HOCOC4DIAL + HO ₂ + CO + CO ₂	k1_R02RC03	Rickard (2022)
G46429	TrGAroCN	BZEPOXMUC + NO ₃ → BZEMUCCO3 + HNO ₃	2.*KN03AL*2.75	Rickard (2022)
G46430	TrGAroC	BZEPOXMUC + O ₃ → EPXC4DIAL + .125 HCHO + .1125 HCOCO ₂ H + .0675 GLYOX + .0675 H ₂ O ₂ + .82 HO ₂ + .57 OH + 1.265 CO + .25 CO ₂	2.00E-18	Rickard (2022)*

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G46431	TrGAroC	BZEPOXMUC + OH → .31 BZEMUCCO3 + .69 BZEMUCO2	6.08E-11	Rickard (2022)
G46432a	TrGAroCN	NCATECO2 + HO2 → NCATECOOH	k_RO2_HO2(temp,6)*(1.-r_CHOHCH2O2_OH)	Rickard (2022)
G46432b	TrGAroCN	NCATECO2 + HO2 → NC4DCO2H + HCOCO2H + HO2 + OH	k_RO2_HO2(temp,6)*r_CHOHCH2O2_OH	Rickard (2022)*
G46433	TrGAroCN	NCATECO2 + NO → NC4DCO2H + HCOCO2H + HO2 + NO2	KRO2NO	Rickard (2022)*
G46434	TrGAroCN	NCATECO2 + NO3 → NC4DCO2H + HCOCO2H + HO2 + NO2	KRO2NO3	Rickard (2022)*
G46435	TrGAroCN	NCATECO2 → NC4DCO2H + HCOCO2H + HO2	k1_RO2ISOPD02	Rickard (2022)*
G46436	TrGAroCN	NPHEN1OOH + OH → NPHEN1O2	9.00E-13	Rickard (2022)
G46437a	TrGAroCN	NPHENO2 + HO2 → NPHENOOH	k_RO2_HO2(temp,6)*(1.-r_CHOHCH2O2_OH)	Rickard (2022)
G46437b	TrGAroCN	NPHENO2 + HO2 → MALDALCO2H + GLYOX + NO2 + OH	k_RO2_HO2(temp,6)*r_CHOHCH2O2_OH	Rickard (2022)*
G46438	TrGAroCN	NPHENO2 + NO → MALDALCO2H + GLYOX + NO2 + NO2	KRO2NO	Rickard (2022)*
G46439	TrGAroCN	NPHENO2 + NO3 → MALDALCO2H + GLYOX + NO2 + NO2	KRO2NO3	Rickard (2022)*
G46440	TrGAroCN	NPHENO2 → MALDALCO2H + GLYOX + NO2	k1_RO2ISOPD02	Rickard (2022)*
G46441	TrGAroC	BENZENE + OH → .352 BZBIPERO2 + .118 BZEPOXMUC + .118 HO2 + .53 PHENOL + .53 HO2	2.3E-12*EXP(-190./temp)	Rickard (2022)*
G46442	TrGAroCN	C5CO2OHPAN + OH → HOCOC4DIAL + CO + CO + NO2	7.66E-11	Rickard (2022)
G46443	TrGAroCN	C5CO2OHPAN → C5CO2OHCO3 + NO2	k_PAN_M	Rickard (2022)
G46444	TrGAroCN	CATEC1O + NO2 → NCATECHOL	k_C6H5O_NO2	Rickard (2022), Platz et al. (1998)
G46445	TrGAroC	CATEC1O + O3 → CATEC1O2	k_C6H5O_03	Rickard (2022), Tao and Li (1999)
G46446	TrGAroC	BZEMUCCO + OH → EPXDLCO3 + GLYOX	9.20E-11	Rickard (2022)
G46447a	TrGAroCN	NNCATECO2 + HO2 → NNCATECOOH	k_RO2_HO2(temp,6)*(1.-r_CHOHCH2O2_OH)	Rickard (2022)
G46447b	TrGAroCN	NNCATECO2 + HO2 → NC4DCO2H + HCOCO2H + NO2 + OH	k_RO2_HO2(temp,6)*r_CHOHCH2O2_OH	Rickard (2022)*

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G46448	TrGAroCN	NNCATECO2 + NO → NC4DCO2H + HCOCO2H + NO2 + NO2	KR02NO	Rickard (2022)*
G46449	TrGAroCN	NNCATECO2 + NO3 → NC4DCO2H + HCOCO2H + NO2 + NO2	KR02NO3	Rickard (2022)*
G46450	TrGAroCN	NNCATECO2 → NC4DCO2H + HCOCO2H + NO2	k1_RO2ISOPD02	Rickard (2022)*
G46451	TrGAroC	BZEMUCCO2H + OH → C5DIALO2 + CO2	4.06E-11	Rickard (2022)
G46452	TrGAroCN	NNCATECOOH + OH → NNCATECO2	k_ROOHRO	Rickard (2022)
G46453	TrGAroCN	NPHEN1O + NO2 → DNPHEN	k_C6H5O_NO2	Rickard (2022), Platz et al. (1998)
G46454	TrGAroCN	NPHEN1O + O3 → NPHEN1O2	k_C6H5O_03	Rickard (2022), Tao and Li (1999)
G46455	TrGAroCN	DNPHEN + NO3 → NDNPHENO2	2.25E-15	Rickard (2022)
G46456	TrGAroCN	DNPHEN + OH → DNPHENO2	3.00E-14	Rickard (2022)
G46457	TrGAroCN	PHENOL + NO3 → .742 C6H5O + .742 HNO3 + .258 NPHENO2	3.8E-12	Rickard (2022)*
G46458	TrGAroC	PHENOL + OH → .06 C6H5O + .8 CATECHOL + .8 HO2 + .14 PHENO2	4.7E-13*EXP(1220./temp)	Rickard (2022)*
G46459	TrGAroCN	PBZQONE + NO3 → NBZQO2	3.00E-13	Rickard (2022)
G46460	TrGAroC	PBZQONE + OH → PBZQO2	4.6E-12	Rickard (2022)
G46461a	TrGAroC	PHENO2 + HO2 → PHENOOH	k_RO2_HO2(temp,6)*(1.-r_CHOHCH2O2_OH)	Rickard (2022)
G46461b	TrGAroC	PHENO2 + HO2 → .71 MALDALCO2H + .71 GLYOX + .29 PBZQONE + HO2 + OH	k_RO2_HO2(temp,6)*r_CHOHCH2O2_OH	Rickard (2022)*
G46462	TrGAroCN	PHENO2 + NO → .71 MALDALCO2H + .71 GLYOX + .29 PBZQONE + HO2 + NO2	KR02NO	Rickard (2022)*
G46463	TrGAroCN	PHENO2 + NO3 → .71 MALDALCO2H + .71 GLYOX + .29 PBZQONE + HO2 + NO2	KR02NO3	Rickard (2022)*
G46464	TrGAroC	PHENO2 → .71 MALDALCO2H + .71 GLYOX + .29 PBZQONE + HO2	k1_RO2ISOPD02	Rickard (2022)*
G46465	TrGAroC	C615CO2OOH + OH → C6125CO + OH	9.42E-11	Rickard (2022)
G46466a	TrGAroC	C5CO2DBCO3 + HO2 → C5CO2DCO3H	KAPH02*(r_CO3_00H+r_CO3_03)	Rickard (2022)
G46466b	TrGAroC	C5CO2DBCO3 + HO2 → CH3C(O) + HCOCOCHO + CO2 + OH	KAPH02*r_CO3_OH	Rickard (2022)
G46467	TrGAroCN	C5CO2DBCO3 + NO → CH3C(O) + HCOCOCHO + CO2 + NO2	KAPNO	Rickard (2022)

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G46468	TrGAroCN	C5CO2DBCO3 + NO ₂ → C5CO2DBPAN	k_CH3C03_N02	Rickard (2022)*
G46469	TrGAroCN	C5CO2DBCO3 + NO ₃ → CH ₃ C(O) + HCOCOCHO + CO ₂ + NO ₂	KR02N03*1.74	Rickard (2022)
G46470	TrGAroC	C5CO2DBCO3 → CH ₃ C(O) + HCOCOCHO + CO ₂	k1_R02RC03	Rickard (2022)
G46471	TrGAroCN	NPHEN1O2 + HO ₂ → NPHEN1OOH	k_R02_H02(temp,6)	Rickard (2022)
G46472a	TrGAroCN	NPHEN1O2 + NO → NPHEN1O + NO ₂	KR02NO	Rickard (2022)
G46472b	TrGAroCN	NPHEN1O2 + NO ₂ → NPHEN1O + NO ₃	k_C6H502_N02	Jagiella and Zabel (2007)*
G46473	TrGAroCN	NPHEN1O2 + NO ₃ → NPHEN1O + NO ₂	KR02N03	Rickard (2022)
G46474	TrGAroCN	NPHEN1O2 → NPHEN1O	k1_R02sR02	Rickard (2022)
G46475	TrGAroCN	NPHENOOH + OH → NPHENO2	1.07E-10	Rickard (2022)
G46476	TrGAroCN	C6H5O + NO ₂ → HOC6H4NO2	k_C6H50_N02	Rickard (2022), Platz et al. (1998)*
G46477	TrGAroC	C6H5O + O ₃ → C6H5O2	k_C6H50_03	Rickard (2022), Tao and Li (1999)
G46478	TrGAroCN	NCATECOOH + OH → NCATECO2	k_ROOHRO	Rickard (2022)
G46479	TrGAroC	PBZQOOH + OH → PBZQCO + OH	1.23E-10	Rickard (2022)
G46480a	TrGAroC	PBZQO2 + HO ₂ → PBZQOOH	k_R02_H02(temp,6)*(1.-r_CHOCH202_OH-r_COCH202_OH)	Rickard (2022)
G46480b	TrGAroC	PBZQO2 + HO ₂ → C5CO2OHCO3 + OH	k_R02_H02(temp,6)*(r_CHOCH202_OH+r_COCH202_OH)	Rickard (2022)*
G46481	TrGAroCN	PBZQO2 + NO → C5CO2OHCO3 + NO ₂	KR02NO	Rickard (2022)*
G46482	TrGAroCN	PBZQO2 + NO ₃ → C5CO2OHCO3 + NO ₂	KR02N03	Rickard (2022)*
G46483	TrGAroC	PBZQO2 → C5CO2OHCO3	k1_R02sR02	Rickard (2022)*
G46484	TrGAroC	BZOBIPOEROH + OH → MALDIALCO3 + GLYOX	8.16E-11	Rickard (2022)
G46485a	TrGAroCN	DNPHEN02 + HO ₂ → DNPHENOOH	k_R02_H02(temp,6)*(1.-r_CHOCH202_OH)	Rickard (2022)
G46485b	TrGAroCN	DNPHEN02 + HO ₂ → NC4DCO2H + HCOCO ₂ H + NO ₂ + OH	k_R02_H02(temp,6)*r_CHOCH202_OH	Rickard (2022)*
G46486	TrGAroCN	DNPHEN02 + NO → NC4DCO2H + HCOCO ₂ H + NO ₂ + NO ₂	KR02NO	Rickard (2022)*
G46487	TrGAroCN	DNPHEN02 + NO ₃ → NC4DCO2H + HCOCO ₂ H + NO ₂ + NO ₂	KR02N03	Rickard (2022)*
G46488	TrGAroCN	DNPHEN02 → NC4DCO2H + HCOCO ₂ H + NO ₂	k1_R02ISOPD02	Rickard (2022)*
G46489	TrGAroC	BZBIPEROOH + OH → BZOBIPOEROH + OH	9.77E-11	Rickard (2022)
G46490a	TrGAroC	BZEMUCO2 + HO ₂ → BZEMUCOOH	k_R02_H02(temp,6)	Rickard (2022)

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G46490b	TrGAroC	BZEMUCO2 + HO ₂ → .5 EPXC4DIAL + .5 GLYOX + .5 HO ₂ + .5 C3DIALO2 + .5 C32OH13CO + OH	k_RO2_HO2(temp,6)	Rickard (2022)*
G46491a	TrGAroCN	BZEMUCO2 + NO → BZEMUCNO3	KR02NO*alpha_AN(10,2,0,1,0,temp,cair)	Rickard (2022)
G46491b	TrGAroCN	BZEMUCO2 + NO → .5 EPXC4DIAL + .5 GLYOX + .5 HO ₂ + .5 C3DIALO2 + .5 C32OH13CO + NO ₂	KR02NO*(1.-alpha_AN(10,2,0,1,0,temp,cair))	Rickard (2022)*
G46492	TrGAroCN	BZEMUCO2 + NO ₃ → .5 EPXC4DIAL + .5 GLYOX + .5 HO ₂ + .5 C3DIALO2 + .5 C32OH13CO + NO ₂	KR02NO3	Rickard (2022)*
G46493	TrGAroC	BZEMUCO2 → .5 EPXC4DIAL + .5 GLYOX + .5 HO ₂ + .5 C3DIALO2 + .5 C32OH13CO	k1_RO2sOR02	Rickard (2022)*
G46494	TrGAroCN	C5CO2DBPAN + OH → HCOCOCHO + CH ₃ CHO + CO ₂ + NO ₂	3.28E-11	Rickard (2022)
G46495	TrGAroCN	C5CO2DBPAN → C5CO2DBCO3 + NO ₂	k_PAN_M	Rickard (2022)
G46496	TrGAroCN	NBZQOOH + OH → NBZQO2	6.68E-11	Rickard (2022)
G46497	TrGAroC	CATEC1OOH + OH → CATEC1O2	k_ROOHRO	Rickard (2022)
G46498	TrGAroC	C6125CO + OH → C5CO14O2 + CO	6.45E-11	Rickard (2022)
G46499a	TrGAroCN	NBZQO2 + HO ₂ → NBZQOOH	k_RO2_HO2(temp,6)*(1.-r_COCH2O2_OH)	Rickard (2022)
G46499b	TrGAroCN	NBZQO2 + HO ₂ → C6CO4DB + NO ₂ + OH	k_RO2_HO2(temp,6)*r_COCH2O2_OH	Rickard (2022)*
G46500	TrGAroCN	NBZQO2 + NO → C6CO4DB + NO ₂ + NO ₂	KR02NO	Rickard (2022)*
G46501	TrGAroCN	NBZQO2 + NO ₃ → C6CO4DB + NO ₂ + NO ₂	KR02NO3	Rickard (2022)*
G46502	TrGAroCN	NBZQO2 → C6CO4DB + NO ₂	k1_RO2sOR02	Rickard (2022)*
G46503	TrGAroCN	DNPHENOOH + OH → DNPHENO2	k_ROOHRO	Rickard (2022)
G46504	TrGAroC	CATEC1O2 + HO ₂ → CATEC1OOH	k_RO2_HO2(temp,6)	Rickard (2022)
G46505a	TrGAroCN	CATEC1O2 + NO → CATEC1O + NO ₂	KR02NO	Rickard (2022)
G46505b	TrGAroCN	CATEC1O2 + NO ₂ → CATEC1O + NO ₃	k_C6H5O2_NO2	Jagiella and Zabel (2007)*
G46506	TrGAroCN	CATEC1O2 + NO ₃ → CATEC1O + NO ₂	KR02NO3	Rickard (2022)
G46507	TrGAroC	CATEC1O2 → CATEC1O	k1_RO2sOR02	Rickard (2022)
G46508	TrGAroC	BZEMUCCO3H + OH → BZEMUCCO3	4.37E-11	Rickard (2022)
G46509	TrGAroC	C6H5OOH + OH → C6H5O2	3.60E-12	Rickard (2022)
G46510	TrGAroC	BZEMUCCOAH + OH → BZEMUCCO + OH	1.31E-10	Rickard (2022)
G46511a	TrGAroC	BZEMUCCO3 + HO ₂ → BZEMUCCO2H + O ₃	KAPH02*r_CO3_03	Rickard (2022)
G46511b	TrGAroC	BZEMUCCO3 + HO ₂ → BZEMUCCO3H	KAPH02*r_CO3_00H	Rickard (2022)
G46511c	TrGAroC	BZEMUCCO3 + HO ₂ → C5DIALO2 + CO ₂ + OH	KAPH02*r_CO3_0H	Rickard (2022)
G46512	TrGAroCN	BZEMUCCO3 + NO → C5DIALO2 + CO ₂ + NO ₂	KAPNO	Rickard (2022)

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G46513	TrGAroCN	BZEMUCCO3 + NO ₂ → BZEMUCPAN	k_CH3C03_N02	Rickard (2022)
G46514	TrGAroCN	BZEMUCCO3 + NO ₃ → C5DIALO2 + CO ₂ + NO ₂	KR02N03*1.74	Rickard (2022)
G46515	TrGAroC	BZEMUCCO3 → C5DIALO2 + CO ₂	k1_R02RC03	Rickard (2022)*
G46516	TrGAroC	C6H5O2 + HO ₂ → C6H5OOH	k_R02_H02(temp, 6)	Rickard (2022)
G46517	TrGAroCN	C6H5O2 + NO → C6H5O + NO ₂	KR02NO	Rickard (2022)
G46518	TrGAroCN	C6H5O2 + NO ₃ → C6H5O + NO ₂	KR02N03	Rickard (2022)
G46519	TrGAroC	C6H5O2 → C6H5O	k1_R02sR02	Rickard (2022)
G46520	TrGAroCN	C6H5O2 + NO ₂ → C6H5O + NO ₃	k_C6H5O2_N02	Jagiella and Zabel (2007)
G46521	TrGAroCN	BZEMUCNO3 + OH → BZEMUCCO + NO ₂	4.38E-11	Rickard (2022)
G46522a	TrGAroC	BZBIPERO2 + HO ₂ → BZBIPEROOH	k_R02_H02(temp, 6)*(1.-r_BIPERO2_OH)	Rickard (2022)
G46522b	TrGAroC	BZBIPERO2 + HO ₂ → OH + GLYOX + HO ₂ + .5 BZFUONE + .5 BZFUONE	k_R02_H02(temp, 6)*r_BIPERO2_OH	Rickard (2022), Birdsall et al. (2010)*
G46523a	TrGAroCN	BZBIPERO2 + NO → BZBIPERNO3	KR02NO*alpha_AN(9, 2, 0, 0, 1, temp, cair)	Rickard (2022)
G46523b	TrGAroCN	BZBIPERO2 + NO → NO ₂ + GLYOX + HO ₂ + .5 BZFUONE + .5 BZFUONE	KR02NO*(1.-alpha_AN(9, 2, 0, 0, 1, temp, cair))	Rickard (2022)*
G46524	TrGAroCN	BZBIPERO2 + NO ₃ → NO ₂ + GLYOX + HO ₂ + .5 BZFUONE + .5 BZFUONE	KR02N03	Rickard (2022)*
G46525	TrGAroC	BZBIPERO2 → GLYOX + HO ₂ + BZFUONE	k1_R02sR02	Rickard (2022)*
G47200	TrGTerCN	CO235C6CHO + NO ₃ → CO235C6CO3 + HNO ₃	KN03AL*5.5	Rickard (2022)
G47201	TrGTerC	CO235C6CHO + OH → CO235C6CO3	6.70E-11	Rickard (2022)
G47202a	TrGTerC	CO235C6CO3 + HO ₂ → C235C6CO3H	KAPH02*(r_CO3_00H+r_CO3_03)	Rickard (2022)
G47202b	TrGTerC	CO235C6CO3 + HO ₂ → CO235C6O2 + CO ₂ + OH	KAPH02*r_CO3_OH	Rickard (2022)
G47203	TrGTerCN	CO235C6CO3 + NO → CO235C6O2 + CO ₂ + NO ₂	KAPNO	Rickard (2022)
G47204	TrGTerCN	CO235C6CO3 + NO ₂ → C7PAN3	k_CH3C03_N02	Rickard (2022)
G47205	TrGTerC	CO235C6CO3 → CO235C6O2 + CO ₂	k1_R02RC03	Rickard (2022)
G47206	TrGTerC	C235C6CO3H + OH → CO235C6CO3	4.75E-12	Rickard (2022)
G47207	TrGTerCN	C7PAN3 + OH → CO235C5CHO + CO + NO ₂	8.83E-13	Rickard (2022)
G47208	TrGTerCN	C7PAN3 → CO235C6CO3 + NO ₂	k_PAN_M	Rickard (2022)
G47209a	TrGTerC	C716O2 + HO ₂ → C716OOH	k_R02_H02(temp, 7)*r_COCH202_00H	Rickard (2022), Sander et al. (2019)
G47209b	TrGTerC	C716O2 + HO ₂ → CO13C4CHO + CH ₃ C(O) + OH	k_R02_H02(temp, 7)*r_COCH202_OH	Rickard (2022), Sander et al. (2019)
G47210	TrGTerCN	C716O2 + NO → CO13C4CHO + CH ₃ C(O) + NO ₂	KR02NO	Rickard (2022)*

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G47211	TrGTerC	C716O2 → CO13C4CHO + CH ₃ C(O)	k1_R02sR02	Rickard (2022)
G47212	TrGTerC	C716OOH + OH → CO235C6CHO + OH	1.20E-10	Rickard (2022)
G47213	TrGTerC	C721O2 + HO ₂ → C721OOH	k_R02_H02(temp, 7)	Rickard (2022)
G47214	TrGTerCN	C721O2 + NO → C722O2 + NO ₂	KR02NO	Rickard (2022)*
G47215	TrGTerC	C721O2 → C722O2	k1_R02pR02	Rickard (2022)
G47216	TrGTerC	C721OOH + OH → C721O2	1.27E-11	Rickard (2022)
G47217	TrGTerC	C722O2 + HO ₂ → C722OOH	k_R02_H02(temp, 7)	Rickard (2022)
G47218	TrGTerCN	C722O2 + NO → CH ₃ COCH ₃ + C44O2 + NO ₂	KR02NO	Rickard (2022)*
G47219	TrGTerC	C722O2 → CH ₃ COCH ₃ + C44O2	k1_R02tR02	Rickard (2022)
G47220	TrGTerC	C722OOH + OH → C722O2	3.31E-11	Rickard (2022)
G47221	TrGTerC	ROO6R3O2 → ROO6R5O2	5.68E10*EXP(-8745./temp)	Vereecken and Peeters (2012)
G47222	TrGTerCN	ROO6R3O2 + NO → ROO6R3O + NO ₂	KR02NO	Vereecken and Peeters (2012)*
G47223	TrGTerC	ROO6R3O2 + HO ₂ → 7 LCARBON	k_R02_H02(temp, 7)	Vereecken and Peeters (2012)*
G47224	TrGTerC	ROO6R3O2 → ROO6R3O	k1_R02sR02	Vereecken and Peeters (2012)
G47225	TrGTerC	ROO6R3O → 7 LCARBON + HO ₂	5.7E10*EXP(-2949./temp)	Vereecken and Peeters (2012)*
G47226	TrGTerC	ROO6R5O2 → 7 LCARBON + OH	9.17E10*EXP(-8706./temp)	Vereecken and Peeters (2012)*
G47400	TrGAroC	TOLUENE + OH → .07 C6H5CH2O2 + .18 CRESOL + .18 HO ₂ + .65 TLBIPERO2 + .10 TLEPOXMUC + .10 HO ₂	1.8E-12*EXP(340./temp)	Rickard (2022)*
G47401	TrGAroC	C6H5CH2O2 + HO ₂ → C6H5CH2OOH	1.5E-13*EXP(1310./temp)	Rickard (2022)
G47402a	TrGAroCN	C6H5CH2O2 + NO → C6H5CH2NO3	KR02NO*alpha_AN(7, 1, 0, 0, 0, temp, cair)	Rickard (2022)*
G47402b	TrGAroCN	C6H5CH2O2 + NO → BENZAL + HO ₂ + NO ₂	KR02NO*(1.-alpha_AN(7, 1, 0, 0, 0, temp, cair))	Rickard (2022)*
G47403	TrGAroCN	C6H5CH2O2 + NO ₃ → BENZAL + HO ₂ + NO ₂	KR02NO3	Rickard (2022)*
G47404	TrGAroC	C6H5CH2O2 → BENZAL + HO ₂	2.*(k_CH3O2*2.4E-14*EXP(1620./temp))**(.5)*R02	Rickard (2022)*
G47405	TrGAroCN	CRESOL + NO ₃ → .103 CRESO2 + .103 HNO ₃ + .506 NCRESO2 + .391 TOL1O + .391 HNO ₃	1.4E-11	Rickard (2022)*
G47406	TrGAroC	CRESOL + OH → .2 CRESO2 + .727 MCATECHOL + .727 HO ₂ + .073 TOL1O	4.65E-11	Rickard (2022)*
G47407a	TrGAroC	TLBIPERO2 + HO ₂ → TLBIPEROOH	k_R02_H02(temp, 7)*(1.-r_BIPERO2_OH)	Rickard (2022)

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G47407b	TrGAroC	$\text{TLBIPERO}_2 + \text{HO}_2 \rightarrow \text{OH} + .6 \text{ GLYOX} + .4 \text{ MGLYOX}$ + $\text{HO}_2 + .2 \text{ C4MDIAL} + .2 \text{ C5DICARB} + .2 \text{ TLFUONE}$ + $.2 \text{ BZFUONE} + .2 \text{ MALDIAL}$	$k_{\text{R02_HO2}}(\text{temp}, 7) * r_{\text{BIPERO2_OH}}$	Rickard (2022), Birdsall et al. (2010)*
G47408a	TrGAroCN	$\text{TLBIPERO}_2 + \text{NO} \rightarrow \text{NO}_2 + .6 \text{ GLYOX} + .4 \text{ MGLYOX}$ + $\text{HO}_2 + .2 \text{ C4MDIAL} + .2 \text{ C5DICARB} + .2 \text{ TLFUONE}$ + $.2 \text{ BZFUONE} + .2 \text{ MALDIAL}$	$K_{\text{R02NO}} * (1 - \alpha_{\text{AN}}(11, 2, 0, 0, 1, \text{temp}, \text{cair}))$	Rickard (2022)*
G47408b	TrGAroCN	$\text{TLBIPERO}_2 + \text{NO} \rightarrow \text{TLBIPERNO}_3$	$K_{\text{R02NO}} * \alpha_{\text{AN}}(11, 2, 0, 0, 1, \text{temp}, \text{cair})$	Rickard (2022)*
G47409	TrGAroCN	$\text{TLBIPERO}_2 + \text{NO}_3 \rightarrow \text{NO}_2 + .6 \text{ GLYOX} + .4 \text{ MGLYOX}$ + $\text{HO}_2 + .2 \text{ C4MDIAL} + .2 \text{ C5DICARB} + .2 \text{ TLFUONE}$ + $.2 \text{ BZFUONE} + .2 \text{ MALDIAL}$	K_{R02NO3}	Rickard (2022)*
G47410	TrGAroC	$\text{TLBIPERO}_2 \rightarrow .6 \text{ GLYOX} + .4 \text{ MGLYOX} + \text{HO}_2 + .2 \text{ C4MDIAL} + .2 \text{ C5DICARB} + .2 \text{ TLFUONE} + .2 \text{ BZFUONE} + .2 \text{ MALDIAL}$	$k_{1,\text{R02sOR02}}$	Rickard (2022)*
G47411	TrGAroCN	$\text{TLEPOXMUC} + \text{NO}_3 \rightarrow \text{TLEMUCCO}_3 + \text{HNO}_3$	$K_{\text{NO3AL}} * 2.75$	Rickard (2022)
G47412	TrGAroC	$\text{TLEPOXMUC} + \text{O}_3 \rightarrow \text{EPXC4DIAL} + .125 \text{ CH}_3\text{CHO} + .695 \text{ CH}_3\text{C(O)} + .57 \text{ CO} + .57 \text{ OH} + .125 \text{ HO}_2 + .1125 \text{ CH}_3\text{COCO}_2\text{H} + .0675 \text{ MGLYOX} + .0675 \text{ H}_2\text{O}_2 + .25 \text{ CO}_2$	$5.00E-18$	Rickard (2022)*
G47413	TrGAroC	$\text{TLEPOXMUC} + \text{OH} \rightarrow .31 \text{ TLEMUCCO}_3 + .69 \text{ TLEMUCCO}_2$	$7.99E-11$	Rickard (2022)*
G47414	TrGAroC	$\text{C6H5CH}_2\text{OOH} + \text{OH} \rightarrow \text{BENZAL} + \text{OH}$	$2.05E-11$	Rickard (2022)
G47415	TrGAroCN	$\text{C6H5CH}_2\text{NO}_3 + \text{OH} \rightarrow \text{BENZAL} + \text{NO}_2$	$6.03E-12$	Rickard (2022)
G47416	TrGAroCN	$\text{BENZAL} + \text{NO}_3 \rightarrow \text{C6H}_5\text{CO}_3 + \text{HNO}_3$	$2.40E-15$	Rickard (2022)
G47417	TrGAroC	$\text{BENZAL} + \text{OH} \rightarrow \text{C6H}_5\text{CO}_3$	$5.9E-12 * \exp(225/\text{temp})$	Rickard (2022)
G47418a	TrGAroC	$\text{CRESO}_2 + \text{HO}_2 \rightarrow \text{CRESOOH}$	$k_{\text{R02_HO2}}(\text{temp}, 7) * (1 - r_{\text{CHOHCH2O2_OH}})$	Rickard (2022)
G47418b	TrGAroC	$\text{CRESO}_2 + \text{HO}_2 \rightarrow .68 \text{ C5CO14OH} + .68 \text{ GLYOX} + \text{HO}_2 + .32 \text{ PTLQONE} + \text{OH}$	$k_{\text{R02_HO2}}(\text{temp}, 7) * r_{\text{CHOHCH2O2_OH}}$	Rickard (2022)*
G47419	TrGAroCN	$\text{CRESO}_2 + \text{NO} \rightarrow .68 \text{ C5CO14OH} + .68 \text{ GLYOX} + \text{HO}_2 + .32 \text{ PTLQONE} + \text{NO}_2$	K_{R02NO}	Rickard (2022)*
G47420	TrGAroCN	$\text{CRESO}_2 + \text{NO}_3 \rightarrow .68 \text{ C5CO14OH} + .68 \text{ GLYOX} + \text{HO}_2 + .32 \text{ PTLQONE} + \text{NO}_2$	K_{R02NO3}	Rickard (2022)*
G47421	TrGAroC	$\text{CRESO}_2 \rightarrow .68 \text{ C5CO14OH} + .68 \text{ GLYOX} + \text{HO}_2 + .32 \text{ PTLQONE}$	$k_{1,\text{R02ISOPD02}}$	Rickard (2022)*

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G47422a	TrGAroCN	NCRESO2 + HO ₂ → NCRESOOH	k_RO2_HO2(temp,7)*(1.-r_-CHOHCH2O2_OH)	Rickard (2022)
G47422b	TrGAroCN	NCRESO2 + HO ₂ → C5CO14OH + GLYOX + NO ₂ + OH	k_RO2_HO2(temp,7)*r_CHOHCH2O2_OH	Rickard (2022)*
G47423	TrGAroCN	NCRESO2 + NO → C5CO14OH + GLYOX + NO ₂ + NO ₂	KRO2NO	Rickard (2022)*
G47424	TrGAroCN	NCRESO2 + NO ₃ → C5CO14OH + GLYOX + NO ₂ + NO ₂	KRO2NO3	Rickard (2022)*
G47425	TrGAroCN	NCRESO2 → C5CO14OH + GLYOX + NO ₂	k1_RO2ISOPD02	Rickard (2022)*
G47426	TrGAroCN	TOL1O + NO ₂ → TOL1OHNO2	k_C6H5O_NO2	Rickard (2022), Platz et al. (1998)*
G47427	TrGAroC	TOL1O + O ₃ → OXYL1O2	k_C6H5O_03	Rickard (2022), Tao and Li (1999)
G47428	TrGAroCN	MCATECHOL + NO ₃ → MCATEC1O + HNO ₃	1.7E-10*1.0	Rickard (2022)
G47429	TrGAroC	MCATECHOL + O ₃ → MC3ODBCO2H + HCOCO ₂ H + HO ₂ + OH	2.8E-17	Rickard (2022)*
G47430	TrGAroC	MCATECHOL + OH → MCATEC1O	2.0E-10*1.0	Rickard (2022)
G47431	TrGAroC	TLBIPEROOH + OH → TLOBIPEROH + OH	9.64E-11	Rickard (2022)
G47432	TrGAroCN	TLBIPERNO3 + OH → TLOBIPEROH + NO ₂	7.16E-11	Rickard (2022)
G47433	TrGAroC	TLOBIPEROH + OH → C5CO14O2 + GLYOX	7.99E-11	Rickard (2022)
G47434a	TrGAroC	TLEMUCCO3 + HO ₂ → C615CO2O2 + CO ₂ + OH	KAPH02*r_CO3_OH	Rickard (2022)
G47434b	TrGAroC	TLEMUCCO3 + HO ₂ → TLEMUCCO2H + O ₃	KAPH02*r_CO3_03	Rickard (2022)
G47434c	TrGAroC	TLEMUCCO3 + HO ₂ → TLEMUCCO3H	KAPH02*r_CO3_OOH	Rickard (2022)
G47435	TrGAroCN	TLEMUCCO3 + NO → C615CO2O2 + CO ₂ + NO ₂	KAPNO	Rickard (2022)
G47436	TrGAroCN	TLEMUCCO3 + NO ₂ → TLEMUCPAN	k_CH3CO3_NO2	Rickard (2022)*
G47437	TrGAroCN	TLEMUCCO3 + NO ₃ → C615CO2O2 + CO ₂ + NO ₂	KRO2NO3*1.74	Rickard (2022)
G47438	TrGAroC	TLEMUCCO3 → C615CO2O2 + CO ₂	k1_RO2RC03	Rickard (2022)*
G47439a	TrGAroC	TLEMUCO2 + HO ₂ → TLEMUCOOH	k_RO2_HO2(temp,7)*(1.-r_-CHOHCH2O2_OH-r_COCH2O2_OH)	Rickard (2022)
G47439b	TrGAroC	TLEMUCO2 + HO ₂ → .5 C3DIALO2 + .5 CO2H3CHO + .5 EPXC4DIAL + .5 MGLYOX + .5 HO ₂ + OH	k_RO2_HO2(temp,7)*(r_CHOHCH2O2_OH+r_COCH2O2_OH)	Rickard (2022)*
G47440a	TrGAroCN	TLEMUCO2 + NO → TLEMUCN03	KRO2NO*alpha_AN(11,2,1,0,0, temp,cair)	Rickard (2022)
G47440b	TrGAroCN	TLEMUCO2 + NO → .5 C3DIALO2 + .5 CO2H3CHO + .5 EPXC4DIAL + .5 MGLYOX + .5 HO ₂ + NO ₂	KRO2NO*(1.-alpha_AN(11,2,1,0,0, temp,cair))	Rickard (2022)*

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G47441	TrGAroCN	TLEMUCO2 + NO ₃ → .5 C3DIALO2 + .5 CO2H3CHO + .5 EPXC4DIAL + .5 MGLYOX + .5 HO ₂ + NO ₂	KR02N03	Rickard (2022)*
G47442	TrGAroC	TLEMUCO2 → .5 C3DIALO2 + .5 CO2H3CHO + .5 EPXC4DIAL + .5 MGLYOX + .5 HO ₂	k1_R02s0R02	Rickard (2022)*
G47443a	TrGAroC	C6H5CO3 + HO ₂ → C6H5CO3H	1.1E-11*EXP(364./temp)*0.65	Roth et al. (2010)
G47443b	TrGAroC	C6H5CO3 + HO ₂ → C6H5O2 + CO ₂ + OH	1.1E-11*EXP(364./temp)*0.20	Roth et al. (2010)
G47443c	TrGAroC	C6H5CO3 + HO ₂ → PHCOOH + O ₃	1.1E-11*EXP(364./temp)*0.15	Roth et al. (2010)
G47444	TrGAroCN	C6H5CO3 + NO → C6H5O2 + CO ₂ + NO ₂	KAPNO	Rickard (2022)
G47445	TrGAroCN	C6H5CO3 + NO ₂ → PBZN	k_CH3C03_N02	Rickard (2022)*
G47446	TrGAroCN	C6H5CO3 + NO ₃ → C6H5O2 + CO ₂ + NO ₂	KR02N03*1.74	Rickard (2022)
G47447	TrGAroC	C6H5CO3 → C6H5O2 + CO ₂	k1_R02RC03	Rickard (2022)*
G47448	TrGAroC	CRESOOH + OH → CRESO2	1.15E-10	Rickard (2022)
G47449	TrGAroCN	NCRESOOH + OH → NCRESO2	1.07E-10	Rickard (2022)
G47450	TrGAroCN	TOL1OHNO2 + NO ₃ → NCRES1O + HNO ₃	3.13E-13*1.0	Rickard (2022)
G47451	TrGAroCN	TOL1OHNO2 + OH → NCRES1O	2.8E-12	Rickard (2022)
G47452	TrGAroC	OXYL1O2 + HO ₂ → OXYL1OOH	k_R02_H02(temp,7)	Rickard (2022)
G47453	TrGAroCN	OXYL1O2 + NO → TOL1O + NO ₂	KR02NO	Rickard (2022)
G47454	TrGAroCN	OXYL1O2 + NO ₂ → TOL1O + NO ₃	k_C6H502_N02	Jagiella and Zabel (2007)*
G47455	TrGAroCN	OXYL1O2 + NO ₃ → TOL1O + NO ₂	KR02N03	Rickard (2022)
G47456	TrGAroC	OXYL1O2 → TOL1O	k1_R02sR02	Rickard (2022)
G47457	TrGAroCN	MCATEC1O + NO ₂ → MNCATECH	k_C6H50_N02	Rickard (2022), Platz et al. (1998)
G47458	TrGAroC	MCATEC1O + O ₃ → MCATEC1O2	k_C6H50_03	Rickard (2022), Tao and Li (1999)
G47459	TrGAroC	TLEMUCCO2H + OH → C615CO2O2 + CO ₂	5.98E-11	Rickard (2022)
G47460	TrGAroC	TLEMUCCO3H + OH → TLEMUCCO3	6.29E-11	Rickard (2022)
G47461	TrGAroCN	TLEMUCPAN + OH → C5DICARB + CO + CO ₂ + NO ₂	5.96E-11	Rickard (2022)
G47462	TrGAroCN	TLEMUCPAN → TLEMUCCO3 + NO ₂	k_PAN_M	Rickard (2022)
G47463	TrGAroC	TLEMUCOOH + OH → TLEMUCCO + OH	7.04E-11	Rickard (2022)
G47464	TrGAroCN	TLEMUCNO3 + OH → TLEMUCCO + NO ₂	3.06E-11	Rickard (2022)
G47465	TrGAroC	TLEMUCCO + OH → CH ₃ C(O) + EPXC4DIAL + CO	4.06E-11	Rickard (2022)
G47466	TrGAroC	C6H5CO3H + OH → C6H5CO3	4.66E-12	Rickard (2022)
G47467	TrGAroC	PHCOOH + OH → C6H5O2 + CO ₂	1.10E-12	Rickard (2022)
G47468	TrGAroCN	PBZN + OH → C6H5OOH + CO + NO ₂	1.06E-12	Rickard (2022)
G47469	TrGAroCN	PBZN → C6H5CO3 + NO ₂	k_PAN_M*0.67	Rickard (2022)

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G47470	TrGAroCN	PTLQONE + NO ₃ → NPTLQO2	1.00E-12	Rickard (2022)
G47471	TrGAroC	PTLQONE + OH → PTLQO2	2.3E-11	Rickard (2022)
G47472	TrGAroCN	NCRES1O + NO ₂ → DNCRES	k_C6H5O_N02	Rickard (2022), Platz et al. (1998)
G47473	TrGAroCN	NCRES1O + O ₃ → NCRES1O2	k_C6H5O_03	Rickard (2022), Tao and Li (1999)
G47474	TrGAroC	OXYL1OOH + OH → OXYL1O2	4.65E-11	Rickard (2022)
G47475	TrGAroCN	MNCATECH + NO ₃ → MNNCATECO2	5.03E-12	Rickard (2022)
G47476	TrGAroCN	MNCATECH + OH → MNCATECO2	6.83E-12	Rickard (2022)
G47477	TrGAroC	MCATEC1O2 + HO ₂ → MCATEC1OOH	k_R02_H02(temp,7)	Rickard (2022)
G47478	TrGAroCN	MCATEC1O2 + NO → MCATEC1O + NO ₂	KR02NO	Rickard (2022)
G47479	TrGAroCN	MCATEC1O2 + NO ₂ → MCATEC1O + NO ₃	k_C6H5O2_N02	Jagiella and Zabel (2007)*
G47480	TrGAroCN	MCATEC1O2 + NO ₃ → MCATEC1O + NO ₂	KR02N03	Rickard (2022)
G47481	TrGAroC	MCATEC1O2 → MCATEC1O	k1_R02sOR02	Rickard (2022)
G47482a	TrGAroCN	NPTLQO2 + HO ₂ → NPTLQOOH	k_R02_H02(temp,7)*(1.-r_COCH202_OH)	Rickard (2022)
G47482b	TrGAroCN	NPTLQO2 + HO ₂ → C7CO4DB + NO ₂ + OH	k_R02_H02(temp,7)*r_COCH202_OH	Rickard (2022)*
G47483	TrGAroCN	NPTLQO2 + NO → C7CO4DB + NO ₂ + NO ₂	KR02NO	Rickard (2022)*
G47484	TrGAroCN	NPTLQO2 + NO ₃ → C7CO4DB + NO ₂ + NO ₂	KR02N03	Rickard (2022)*
G47485	TrGAroCN	NPTLQO2 → C7CO4DB + NO ₂	k1_R02sOR02	Rickard (2022)*
G47486a	TrGAroC	PTLQO2 + HO ₂ → PTLQOOH	k_R02_H02(temp,7)*(1.-r_COCH202_OH-r_COCH202_OH)	Rickard (2022)
G47486b	TrGAroC	PTLQO2 + HO ₂ → C6CO2OHCO3 + OH	k_R02_H02(temp,7)*(r_COCH202_OH+r_COCH202_OH)	Rickard (2022)*
G47487	TrGAroCN	PTLQO2 + NO → C6CO2OHCO3 + NO ₂	KR02NO	Rickard (2022)*
G47488	TrGAroCN	PTLQO2 + NO ₃ → C6CO2OHCO3 + NO ₂	KR02N03	Rickard (2022)*
G47489	TrGAroC	PTLQO2 → C6CO2OHCO3	k1_R02sOR02	Rickard (2022)*
G47490	TrGAroCN	DNCRES + NO ₃ → NDNCRESO2	7.83E-15	Rickard (2022)
G47491	TrGAroCN	DNCRES + OH → DNCRESO2	5.10E-14	Rickard (2022)
G47492	TrGAroCN	NCRES1O2 + HO ₂ → NCRES1OOH	k_R02_H02(temp,7)	Rickard (2022)
G47493	TrGAroCN	NCRES1O2 + NO → NCRES1O + NO ₂	KR02NO	Rickard (2022)
G47494	TrGAroCN	NCRES1O2 + NO ₂ → NCRES1O + NO ₃	k_C6H5O2_N02	Jagiella and Zabel (2007)*
G47495	TrGAroCN	NCRES1O2 + NO ₃ → NCRES1O + NO ₂	KR02N03	Rickard (2022)
G47496	TrGAroCN	NCRES1O2 → NCRES1O	k1_R02sOR02	Rickard (2022)

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G47497a	TrGAroCN	MNNCATECO2 + HO ₂ → MNNCATCOOH	k_RO2_HO2(temp,7)*(1.-r_-CHOHCH2O2_OH)	Rickard (2022)
G47497b	TrGAroCN	MNNCATECO2 + HO ₂ → NC4MDCO2HN + HCOCO ₂ H + NO ₂ + OH	k_RO2_HO2(temp,7)*r_CHOHCH2O2_OH	Rickard (2022)*
G47498	TrGAroCN	MNNCATECO2 + NO → NC4MDCO2HN + HCOCO ₂ H + NO ₂ + NO ₂	KR02NO	Rickard (2022)*
G47499	TrGAroCN	MNNCATECO2 + NO ₃ → NC4MDCO2HN + HCOCO ₂ H + NO ₂ + NO ₂	KR02N03	Rickard (2022)*
G47500	TrGAroCN	MNNCATECO2 → NC4MDCO2HN + HCOCO ₂ H + NO ₂	k1_RO2ISOPD02	Rickard (2022)
G47501a	TrGAroCN	MNCATECO2 + HO ₂ → MNCATECOOH	k_RO2_HO2(temp,7)*(1.-r_-CHOHCH2O2_OH)	Rickard (2022)
G47501b	TrGAroCN	MNCATECO2 + HO ₂ → NC4MDCO2HN + HCOCO ₂ H + HO ₂ + OH	k_RO2_HO2(temp,7)*r_CHOHCH2O2_OH	Rickard (2022)*
G47502	TrGAroCN	MNCATECO2 + NO → NC4MDCO2HN + HCOCO ₂ H + HO ₂ + NO ₂	KR02NO	Rickard (2022)*
G47503	TrGAroCN	MNCATECO2 + NO ₃ → NC4MDCO2HN + HCOCO ₂ H + HO ₂ + NO ₂	KR02N03	Rickard (2022)*
G47504	TrGAroCN	MNCATECO2 → NC4MDCO2HN + HCOCO ₂ H + HO ₂	k1_RO2ISOPD02	Rickard (2022)*
G47505	TrGAroC	MCATEC1OOH + OH → MCATEC1O2	2.05E-10	Rickard (2022)
G47506	TrGAroCN	NPTLQOOH + OH → NPTLQO2	8.56E-11	Rickard (2022)
G47507	TrGAroC	PTLQOOH + OH → PTLQCO + OH	1.42E-10	Rickard (2022)
G47508	TrGAroC	PTLQCO + OH → C6CO2OHCO3	7.95E-11	Rickard (2022)
G47509a	TrGAroCN	NDNCRESO2 + HO ₂ → NDNCRESOOH	k_RO2_HO2(temp,7)*(1.-r_-CHOHCH2O2_OH)	Rickard (2022)
G47509b	TrGAroCN	NDNCRESO2 + HO ₂ → NC4MDCO2HN + HNO ₃ + 2 CO + NO ₂ + OH	k_RO2_HO2(temp,7)*r_CHOHCH2O2_OH	Rickard (2022)*
G47510	TrGAroCN	NDNCRESO2 + NO → NC4MDCO2HN + HNO ₃ + 2 CO + NO ₂ + NO ₂	KR02NO	Rickard (2022)*
G47511	TrGAroCN	NDNCRESO2 + NO ₃ → NC4MDCO2HN + HNO ₃ + 2 CO + NO ₂ + NO ₂	KR02N03	Rickard (2022)*
G47512	TrGAroCN	NDNCRESO2 → NC4MDCO2HN + HNO ₃ + 2 CO + NO ₂	k1_RO2ISOPD02	Rickard (2022)*
G47513a	TrGAroCN	DNCRESO2 + HO ₂ → DNCRESOOH	k_RO2_HO2(temp,7)*(1.-r_-CHOHCH2O2_OH)	Rickard (2022)
G47513b	TrGAroCN	DNCRESO2 + HO ₂ → NC4MDCO2HN + HCOCO ₂ H + NO ₂ + OH	k_RO2_HO2(temp,7)*r_CHOHCH2O2_OH	Rickard (2022)*

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G47514	TrGAroCN	DNCRESO2 + NO → NC4MDCO2HN + HCOCO ₂ H + NO ₂ + NO ₂	KR02NO	Rickard (2022)*
G47515	TrGAroCN	DNCRESO2 + NO ₃ → NC4MDCO2HN + HCOCO ₂ H + NO ₂ + NO ₂	KR02NO3	Rickard (2022)*
G47516	TrGAroCN	DNCRESO2 → NC4MDCO2HN + HCOCO ₂ H + NO ₂	k1_RO2ISOPD02	Rickard (2022)*
G47517	TrGAroCN	NCRES1OOH + OH → NCRES1O2	1.53E-12	Rickard (2022)
G47518	TrGAroCN	MNNCATCOOH + OH → MNNCATECO2	k_ROOHRO	Rickard (2022)
G47519	TrGAroCN	MNCATECOOH + OH → MNCATECO2	k_ROOHRO	Rickard (2022)
G47520	TrGAroC	C7CO4DB + OH → CO + CO + CH ₃ C(O) + HCOCOCHO	9.58E-11	Rickard (2022)
G47521a	TrGAroC	C6CO2OHCO3 + HO ₂ → C5134CO2OH + HO ₂ + CO + CO ₂ + OH	KAPH02*r_CO3_OH	Rickard (2022)
G47521b	TrGAroC	C6CO2OHCO3 + HO ₂ → C6COOHCO3H	KAPH02*(r_CO3_00H+r_CO3_03)	Rickard (2022)
G47522	TrGAroCN	C6CO2OHCO3 + NO → C5134CO2OH + HO ₂ + CO + CO ₂ + NO ₂	KAPNO	Rickard (2022)
G47523	TrGAroCN	C6CO2OHCO3 + NO ₂ → C6CO2OHPAN	k_CH3CO3_N02	Rickard (2022)
G47524	TrGAroCN	C6CO2OHCO3 + NO ₃ → C5134CO2OH + HO ₂ + CO + CO ₂ + NO ₂	KR02NO3*1.74	Rickard (2022)
G47525	TrGAroC	C6CO2OHCO3 → C5134CO2OH + HO ₂ + CO + CO ₂	k1_RO2RC03	Rickard (2022)
G47526	TrGAroCN	NDNCRESOOH + OH → NDNCRESO2	k_ROOHRO	Rickard (2022)
G47527	TrGAroCN	DNCRESOOH + OH → DNCRESO2	k_ROOHRO	Rickard (2022)
G47528	TrGAroC	C6COOHCO3H + OH → C6CO2OHCO3	9.29E-11	Rickard (2022)
G47529	TrGAroCN	C6CO2OHPAN + OH → C5134CO2OH + CO + CO + NO ₂	8.96E-11	Rickard (2022)
G47530	TrGAroCN	C6CO2OHPAN → C6CO2OHCO3 + NO ₂	k_PAN_M	Rickard (2022)
G48200	TrGTerC	C85O2 → C86O2	k1_RO2tR02	Rickard (2022)
G48201	TrGTerC	C85O2 + HO ₂ → C85OOH	k_RO2_HO2(temp,8)	Rickard (2022)
G48202	TrGTerCN	C85O2 + NO → C86O2 + NO ₂	KR02NO	Rickard (2022)*
G48203	TrGTerC	C85OOH + OH → C85O2	1.29E-11	Rickard (2022)
G48204	TrGTerC	C86O2 → C511O2 + CH ₃ COCH ₃	k1_RO2tR02	Rickard (2022)
G48205	TrGTerCN	C86O2 + NO → C511O2 + CH ₃ COCH ₃ + NO ₂	KR02NO	Rickard (2022)*
G48206	TrGTerC	C86O2 + HO ₂ → C86OOH	k_RO2_HO2(temp,8)	Rickard (2022)
G48207	TrGTerC	C86OOH + OH → C86O2	3.45E-11	Rickard (2022)
G48208	TrGTerC	C811O2 → C812O2	k1_RO2pR02	Rickard (2022)
G48209	TrGTerC	C811O2 + HO ₂ → 8 LCARBON	k_RO2_HO2(temp,8)	Rickard (2022)

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G48210	TrGTerCN	C811O2 + NO → C812O2 + NO2	KR02NO	Rickard (2022)*
G48211	TrGTerC	C812O2 → C813O2	k1_R02tR02	Rickard (2022)
G48212	TrGTerCN	C812O2 + NO → C813O2 + NO2	KR02NO	Rickard (2022)*
G48213	TrGTerC	C812O2 + HO2 → C812OOH	k_R02_HO2(temp, 8)	Rickard (2022)
G48214	TrGTerC	C812OOH + OH → C812O2	1.09E-11	Rickard (2022)
G48215	TrGTerC	C813O2 → CH3COCH3 + C512O2	k1_R02tR02	Rickard (2022)
G48216	TrGTerCN	C813O2 + NO → CH3COCH3 + C512O2 + NO2	KR02NO	Rickard (2022)*
G48217	TrGTerC	C813O2 + HO2 → C813OOH	k_R02_HO2(temp, 8)	Rickard (2022)
G48218	TrGTerC	C813OOH + OH → C813O2	1.86E-11	Rickard (2022)
G48219	TrGTerCN	C721CHO + NO3 → C721CO3 + HNO3	KN03AL*8.5	Rickard (2022)
G48220	TrGTerC	C721CHO + OH → C721CO3	2.63E-11	Rickard (2022)
G48221a	TrGTerC	C721CO3 + HO2 → C721CO3H	KAPHO2*r_CO3_00H	Rickard (2022)
G48221b	TrGTerC	C721CO3 + HO2 → C721O2 + CO2 + OH	KAPHO2*r_CO3_OH	Rickard (2022)
G48221c	TrGTerC	C721CO3 + HO2 → NORPINIC + O3	KAPHO2*r_CO3_03	Rickard (2022)
G48222	TrGTerCN	C721CO3 + NO → C721O2 + CO2 + NO2	KAPNO	Rickard (2022)*
G48223	TrGTerCN	C721CO3 + NO2 → C721PAN	k_CH3CO3_N02	Rickard (2022)
G48224	TrGTerCN	C721CO3 + NO3 → C721O2 + CO2 + NO2	KR02N03*1.74	Rickard (2022)
G48225	TrGTerC	C721CO3 → C721O2 + CO2	k1_R02RC03*0.9	Sander et al. (2019)
G48226	TrGTerC	C721CO3 → NORPINIC	k1_R02RC03*0.1	Sander et al. (2019)
G48227	TrGTerC	C721CO3H + OH → C721CO3	9.65E-12	Rickard (2022)
G48228	TrGTerC	NORPINIC + OH → C721O2 + CO2	6.57E-12	Rickard (2022)
G48229	TrGTerCN	C721PAN + OH → C721OOH + CO + NO2	2.96E-12	Rickard (2022)
G48230	TrGTerCN	C721PAN → C721CO3 + NO2	k_PAN_M	Rickard (2022)
G48231	TrGTerC	C8BC + OH → C8BCO2	3.04E-12	Rickard (2022)
G48232	TrGTerC	C8BCO2 + HO2 → C8BCOOH	k_R02_HO2(temp, 8)	Rickard (2022)
G48233a	TrGTerCN	C8BCO2 + NO → C89O2 + NO2	KR02NO*(1.-alpha_AN(8, 2, 0, 0, 0, temp, cair))	Rickard (2022)
G48233b	TrGTerCN	C8BCO2 + NO → C8BCNO3	KR02NO*alpha_AN(8, 2, 0, 0, 0, temp, cair)	Rickard (2022)
G48234	TrGTerC	C8BCO2 → C89O2	k1_R02sR02	Rickard (2022)
G48235	TrGTerC	C8BCOOH + OH → C8BCCO + OH	1.62E-11	Rickard (2022)
G48236	TrGTerCN	C8BCNO3 + OH → C8BCCO + NO2	1.84E-12	Rickard (2022)
G48237	TrGTerC	C8BCCO + OH → C89O2	3.94E-12	Rickard (2022)
G48238	TrGTerC	C89O2 + HO2 → C89OOH	k_R02_HO2(temp, 8)	Rickard (2022)

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G48239a	TrGTerCN	C89O2 + NO → C810O2 + NO ₂	KR02NO*(1.-alpha_AN(7,2,0,0,0, temp,cair))	Rickard (2022)
G48239b	TrGTerCN	C89O2 + NO → C89NO3	KR02NO*alpha_AN(7,2,0,0,0,temp, cair)	Rickard (2022)
G48240	TrGTerCN	C89O2 + NO ₃ → C810O2 + NO ₂	KR02NO3	Rickard (2022)
G48241	TrGTerC	C89O2 → C810O2	k1_R02tR02	Rickard (2022)
G48242	TrGTerC	C89OOH + OH → C89O2	3.61E-11	Rickard (2022)
G48243	TrGTerCN	C89NO3 + OH → CH ₃ COCH ₃ + CO13C4CHO + NO ₂	2.56E-11	Rickard (2022)
G48244	TrGTerC	C810O2 + HO ₂ → C810OOH	k_R02_H02(temp,8)	Rickard (2022)
G48245a	TrGTerCN	C810O2 + NO → CH ₃ COCH ₃ + C514O2 + NO ₂	KR02NO*(1.-alpha_AN(10,3,0,0,0, temp,cair))	Rickard (2022)
G48245b	TrGTerCN	C810O2 + NO → C810NO3	KR02NO*alpha_AN(10,3,0,0,0, temp,cair)	Rickard (2022)
G48246	TrGTerCN	C810O2 + NO ₃ → CH ₃ COCH ₃ + C514O2 + NO ₂	KR02NO3	Rickard (2022)
G48247	TrGTerC	C810O2 → CH ₃ COCH ₃ + C514O2	k1_R02tR02	Rickard (2022)
G48248	TrGTerC	C810OOH + OH → C810O2	8.35E-11	Rickard (2022)
G48249	TrGTerCN	C810NO3 + OH → CH ₃ COCH ₃ + CO13C4CHO + NO ₂	4.96E-11	Rickard (2022)
G48400a	TrGAroC	LXYL + OH → TLEPOXMUC + HO ₂ + LCARBON	0.401E-11	Rickard (2022)*
G48400b	TrGAroC	LXYL + OH → C6H5CH2O2 + LCARBON	0.101E-11	Rickard (2022)*
G48400c	TrGAroC	LXYL + OH → CRESOL + LCARBON	0.261E-11	Rickard (2022)*
G48400d	TrGAroC	LXYL + OH → TLBIPERO2 + HO ₂ + LCARBON	0.932E-11	Rickard (2022)*
G48401	TrGAroCN	LXYL + NO ₃ → C6H5CH2O2 + HNO ₃ + LCARBON	3.9E-16	Rickard (2022)*
G48402	TrGAroC	EBENZ + OH → .10 TLEPOXMUC + .07 C6H5CH2O2 + .18 CRESOL + .65 TLBIPERO2 + .28 HO ₂ + LCARBON	7.00E-12	Rickard (2022)*
G48403	TrGAroCN	EBENZ + NO ₃ → C6H5CH2O2 + HNO ₃ + LCARBON	1.20E-16	Rickard (2022)*
G48404	TrGAroCN	STYRENE + NO ₃ → NSTYRENO2	1.50E-12	Rickard (2022)
G48405	TrGAroC	STYRENE + O ₃ → .545 HCHO + .1 BENZENE + .28 C6H5O2 + .56 CO + .36 OH + .28 HO ₂ + .075 PHCOOH + .545 BENZAL + .09 H ₂ O ₂ + .075 HCOOH + .2 CO ₂	1.70E-17	Rickard (2022)*
G48406	TrGAroC	STYRENE + OH → STYRENO2	5.80E-11	Rickard (2022)
G48407	TrGAroCN	NSTYRENO2 + HO ₂ → NSTYRENOOH	k_R02_H02(temp,8)	Rickard (2022)
G48408	TrGAroCN	NSTYRENO2 + NO → NO ₂ + NO ₂ + HCHO + BENZAL	KR02NO	Rickard (2022)*
G48409	TrGAroCN	NSTYRENO2 + NO ₃ → NO ₂ + NO ₂ + HCHO + BENZAL	KR02NO3	Rickard (2022)*
G48410	TrGAroCN	NSTYRENO2 → NO ₂ + HCHO + BENZAL	k1_R02sR02	Rickard (2022)*

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G48411	TrGAroCN	NSTYRENOOH + OH → NSTYRENO2	6.16E-11	Rickard (2022)
G48412a	TrGAroC	STYRENO2 + HO ₂ → STYRENOOH	k_RO2_HO2(temp,8)*(1.-r_CHOHCH2O2_OH)	Rickard (2022)
G48412b	TrGAroC	STYRENO2 + HO ₂ → HO ₂ + OH + HCHO + BENZAL	k_RO2_HO2(temp,8)*r_CHOHCH2O2_OH	Rickard (2022)*
G48413	TrGAroCN	STYRENO2 + NO → NO ₂ + HO ₂ + HCHO + BENZAL	KRO2NO	Rickard (2022)*
G48414	TrGAroCN	STYRENO2 + NO ₃ → NO ₂ + HO ₂ + HCHO + BENZAL	KRO2NO3	Rickard (2022)*
G48415	TrGAroC	STYRENO2 → HO ₂ + HCHO + BENZAL	k1_RO2sR02	Rickard (2022)*
G48416	TrGAroC	STYRENOOH + OH → STYRENO2	6.16E-11	Rickard (2022)
G49200	TrGTerC	C96O2 → C97O2	k1_RO2pR02	Rickard (2022)
G49201	TrGTerC	C96O2 + HO ₂ → C96OOH	k_RO2_HO2(temp,9)	Rickard (2022)
G49202a	TrGTerCN	C96O2 + NO → C97O2 + NO ₂	KRO2NO*(1.-alpha_AN(10,1,0,0,0,temp,cair))	Rickard (2022)
G49202b	TrGTerCN	C96O2 + NO → C96NO3	KRO2NO*alpha_AN(10,1,0,0,0,temp,cair)	Rickard (2022)
G49203	TrGTerCN	C96NO3 + OH → NORPINAL + NO ₂	2.88E-12	Rickard (2022)
G49204a	TrGTerC	C96OOH + OH → C96O2	k_ROOHR0	Rickard (2022)
G49205b	TrGTerC	C96OOH + OH → NORPINAL + OH	1.30E-11	Rickard (2022)
G49206	TrGTerC	C97O2 → C98O2	k1_RO2tR02	Rickard (2022)
G49207	TrGTerCN	C97O2 + NO → C98O2 + NO ₂	KRO2NO	Rickard (2022)*
G49208a	TrGTerC	C97O2 + HO ₂ → C97OOH	k_RO2_HO2(temp,9)*r_COCH2O2_OOH	Rickard (2022), Sander et al. (2019)
G49208b	TrGTerC	C97O2 + HO ₂ → C98O2 + OH	k_RO2_HO2(temp,9)*r_COCH2O2_OH	Rickard (2022), Sander et al. (2019)
G49209	TrGTerC	C97OOH + OH → C97O2	1.05E-11	Rickard (2022)
G49210	TrGTerC	C98O2 → C614O2 + CH ₃ COCH ₃	k1_RO2tR02	Rickard (2022)
G49211a	TrGTerCN	C98O2 + NO → C614O2 + CH ₃ COCH ₃ + NO ₂	KRO2NO*(1.-alpha_AN(12,3,0,0,0,temp,cair))	Rickard (2022)
G49211b	TrGTerCN	C98O2 + NO → 9 LCARBON + LNITROGEN	KRO2NO*alpha_AN(12,3,0,0,0,temp,cair)	Rickard (2022)
G49212	TrGTerC	C98O2 + HO ₂ → C98OOH	k_RO2_HO2(temp,9)	Rickard (2022)
G49213	TrGTerC	C98OOH + OH → C98O2	2.05E-11	Rickard (2022)
G49214	TrGTerC	NORPINAL + OH → C85CO3	2.64E-11	Rickard (2022)
G49215	TrGTerCN	NORPINAL + NO ₃ → C85CO3 + HNO ₃	KNO3AL*8.5	Rickard (2022)
G49216	TrGTerC	C85CO3 → C85O2 + CO ₂	k1_RO2RC03	Rickard (2022)
G49217	TrGTerCN	C85CO3 + NO → C85O2 + CO ₂ + NO ₂	KAPNO	Rickard (2022)

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G49218	TrGTerCN	C85CO3 + NO ₂ → C9PAN2	k_CH3CO3_N02	Rickard (2022)
G49219a	TrGTerC	C85CO3 + HO ₂ → C85CO3H	KAPH02*(r_CO3_00H+r_CO3_03)	Rickard (2022)
G49219b	TrGTerC	C85CO3 + HO ₂ → C85O ₂ + CO ₂ + OH	KAPH02*r_CO3_OH	Rickard (2022)
G49220	TrGTerCN	C9PAN2 → C85CO3 + NO ₂	k_PAN_M	Rickard (2022)
G49221	TrGTerCN	C9PAN2 + OH → C85OOH + CO + NO ₂	6.60E-12	Rickard (2022)
G49222	TrGTerC	C85CO3H + OH → C85CO3	1.02E-11	Rickard (2022)
G49223a	TrGTerC	C89CO3 → .8 C811CO3 + .2 C89O ₂ + .2 CO ₂	k1_RO2RC03*0.9	Sander et al. (2019)
G49223b	TrGTerC	C89CO3 → C89CO2H	k1_RO2RC03*0.1	Sander et al. (2019)
G49224a	TrGTerC	C89CO3 + HO ₂ → C89CO3H	KAPH02*r_CO3_00H	Rickard (2022)
G49224b	TrGTerC	C89CO3 + HO ₂ → C89CO2H + O ₃	KAPH02*r_CO3_03	Rickard (2022)
G49224c	TrGTerC	C89CO3 + HO ₂ → .80 C811CO3 + .20 C89O ₂ + .2 CO ₂ + OH	KAPH02*r_CO3_OH	Rickard (2022)
G49225	TrGTerCN	C89CO3 + NO ₂ → C89PAN	k_CH3CO3_N02	Rickard (2022)
G49226	TrGTerCN	C89CO3 + NO → .8 C811CO3 + .2 C89O ₂ + .2 CO ₂ + NO ₂	KAPNO	Rickard (2022)
G49227	TrGTerC	C89CO2H + OH → .8 C811CO3 + .2 C89O ₂ + .2 CO ₂	2.69E-11	Rickard (2022)
G49228	TrGTerC	C89CO3H + OH → C89CO3	3.00E-11	Rickard (2022)
G49229	TrGTerCN	C89PAN → C89CO3 + NO ₂	k_PAN_M	Rickard (2022)
G49230	TrGTerCN	C89PAN + OH → CH ₃ COCH ₃ + CO13C4CHO + CO + NO ₂	2.52E-11	Rickard (2022)
G49231a	TrGTerC	C811CO3 → C811O ₂ + CO ₂	k1_RO2RC03*0.9	Sander et al. (2019)
G49231b	TrGTerC	C811CO3 → PINIC	k1_RO2RC03*0.1	Sander et al. (2019)
G49232a	TrGTerC	C811CO3 + HO ₂ → C811CO3H	KAPH02*r_CO3_00H	Rickard (2022)
G49232b	TrGTerC	C811CO3 + HO ₂ → PINIC + O ₃	KAPH02*r_CO3_03	Rickard (2022)
G49232c	TrGTerC	C811CO3 + HO ₂ → C811O ₂ + CO ₂ + OH	KAPH02*r_CO3_OH	Rickard (2022)
G49233	TrGTerCN	C811CO3 + NO → C811O ₂ + CO ₂ + NO ₂	KAPNO	Rickard (2022)
G49234	TrGTerCN	C811CO3 + NO ₂ → C811PAN	k_CH3CO3_N02	Rickard (2022)
G49235	TrGTerC	PINIC + OH → C811O ₂ + CO ₂	7.29E-12	Rickard (2022)
G49236	TrGTerC	NOPINONE + OH → NOPINDO2	1.55E-11	Capouet et al. (2008), Rickard (2022)
G49237a	TrGTerC	NOPINDO2 + HO ₂ → NOPINDOOH	k_RO2_H02(temp,9)*r_COCH2O2_00H	Rickard (2022), Sander et al. (2019)
G49237b	TrGTerC	NOPINDO2 + HO ₂ → C89CO3 + OH	k_RO2_H02(temp,9)*r_COCH2O2_OH	Rickard (2022), Sander et al. (2019)
G49238	TrGTerCN	NOPINDO2 + NO → C89CO3 + NO ₂	KRO2NO	Rickard (2022)*

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G49239	TrGTerC	NOPINDO2 → C89CO3	k1_R02p0R02	Rickard (2022)
G49240	TrGTerC	NOPINDOOH → NOPINDCO	2.63E-11	Rickard (2022)
G49241	TrGTerC	NOPINDCO + OH → C89CO3	3.07E-12	Rickard (2022)
G49242	TrGTerC	NOPINOO → NOPINONE + H ₂ O ₂	6.00E-18*c(ind_H2O)	Rickard (2022)
G49243	TrGTerC	NOPINOO + CO → NOPINONE + CO ₂	1.2E-15	Rickard (2022)
G49244	TrGTerCN	NOPINOO + NO → NOPINONE + NO ₂	1.E-14	Rickard (2022)
G49245	TrGTerCN	NOPINOO + NO ₂ → NOPINONE + NO ₃	1.E-15	Rickard (2022)
G49246	TrGTerC	NORPINENOL + OH → HCOOH + OH + C86O2	k_CH2CHOH_OH_HCOOH	Sander et al. (2019), So et al. (2014)*
G49247	TrGTerC	NORPINENOL + HCOOH → NORPINAL + HCOOH	k_CH2CHOH_HCOOH	Sander et al. (2019), da Silva (2010)*
G49248	TrGTerC	NORPINAL + HCOOH → NORPINENOL + HCOOH	k_ALD_HCOOH	Sander et al. (2019), da Silva (2010)*
G49249	TrGTerC	C811CO3H + OH → C811CO3	1.04E-11	Rickard (2022)
G49250	TrGTerCN	C811PAN → C811CO3 + NO ₂	k_PAN_M	Rickard (2022)
G49251	TrGTerCN	C811PAN + OH → C721CHO + CO + NO ₂	6.77E-12	Rickard (2022)
G49400a	TrGAroC	LTMB + OH → TLEPOXMUC + HO ₂ + 2 LCARBON	0.827E-11	Rickard (2022)*
G49400b	TrGAroC	LTMB + OH → C6H5CH2O2 + 2 LCARBON	0.189E-11	Rickard (2022)*
G49400c	TrGAroC	LTMB + OH → CRESOL + 2 LCARBON	0.141E-11	Rickard (2022)*
G49400d	TrGAroC	LTMB + OH → TLBIPERO2 + HO ₂ + 2 LCARBON	2.917E-11	Rickard (2022)*
G49401	TrGAroCN	LTMB + NO ₃ → C6H5CH2O2 + HNO ₃ + 2 LCARBON	1.52E-15	Rickard (2022)*
G40200	TrGTerC	APINENE + OH → .75 LAPINABO2 + .15 MENTHEN6ONE + .15 HO ₂ + .10 ROO6R1O2	1.2E-11*EXP(440./temp)	Atkinson et al. (2006)*
G40201a	TrGTerCN	LAPINABO2 + NO → PINAL + HO ₂ + NO ₂	KR02NO*(1.-(.65*alpha_AN(11,3,0,0,0,temp,cair)+.35*alpha_AN(11,2,0,0,0,temp,cair)))	Rickard (2022), Sander et al. (2019)
G40201b	TrGTerCN	LAPINABO2 + NO → LAPINABNO3	KR02NO*(.65*alpha_AN(11,3,0,0,0,temp,cair)+.35*alpha_AN(11,2,0,0,0,temp,cair))	Rickard (2022), Sander et al. (2019)
G40202a	TrGTerC	LAPINABO2 + HO ₂ → LAPINABOOH	k_R02_H02(temp,10)*(1.-r_CHOCH2O2_OH)	Rickard (2022), Sander et al. (2019)
G40202b	TrGTerC	LAPINABO2 + HO ₂ → PINAL + HO ₂ + OH	k_R02_H02(temp,10)*r_CHOCH2O2_OH	Rickard (2022), Sander et al. (2019)
G40203	TrGTerC	LAPINABO2 → PINAL + HO ₂	R02*(0.65*k1_R02t0R02+.35*k1_R02s0R02)	Rickard (2022)*

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G40204	TrGTerC	LAPINABOOH + OH → .35 LAPINABO2 + .65 C96CO3	2.77E-11	Rickard (2022)*
G40205	TrGTerCN	LAPINABNO3 + OH → .35 PINAL + .65 C96CO3 + NO ₂	4.29E-12	Rickard (2022)*
G40206	TrGTerC	MENTHEN6ONE + OH → OHMENTHEN6ONEO2	6.46E-11	Vereecken et al. (2007)*
G40207	TrGTerCN	OHMENTHEN6ONEO2 + NO → 2OHMENTHEN6ONE + HO ₂ + NO ₂	KR02NO	Vereecken et al. (2007)*
G40208	TrGTerC	OHMENTHEN6ONEO2 + HO ₂ → 2OHMENTHEN6ONE	k_RO2_HO2(temp, 10)	Vereecken et al. (2007)
G40209	TrGTerC	OHMENTHEN6ONEO2 → 2OHMENTHEN6ONE + HO ₂	k1_RO2tOR02	Vereecken et al. (2007)
G40210	TrGTerC	2OHMENTHEN6ONE + OH → 10 LCARBON	1E-11	Vereecken et al. (2007)
G40211	TrGTerC	PINAL + OH → .772 C96CO3 + .228 PINALO2	5.2E-12*EXP(600./temp)	Wallington et al. (2018)*
G40212	TrGTerCN	PINAL + NO ₃ → C96CO3 + HNO ₃	2.0E-14	Wallington et al. (2018)*
G40213a	TrGTerC	C96CO3 → C96O2 + CO ₂	k1_RO2RC03*0.9	Rickard (2022)
G40213b	TrGTerC	C96CO3 → PINONIC	k1_RO2RC03*0.1	Rickard (2022)
G40214a	TrGTerC	C96CO3 + HO ₂ → PERPINONIC	KAPH02*r_C03_00H	Rickard (2022)
G40214b	TrGTerC	C96CO3 + HO ₂ → PINONIC + O ₃	KAPH02*r_C03_03	Rickard (2022)
G40214c	TrGTerC	C96CO3 + HO ₂ → C96O2 + OH + CO ₂	KAPH02*r_C03_OH	Rickard (2022)
G40215	TrGTerCN	C96CO3 + NO ₂ → C10PAN2	k_CH3C03_N02	Rickard (2022)
G40216	TrGTerCN	C96CO3 + NO → C96O2 + NO ₂ + CO ₂	KAPNO	Rickard (2022)
G40217	TrGTerCN	C96CO3 + NO ₃ → C96O2 + NO ₂ + CO ₂	KR02N03*1.74	Rickard (2022)
G40218	TrGTerCN	C10PAN2 → C96CO3 + NO ₂	k_PAN_M	Rickard (2022)
G40219	TrGTerCN	C10PAN2 + OH → NORPINAL + CO + NO ₂	3.66E-12	Rickard (2022)
G40220	TrGTerC	PINONIC + OH → C96O2 + CO ₂	6.65E-12	Rickard (2022)
G40221	TrGTerC	PERPINONIC + OH → C96CO3	9.73E-12	Rickard (2022)
G40222	TrGTerC	PINALO2 + HO ₂ → PINALOOH	k_RO2_HO2(temp, 10)	Rickard (2022)
G40223a	TrGTerCN	PINALO2 + NO → C106O2 + NO ₂	KR02NO*(1.-alpha_AN(12,3,0,1,0, temp, cair))	Rickard (2022), Sander et al. (2019)
G40223b	TrGTerCN	PINALO2 + NO → PINALNO3	KR02NO*alpha_AN(12,3,0,1,0, temp, cair)	Rickard (2022), Sander et al. (2019)
G40224	TrGTerC	PINALO2 → C106O2	k1_RO2tR02	Rickard (2022)
G40225	TrGTerC	PINALOOH + OH → PINALO2	2.75E-11	Rickard (2022)
G40226	TrGTerCN	PINALNO3 + OH → CO235C6CHO + CH ₃ COCH ₃ + NO ₂	2.25E-11	Rickard (2022)
G40227	TrGTerC	C106O2 + HO ₂ → C106OOH	k_RO2_HO2(temp, 10)	Rickard (2022)
G40228a	TrGTerCN	C106O2 + NO → C716O2 + CH ₃ COCH ₃ + NO ₂	KR02NO*0.875*(1.-alpha_AN(13,3,0, 0,0,temp, cair))	Rickard (2022), Sander et al. (2019)

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G40228b	TrGTerCN	C106O2 + NO → C106NO3	KR02NO*0.875*alpha_AN(13,3,0,0, 0, temp, cair)	Rickard (2022), Sander et al. (2019)
G40229	TrGTerC	C106O2 → C716O2 + CH ₃ COCH ₃	k1_R02tR02	Rickard (2022)
G40230	TrGTerC	C106OOH + OH → C106O2	8.01E-11	Rickard (2022)
G40231	TrGTerCN	C106NO3 + OH → CO235C6CHO + CH ₃ COCH ₃ + NO ₂	7.03E-11	Rickard (2022)
G40232	TrGTerC	APINENE + O ₃ → .09 APINBOO + .08 PINONIC + .77 OH + .33 NORPINAL + .33 CO + .33 HO ₂ + .06 APINAOO + .44 C109O2	8.05E-16*EXP(-640./temp)	Wallington et al. (2018)*
G40233	TrGTerC	APINAOO → PINAL + H ₂ O ₂	1.00E-17*c(ind_H2O)	Rickard (2022)
G40234	TrGTerC	APINAOO + CO → PINAL + CO ₂	1.20E-15	Rickard (2022)
G40235	TrGTerCN	APINAOO + NO → PINAL + NO ₂	1.00E-14	Rickard (2022)
G40236	TrGTerCN	APINAOO + NO ₂ → PINAL + NO ₃	1.00E-15	Rickard (2022)
G40237a	TrGTerC	APINBOO → PINONIC	1.00E-17*c(ind_H2O)*(0.08+0.15)	Rickard (2022)
G40237b	TrGTerC	APINBOO → PINAL + H ₂ O ₂	1.00E-17*c(ind_H2O)*0.77	Rickard (2022)
G40238	TrGTerC	APINBOO + CO → PINAL + CO ₂	1.20E-15	Rickard (2022)
G40239	TrGTerCN	APINBOO + NO → PINAL + NO ₂	1.00E-14	Rickard (2022)
G40240	TrGTerCN	APINBOO + NO ₂ → PINAL + NO ₃	1.00E-15	Rickard (2022)
G40241	TrGTerC	C109O2 → C89CO3 + HCHO	k1_R02p0R02	Rickard (2022)
G40242	TrGTerCN	C109O2 + NO → C89CO3 + HCHO + NO ₂	KR02NO	Rickard (2022)*
G40243a	TrGTerC	C109O2 + HO ₂ → C109OOH	k_R02_H02(temp,10)*r_COCH202_00H	Rickard (2022), Sander et al. (2019)
G40243b	TrGTerC	C109O2 + HO ₂ → C89CO3 + HCHO + OH	k_R02_H02(temp,10)*r_COCH202_OH	Rickard (2022), Sander et al. (2019)
G40244	TrGTerC	C109OOH + OH → C109CO + OH	5.47E-11	Rickard (2022)
G40245	TrGTerC	C109CO + OH → C89CO3 + CO	5.47E-11	Rickard (2022)
G40246	TrGTerCN	APINENE + NO ₃ → LNAPINABO2	1.2E-12*EXP(490./temp)	Wallington et al. (2018)*
G40247	TrGTerCN	LNAPINABO2 → PINAL + NO ₂	(0.65*k1_R02tR02 + 0.35*k1_R02sR02)	Rickard (2022)
G40248	TrGTerCN	LNAPINABO2 + NO → PINAL + NO ₂ + NO ₂	KR02NO	Rickard (2022)*
G40249	TrGTerCN	LNAPINABO2 + HO ₂ → LNAPINABOOH	k_R02_H02(temp,10)	Rickard (2022)
G40250	TrGTerCN	LNAPINABO2 + NO ₃ → PINAL + NO ₂ + NO ₂	KR02N03	Rickard (2022)
G40251	TrGTerCN	LNAPINABOOH + OH → LNAPINABO2	(.65*6.87E-12+.35*1.23E-11)	Rickard (2022)
G40252a	TrGTerC	BPINENE + OH → BPINAO2	1.47E-11*EXP(467./temp) *(0.8326*0.3+0.068)/(0.8326+0.068)	Gill and Hites (2002)*

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G40252b	TrGTerC	BPINENE + OH → ROO6R1O2	1.47E-11*EXP(467./temp) *0.8326*0.7/(0.8326+0.068)	Gill and Hites (2002)*
G40253a	TrGTerC	BPINAO2 + HO ₂ → BPINAOOH	k_RO2_HO2(temp,10)*r_COCH2O2_0OH	Rickard (2022), Sander et al. (2019)
G40253b	TrGTerC	BPINAO2 + HO ₂ → NOPINONE + HCHO + HO ₂ + OH	k_RO2_HO2(temp,10)*r_COCH2O2_OH	Rickard (2022), Sander et al. (2019)
G40254a	TrGTerCN	BPINAO2 + NO → NOPINONE + HCHO + HO ₂ + NO ₂	KRO2NO*(1.-alpha_AN(11,3,0,0,0, temp,cair))	Rickard (2022), Sander et al. (2019)
G40254b	TrGTerCN	BPINAO2 + NO → BPINANO3	KRO2NO*alpha_AN(11,3,0,0,0, temp,cair)	Rickard (2022), Sander et al. (2019)
G40255	TrGTerC	BPINAO2 → NOPINONE + HCHO + HO ₂	k1_RO2tORO2	Rickard (2022)
G40256	TrGTerC	BPINAOOH + OH → BPINAO2	1.33E-11	Rickard (2022)
G40257	TrGTerCN	BPINANO3 + OH → NOPINONE + HCHO + NO ₂	4.70E-12	Rickard (2022)
G40258a	TrGTerCN	ROO6R1O2 + NO → ROO6R3O2 + CH ₃ COCH ₃ + NO ₂	KRO2NO*(1.-alpha_AN(13,3,0,0,0, temp,cair))	Vereecken and Peeters (2012)
G40258b	TrGTerCN	ROO6R1O2 + NO → ROO6R1NO3	KRO2NO*alpha_AN(13,3,0,0,0, temp,cair)	Vereecken and Peeters (2012)
G40259	TrGTerC	ROO6R1O2 + HO ₂ → 10 LCARBON	k_RO2_HO2(temp,10)	Vereecken and Peeters (2012)*
G40260	TrGTerC	ROO6R1O2 → ROO6R3O2 + CH ₃ COCH ₃	k1_RO2tORO2	Vereecken and Peeters (2012)
G40261a	TrGTerCN	RO6R1O2 + NO → RO6R3O2 + NO ₂	KRO2NO*(1.-alpha_AN(12,3,0,0,0, temp,cair))	Vereecken and Peeters (2012)
G40261b	TrGTerCN	RO6R1O2 + NO → RO6R1NO3	KRO2NO*alpha_AN(12,3,0,0,0, temp,cair)	Vereecken and Peeters (2012)
G40262	TrGTerC	RO6R1O2 + HO ₂ → 10 LCARBON	k_RO2_HO2(temp,10)	Vereecken and Peeters (2012)*
G40263	TrGTerC	RO6R1O2 → RO6R3O2	k1_RO2sORO2	Vereecken and Peeters (2012)
G40264a	TrGTerCN	RO6R3O2 + NO → 9 LCARBON + HCHO + HO ₂ + NO ₂	KRO2NO*(1.-alpha_AN(12,3,0,0,0, temp,cair))	Vereecken and Peeters (2012)
G40264b	TrGTerCN	RO6R3O2 + NO → 10 LCARBON + LNITROGEN	KRO2NO*alpha_AN(12,3,0,0,0, temp,cair)	Vereecken and Peeters (2012)
G40265	TrGTerC	RO6R3O2 + HO ₂ → 10 LCARBON	k_RO2_HO2(temp,10)	Vereecken and Peeters (2012)
G40266	TrGTerC	RO6R3O2 → 9 LCARBON + HCHO + HO ₂	k1_RO2sRO2	Vereecken and Peeters (2012)*
G40267a	TrGTerC	BPINENE + O ₃ → NOPINONE + .63 CO + .37 CH ₂ OO +.16 OH + .16 HO ₂	1.35E-15*EXP(-1270./temp) *.051/(1.-.027)	Wallington et al. (2018)*
G40267b	TrGTerC	BPINENE + O ₃ → NOPINOO + CO ₂	1.35E-15*EXP(-1270./temp) *.368/(1.-.027)	Nguyen et al. (2009), Wallington et al. (2018)

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G40267c	TrGTerC	BPINENE + O ₃ → NOPINDO ₂ + CO ₂ + OH	1.35E-15*EXP(-1270./temp) *.283/(1.-.027)	Nguyen et al. (2009), Wallington et al. (2018)
G40267d	TrGTerC	BPINENE + O ₃ → C8BC + 2 CO ₂	1.35E-15*EXP(-1270./temp) *(.104+.167)/(1.-.027)	Nguyen et al. (2009), Wallington et al. (2018)
G40268	TrGTerCN	BPINENE + NO ₃ → LNPINABO ₂	2.51E-12	Wallington et al. (2018)*
G40269	TrGTerCN	LNPINABO ₂ + HO ₂ → LNPINABOOH	k_RO2_HO2(temp, 10)	Rickard (2022)
G40270	TrGTerCN	LNPINABO ₂ + NO → NOPINONE + HCHO + NO ₂ + NO ₂	KRO2NO	Rickard (2022)*
G40271	TrGTerCN	LNPINABO ₂ + NO ₃ → NOPINONE + HCHO + NO ₂ + NO ₂	KRO2NO3	Rickard (2022)
G40272a	TrGTerCN	LNPINABO ₂ → NOPINONE + HCHO + NO ₂	k1_RO2tR02*0.7	Rickard (2022)
G40272b	TrGTerCN	LNPINABO ₂ → BPINANO ₃	k1_RO2tR02*0.3	Rickard (2022)
G40273	TrGTerCN	LNPINABOOH + OH → LNPINABO ₂	9.58E-12	Rickard (2022)
G40274	TrGTerCN	ROO6R1NO ₃ + OH → ROO6R3O ₂ + CH ₃ COCH ₃ + NO ₂	9.16E-13	Vereecken and Peeters (2012), Gill and Hites (2002)*
G40275	TrGTerCN	RO6R1NO ₃ + OH → 9 LCARBON + HCHO + HO ₂ + NO ₂	9.16E-13	Vereecken and Peeters (2012), Gill and Hites (2002)
G40276	TrGTerC	PINEOL + OH → HCOOH + OH + NORPINAL	k_CH2CHOH_OH_HCOOH	Sander et al. (2019), So et al. (2014)*
G40277	TrGTerC	PINEOL + HCOOH → PINAL + HCOOH	k_CH2CHOH_HCOOH	Sander et al. (2019), da Silva (2010)*
G40278	TrGTerC	PINAL + HCOOH → PINEOL + HCOOH	k_ALD_HCOOH	Sander et al. (2019), da Silva (2010)*
G40279a	TrGC	CARENE + OH → LAPINABO ₂	8.8E-11*(.50+.25)	Atkinson and Arey (2003)
G40279b	TrGC	CARENE + OH → MENTHEN6ONE + HO ₂	8.8E-11*.25*.60	Atkinson and Arey (2003)
G40279c	TrGC	CARENE + OH → ROO6R1O ₂	8.8E-11*.25*.40	Atkinson and Arey (2003)
G40280a	TrGC	CARENE + O ₃ → APINBOO	3.7E-17*.50*.18	Atkinson and Arey (2003)
G40280b	TrGC	CARENE + O ₃ → PINONIC	3.7E-17*.50*.16	Atkinson and Arey (2003)
G40280c	TrGC	CARENE + O ₃ → OH + NORPINAL + CO + HO ₂	3.7E-17*.50*.66	Atkinson and Arey (2003)
G40280d	TrGC	CARENE + O ₃ → APINAOO	3.7E-17*.50*.12	Atkinson and Arey (2003)
G40280e	TrGC	CARENE + O ₃ → OH + C109O ₂	3.7E-17*.50*(.22+.66)	Atkinson and Arey (2003)
G40281	TrGCN	CARENE + NO ₃ → LNAPINABO ₂	9.1E-12	Atkinson and Arey (2003)
G40282a	TrGTerC	SABINENE + OH → BPINA02	1.47E-11*EXP(467./temp) *(0.8326*0.3+0.068)/(0.8326+0.068)	Gill and Hites (2002)*

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G40282b	TrGTerC	SABINENE + OH → ROO6R1O2	1.47E-11*EXP(467./temp) *0.8326*0.7/(0.8326+0.068)	Vereecken and Peeters (2012), Gill and Hites (2002)*
G40283a	TrGTerC	SABINENE + O ₃ → NOPINONE + .63 CO + .37 HOCH ₂ OOH + .16 OH + .16 HO ₂	1.35E-15*EXP(-1270./temp) *.051/(1.-.027)	Wallington et al. (2018)*
G40283b	TrGTerC	SABINENE + O ₃ → NOPINOO + CO ₂	1.35E-15*EXP(-1270./temp) *.368/(1.-.027)	Nguyen et al. (2009), Wallington et al. (2018)
G40283c	TrGTerC	SABINENE + O ₃ → NOPINDO2 + CO ₂ + OH	1.35E-15*EXP(-1270./temp) *.283/(1.-.027)	Nguyen et al. (2009), Wallington et al. (2018)
G40283d	TrGTerC	SABINENE + O ₃ → C8BC + 2 CO ₂	1.35E-15*EXP(-1270./temp) *(.104+.167)/(1.-.027)	Nguyen et al. (2009), Wallington et al. (2018)
G40284	TrGTerCN	SABINENE + NO ₃ → LNBPINABO2	2.51E-12	Wallington et al. (2018)*
G40285a	TrGTerC	CAMPHENENE + OH → BPINAO2	1.47E-11*EXP(467./temp) *(0.8326*0.3+0.068)/(0.8326+0.068)	Gill and Hites (2002)*
G40285b	TrGTerC	CAMPHENENE + OH → ROO6R1O2	1.47E-11*EXP(467./temp) *0.8326*0.7/(0.8326+0.068)	Vereecken and Peeters (2012), Gill and Hites (2002)*
G40286a	TrGTerC	CAMPHENENE + O ₃ → NOPINONE + .63 CO + .37 HOCH ₂ OOH + .16 OH + .16 HO ₂	1.35E-15*EXP(-1270./temp) *.051/(1.-.027)	Wallington et al. (2018)*
G40286b	TrGTerC	CAMPHENENE + O ₃ → NOPINOO + CO ₂	1.35E-15*EXP(-1270./temp) *.368/(1.-.027)	Nguyen et al. (2009), Wallington et al. (2018)
G40286c	TrGTerC	CAMPHENENE + O ₃ → NOPINDO2 + CO ₂ + OH	1.35E-15*EXP(-1270./temp) *.283/(1.-.027)	Nguyen et al. (2009), Wallington et al. (2018)
G40286d	TrGTerC	CAMPHENENE + O ₃ → C8BC + 2 CO ₂	1.35E-15*EXP(-1270./temp) *(.104+.167)/(1.-.027)	Nguyen et al. (2009), Wallington et al. (2018)
G40287	TrGTerCN	CAMPHENENE + NO ₃ → LNBPINABO2	2.51E-12	Wallington et al. (2018)*
G40400	TrGAroC	LHAROM + OH → .14 TLEPOXMUC + .03 C ₆ H ₅ CH ₂ O ₂ + .04 CRESOL + .79 TLBIPERO2 + .18 HO ₂ + 4 LCARBON	5.67E-11	Rickard (2022)*
G40401	TrGAroCN	LHAROM + NO ₃ → C ₆ H ₅ CH ₂ O ₂ + HNO ₃ + 4 LCARBON	2.60E-15	Rickard (2022)*
G6100	UpStTrGCl	Cl + O ₃ → ClO + O ₂	2.8E-11*EXP(-250./temp)	Atkinson et al. (2007)
G6102a	StTrGCl	ClO + ClO → Cl ₂ + O ₂	1.0E-12*EXP(-1590./temp)	Atkinson et al. (2007)
G6102b	StTrGCl	ClO + ClO → 2 Cl + O ₂	3.0E-11*EXP(-2450./temp)	Atkinson et al. (2007)
G6102c	StTrGCl	ClO + ClO → Cl + OCLO	3.5E-13*EXP(-1370./temp)	Atkinson et al. (2007)
G6102d	StTrGCl	ClO + ClO → Cl ₂ O ₂	k_ClO_ClO	Burkholder et al. (2015)

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G6103	StTrGCl	$\text{Cl}_2\text{O}_2 \rightarrow \text{ClO} + \text{ClO}$	$k_{\text{ClO}}/\text{ClO}/(2.16\text{E}-27*\text{EXP}(8537./\text{temp}))$	Burkholder et al. (2015)*
G69MS	StTrGCl	$\text{ClO} + \text{OCLO} \rightarrow \text{Cl}_2\text{O}_3$	1.2E-12	Atkinson et al. (2007)
G699MS	StTrGCl	$\text{Cl} + \text{Cl}_2\text{O} \rightarrow \text{Cl}_2 + \text{ClO}$	$6.2\text{E}-11*\text{EXP}(130./\text{temp})$	Atkinson et al. (2007)
G6202	StTrGCl	$\text{Cl} + \text{H}_2\text{O}_2 \rightarrow \text{HCl} + \text{HO}_2$	$1.1\text{E}-11*\text{EXP}(-980./\text{temp})$	Atkinson et al. (2007)
G6204	StTrGCl	$\text{ClO} + \text{HO}_2 \rightarrow \text{HOCl} + \text{O}_2$	$2.2\text{E}-12*\text{EXP}(340./\text{temp})$	Atkinson et al. (2007)*
G6205	StTrGCl	$\text{HCl} + \text{OH} \rightarrow \text{Cl} + \text{H}_2\text{O}$	$1.7\text{E}-12*\text{EXP}(-230./\text{temp})$	Atkinson et al. (2007)
G67MS	StTrGCl	$\text{Cl}_2 + \text{OH} \rightarrow \text{HOCl} + \text{Cl}$	$3.6\text{E}-12*\text{EXP}(-1200./\text{temp})$	Atkinson et al. (2007)
G6300	UpStTrGClN	$\text{ClO} + \text{NO} \rightarrow \text{NO}_2 + \text{Cl}$	$6.2\text{E}-12*\text{EXP}(295./\text{temp})$	Atkinson et al. (2007)
G6301	StTrGClN	$\text{ClO} + \text{NO}_2 \rightarrow \text{ClNO}_3$	$k_{\text{3rd_iupac}}(\text{temp}, \text{cair}, 1.6\text{E}-31, 3.4, 7.\text{E}-11, 0., 0.4)$	Atkinson et al. (2007)
G6302	TrGClN	$\text{ClNO}_3 \rightarrow \text{ClO} + \text{NO}_2$	$6.918\text{E}-7*\text{EXP}(-10909./\text{temp})*\text{cair}$	Anderson and Fahey (1990)
G6304	StTrGClN	$\text{ClNO}_3 + \text{Cl} \rightarrow \text{Cl}_2 + \text{NO}_3$	$6.2\text{E}-12*\text{EXP}(145./\text{temp})$	Atkinson et al. (2007)
G63MS	UpStTrGClN	$\text{Cl} + \text{NO} \rightarrow \text{CINO}$	$7.7\text{E}-32*(\text{temp}/298.)**(-1.80)*\text{cair}$	Burkholder et al. (2019)
G64MS	UpStTrGClN	$\text{Cl} + \text{NO}_2 \rightarrow \text{ClONO}$	1.6E-11	Burkholder et al. (2015)
G65MS	UpStTrGClN	$\text{Cl} + \text{NO}_2 \rightarrow \text{ClNO}_2$	3.6E-12	Burkholder et al. (2015)
G66MS	UpStTrGClN	$\text{Cl} + \text{NO}_3 \rightarrow \text{ClO} + \text{NO}_2$	2.4E-11	Burkholder et al. (2019)
G68MS	StTrGClN	$\text{CINO}_2 + \text{OH} \rightarrow \text{HOCl} + \text{NO}_2$	$2.4\text{E}-12*\text{EXP}(-1250./\text{temp})$	Atkinson et al. (2007)
G688MS	UpStTrGClN	$\text{OCLO} + \text{NO} \rightarrow \text{NO}_2 + \text{ClO}$	$1.1\text{E}-13*\text{EXP}(350./\text{temp})$	Atkinson et al. (2007)
G689MS	UpStTrGClN	$\text{ClONO} \rightarrow \text{ClNO}_2$	2.3E-5	Janowski et al. (1977)
G6400	StTrGCl	$\text{Cl} + \text{CH}_4 \rightarrow \text{HCl} + \text{CH}_3$	$6.6\text{E}-12*\text{EXP}(-1240./\text{temp})$	Atkinson et al. (2006)
G6400MS	TrGCCl	$\text{Cl} + \text{C}_3\text{H}_8 \rightarrow \text{iC}_3\text{H}_7\text{O}_2 + \text{HCl}$	$1.4\text{E}-10*0.43*\text{EXP}(75./\text{temp})$	Rickard (2022)
G6401MS	TrGCCl	$\text{Cl} + \text{C}_3\text{H}_8 \rightarrow \text{C}_3\text{H}_7\text{O}_2 + \text{HCl}$	$1.4\text{E}-10*0.59*\text{EXP}(-90./\text{temp})$	Rickard (2022)
G6402MS	TrGCCl	$\text{Cl} + \text{iC}_4\text{H}_{10} \rightarrow \text{IC}_4\text{H}_9\text{O}_2 + \text{HCl}$	1.43E-10*0.564	Rickard (2022)
G6403MS	TrGCCl	$\text{Cl} + \text{iC}_4\text{H}_{10} \rightarrow \text{TC}_4\text{H}_9\text{O}_2 + \text{HCl}$	1.43E-10*0.436	Rickard (2022)
G6404MS	TrGCCl	$\text{Cl} + \text{C}_4\text{H}_{10} \rightarrow \text{LC}_4\text{H}_9\text{O}_2 + \text{HCl}$	2.05E-10	Atkinson et al. (2006)
G6401	StTrGCl	$\text{Cl} + \text{HCHO} \rightarrow \text{HCl} + \text{CO} + \text{HO}_2$	$8.1\text{E}-11*\text{EXP}(-34./\text{temp})$	Atkinson et al. (2006)
G6402	StTrGCl	$\text{Cl} + \text{CH}_3\text{OOH} \rightarrow \text{HCHO} + \text{HCl} + \text{OH}$	5.9E-11	Atkinson et al. (2006)*
G6403	StTrGCl	$\text{ClO} + \text{CH}_3\text{O}_2 \rightarrow \text{HO}_2 + \text{Cl} + \text{HCHO}$	$1.8\text{E}-12*\text{EXP}(-600./\text{temp})$	Burkholder et al. (2015)
G6408	StTrGCCl	$\text{CH}_3\text{CCl}_3 + \text{OH} \rightarrow 2 \text{LCARBON} + \text{H}_2\text{O} + 3 \text{Cl}$	$1.64\text{E}-12*\text{EXP}(-1520./\text{temp})$	Burkholder et al. (2015)
G6409	TrGCCl	$\text{Cl} + \text{C}_2\text{H}_4 \rightarrow \text{HOCH}_2\text{CH}_2\text{O}_2 + \text{HCl}$	$k_{\text{3rd_iupac}}(\text{temp}, \text{cair}, 1.85\text{E}-29, 3.3, 6.0\text{E}-10, 0.0, 0.4)$	Atkinson et al. (2006)*
G6410	TrGCCl	$\text{Cl} + \text{CH}_3\text{CHO} \rightarrow \text{HCl} + \text{CH}_3\text{C(O)}$	8.0e-11	Atkinson et al. (2006)
G6411	TrGCCl	$\text{C}_2\text{H}_2 + \text{Cl} \rightarrow \text{LCARBON} + \text{CH}_3 + \text{HCl}$	$k_{\text{3rd_iupac}}(\text{temp}, \text{cair}, 6.1\text{e}-30, 3.0, 2.0\text{e}-10, 0., 0.6)$	Atkinson et al. (2006)

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G6412	TrGCCl	$\text{C}_2\text{H}_6 + \text{Cl} \rightarrow \text{C}_2\text{H}_5\text{O}_2 + \text{HCl}$	8.3E-11*EXP(-100./temp)	Atkinson et al. (2006)
G6413	StTrGCIN	$\text{Cl} + \text{CH}_3\text{ONO}_2 \rightarrow \text{HCl} + \text{HCHO} + \text{NO}_2$	1.3E-11*EXP(-1200./temp)	Burkholder et al. (2015)
G6414	StTrGCIN	$\text{Cl} + \text{CH}_3\text{ONO} \rightarrow \text{HCl} + \text{HCHO} + \text{NO}$	2.1E-12	Sokolov et al. (1999)
G6415	StTrGCl	$\text{Cl} + \text{CH}_3\text{O}_2 \rightarrow .5 \text{ ClO} + .5 \text{ CH}_3\text{O} + .5 \text{ HCl} + .5 \text{ CH}_2\text{OO}$	1.6E-10	Burkholder et al. (2015)
G6416	TrGCClN	$\text{Cl} + \text{CH}_3\text{CN} \rightarrow \text{NCCH}_2\text{O}_2 + \text{HCl}$	1.6E-11*EXP(-2104./temp)	Tyndall et al. (1996), Tyndall et al. (2001b), Sander et al. (2019)
G641MS	StTrGCCl	$\text{Cl} + \text{BENZENE} \rightarrow \text{C}_6\text{H}_5\text{O}_2 + \text{HCl}$	1.3E-16	Sokolov et al. (1998)
G642MS	StTrGCCl	$\text{Cl} + \text{TOLUENE} \rightarrow \text{C}_6\text{H}_5\text{CH}_2\text{O}_2 + \text{HCl}$	6.2E-11	Wang et al. (2005)
G692MS	StTrGCCl	$\text{Cl} + \text{LXYL} \rightarrow \text{C}_6\text{H}_5\text{CH}_2\text{O}_2 + \text{LCARBON} + \text{HCl}$	1.5E-10	Shi and Bernhard (1997)
G643DT	TrGCCl	$\text{Cl} + \text{C}_5\text{H}_8 \rightarrow .63 \text{ LISOPAB} + .30 \text{ LISOPCD} + .07 \text{ LISOPEFO}_2 + \text{HCl}$	7.6E-11*EXP(500./temp) * 1.1*EXP(-595./temp)	Wennberg et al. (2018)*
G644DT	TrGCCl	$\text{Cl} + \text{C}_5\text{H}_8 \rightarrow .63 \text{ LISOPAB} + .30 \text{ LISOPCD} + .07 \text{ LISOPEFO}_2 + \text{LCHLORINE}$	7.6E-11*EXP(500./temp) * (1.-1.1*EXP(-595./temp))	Wennberg et al. (2018)*
G645DT	StTrGCl	$\text{Cl} + \text{CH}_3\text{OH} \rightarrow \text{HOCH}_2\text{O}_2 + \text{HCl}$	7.1E-11*EXP(-75./temp)	Atkinson et al. (2006)
G646DT	StTrGCCl	$\text{Cl} + \text{C}_2\text{H}_5\text{OH} \rightarrow \text{HOCH}_2\text{CH}_2\text{O}_2 + \text{HCl}$	6.0E-11*EXP(155./temp) * 0.28*EXP(-350./temp)	Atkinson et al. (2006)
G647DT	StTrGCCl	$\text{Cl} + \text{C}_2\text{H}_5\text{OH} \rightarrow \text{C}_2\text{H}_5\text{O}_2 + \text{HCl}$	6.0E-11*EXP(155./temp) * (1. - 0.28*EXP(-350./temp))	Atkinson et al. (2006)
G648DT	StTrGCCl	$\text{Cl} + \text{HOCH}_2\text{CHO} \rightarrow \text{HOCHCHO} + \text{HCl}$	8.0E-12/0.9 *0.35	Niki et al. (1987), Atkinson et al. (2006)
G649DT	StTrGCCl	$\text{Cl} + \text{HOCH}_2\text{CHO} \rightarrow \text{HOCH}_2\text{CO} + \text{HCl}$	8.0E-12/0.9 *(1.-0.35)	Niki et al. (1987), Atkinson et al. (2006)
G650DT	StTrGCCl	$\text{Cl} + \text{GLYOX} \rightarrow \text{HCOCO} + \text{HCl}$	3.8E-11	Niki et al. (1985)
G651DT	StTrGCCl	$\text{Cl} + \text{MGLYOX} \rightarrow \text{CH}_3\text{C(O)} + \text{CO} + \text{HCl}$	4.8E-11	Green et al. (1990)
G652DT	StTrGCCl	$\text{Cl} + \text{C}_2\text{H}_5\text{CHO} \rightarrow \text{C}_2\text{H}_5\text{CO}_3 + \text{HCl}$	1.3E-10	Atkinson et al. (2006)*
G653DT	StTrGCCl	$\text{Cl} + \text{CH}_3\text{COCH}_3 \rightarrow \text{CH}_3\text{COCH}_2\text{O}_2 + \text{HCl}$	1.5E-11*EXP(-590./temp)	Atkinson et al. (2006)
G654DT	StTrGCCl	$\text{Cl} + \text{MEK} \rightarrow \text{LMEKO}_2 + \text{HCl}$	3.05E-11*EXP(80./temp)	Atkinson et al. (2006)*
G655MS	StTrGCCl	$\text{Cl} + \text{BENZAL} \rightarrow \text{C}_6\text{H}_5\text{CO}_3 + \text{HCl}$	1.0E-10	Thiault et al. (2002)
G7100	StTrGBr	$\text{Br} + \text{O}_3 \rightarrow \text{BrO} + \text{O}_2$	1.7E-11*EXP(-800./temp)	Atkinson et al. (2007)
G7102a	StTrGBr	$\text{BrO} + \text{BrO} \rightarrow 2 \text{ Br} + \text{O}_2$	2.7E-12	Atkinson et al. (2007)
G7102b	StTrGBr	$\text{BrO} + \text{BrO} \rightarrow \text{Br}_2 + \text{O}_2$	2.9E-14*EXP(840./temp)	Atkinson et al. (2007)
G7200	StTrGBr	$\text{Br} + \text{HO}_2 \rightarrow \text{HBr} + \text{O}_2$	7.7E-12*EXP(-450./temp)	Atkinson et al. (2007)
G7201	StTrGBr	$\text{BrO} + \text{HO}_2 \rightarrow \text{HOBr} + \text{O}_2$	4.5E-12*EXP(500./temp)	Atkinson et al. (2007)

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G7202	StTrGBr	HBr + OH → Br + H ₂ O	6.7E-12*EXP(155./temp)	Atkinson et al. (2007)
G7204	StTrGBr	Br ₂ + OH → HOBr + Br	2.0E-11*EXP(240./temp)	Atkinson et al. (2007)
G7300	TrGBrN	Br + BrNO ₃ → Br ₂ + NO ₃	4.9E-11	Orlando and Tyndall (1996)
G7301	StTrGBrN	BrO + NO → Br + NO ₂	8.7E-12*EXP(260./temp)	Atkinson et al. (2007)
G7302	StTrGBrN	BrO + NO ₂ → BrNO ₃	k_BrO_NO2	Atkinson et al. (2007)*
G7303	TrGBrN	BrNO ₃ → BrO + NO ₂	k_BrO_NO2/(5.44E-9*EXP(14192./temp)*1.E6*R_gas*temp/(atm2Pa*N_A))	Orlando and Tyndall (1996), Atkinson et al. (2007)*
G7400	StTrGBr	Br + HCHO → HBr + CO + HO ₂	7.7E-12*EXP(-580./temp)	Atkinson et al. (2006)
G7401	TrGBr	Br + CH ₃ OOH → CH ₃ O ₂ + HBr	2.6E-12*EXP(-1600./temp)	Kondo and Benson (1984)
G7402	TrGBr	BrO + CH ₃ O ₂ → HOBr + CH ₂ OO	2.42E-14*EXP(1617./temp)	Shalcross et al. (2015)
G7403	StTrGBr	CH ₃ Br + OH → LCARBON + H ₂ O + Br	1.42E-12*EXP(-1150./temp)	Burkholder et al. (2015)
G7404	TrGBrC	Br + C ₂ H ₄ → HOCH ₂ CH ₂ O ₂ + HBr	2.8E-13*EXP(224./temp)/(1.+1.13E24*EXP(-3200./temp)/C(ind_02))	Atkinson et al. (2006)*
G7405	TrGBrC	Br + CH ₃ CHO → HBr + CH ₃ C(O)	1.8e-11*EXP(-460./temp)	Atkinson et al. (2006)
G7406	TrGBrC	Br + C ₂ H ₂ → LCARBON + CH ₃ O ₂ + HBr	6.35e-15*EXP(440./temp)	Atkinson et al. (2006)
G7407	TrGBr	CHBr ₃ + OH → LCARBON + H ₂ O + 3 Br	9.0E-13*EXP(-360./temp)	Burkholder et al. (2015)*
G7408	TrGBr	CH ₂ Br ₂ + OH → LCARBON + H ₂ O + 2 Br	2.0E-12*EXP(-840./temp)	Burkholder et al. (2015)*
G7600	TrGBrCl	Br + BrCl → Br ₂ + Cl	3.32E-15	Manion et al. (2015)
G7601	TrGBrCl	Br + Cl ₂ → BrCl + Cl	1.10E-15	Dolson and Leone (1987)
G7602	TrGBrCl	Br ₂ + Cl → BrCl + Br	2.3E-10*EXP(135./temp)	Bedjanian et al. (1998)
G7603a	StTrGBrCl	BrO + ClO → Br + OClO	1.6E-12*EXP(430./temp)	Atkinson et al. (2007)
G7603b	StTrGBrCl	BrO + ClO → Br + Cl + O ₂	2.9E-12*EXP(220./temp)	Atkinson et al. (2007)
G7603c	StTrGBrCl	BrO + ClO → BrCl + O ₂	5.8E-13*EXP(170./temp)	Atkinson et al. (2007)
G7604	TrGBrCl	BrCl + Cl → Br + Cl ₂	1.45E-11	Clyne and Cruse (1972)
G7605	TrGBrCl	CHCl ₂ Br + OH → LCARBON + 2 Cl + H ₂ O + Br	2.0E-12*EXP(-840./temp)	see note*
G7606	TrGBrCl	CHClBr ₂ + OH → LCARBON + Cl + H ₂ O + 2 Br	2.0E-12*EXP(-840./temp)	see note*
G7607	TrGBrCl	CH ₂ ClBr + OH → LCARBON + Cl + H ₂ O + Br	2.1E-12*EXP(-880./temp)	Burkholder et al. (2015)*
G9200	StTrGS	SO ₂ + OH → H ₂ SO ₄ + HO ₂	k_3rd(temp, cair, 3.3E-31, 4.3, 1.6E-12, 0., 0.6)	Burkholder et al. (2015)
G9400a	TrGCS	DMS + OH → CH ₃ SO ₂ + HCHO	1.13E-11*EXP(-253./temp)	Atkinson et al. (2004)*
G9400b	TrGCS	DMS + OH → DMSO + HO ₂	k_DMS_OH	Atkinson et al. (2004)*
G9401	TrGCNS	DMS + NO ₃ → CH ₃ SO ₂ + HNO ₃ + HCHO	1.9E-13*EXP(520./temp)	Atkinson et al. (2004)

Table 1: Gas phase reactions (... continued)

#	labels	reaction	rate coefficient	reference
G9402	TrGCS	DMSO + OH → .6 SO ₂ + HCHO + .6 CH ₃ + .4 HO ₂ + .4 CH ₃ SO ₃ H	1.E-10	Hynes and Wine (1996)*
G9403	TrGS	CH ₃ SO ₂ → SO ₂ + CH ₃	1.8E13*EXP(-8661./temp)	Barone et al. (1995)
G9404	TrGS	CH ₃ SO ₂ + O ₃ → CH ₃ SO ₃	3.E-13	Barone et al. (1995)
G9405	TrGS	CH ₃ SO ₃ + HO ₂ → CH ₃ SO ₃ H	5.E-11	Barone et al. (1995)
G9408	StTrGS	CH ₂ OO + SO ₂ → H ₂ SO ₄ + HCHO	k_CH2OO_SO2	Welz et al. (2012), Stone et al. (2014)*
G9409	TrGTerCS	NOPINOO + SO ₂ → NOPINONE + H ₂ SO ₄	7.E-14	Rickard (2022)
G9410	TrGTerCS	APINAOO + SO ₂ → PINAL + H ₂ SO ₄	7.00E-14	Rickard (2022)
G9411	TrGTerCS	APINBOO + SO ₂ → PINAL + H ₂ SO ₄	7.00E-14	Rickard (2022)
G9412	TrGTerCS	MBOOO + SO ₂ → IBUTALOH + H ₂ SO ₄	7.00E-14	Rickard (2022)
G9600	TrGCCIS	DMS + Cl → CH ₃ SO ₂ + HCl + HCHO	3.3E-10	Atkinson et al. (2004)
G9700	TrGBrCS	DMS + Br → CH ₃ SO ₂ + HBr + HCHO	9.E-11*EXP(-2386./temp)	Jefferson et al. (1994)
G9701	TrGBrCS	DMS + BrO → DMSO + Br	4.4E-13	Ingham et al. (1999)

General notes

Three-body reactions

Rate coefficients for three-body reactions are defined via the function `k_3rd(T, M, k0300, n, kinf300, m, fc)`. In the code, the temperature T is called `temp` and the concentration of “air molecules” M is called `cair`. Using the auxiliary variables $k_0(T)$, $k_{\text{inf}}(T)$, and k_{ratio} , `k_3rd` is defined as:

$$k_0(T) = k_0^{300} \times \left(\frac{300\text{K}}{T}\right)^n \quad (1)$$

$$k_{\text{inf}}(T) = k_{\text{inf}}^{300} \times \left(\frac{300\text{K}}{T}\right)^m \quad (2)$$

$$k_{\text{ratio}} = \frac{k_0(T)M}{k_{\text{inf}}(T)} \quad (3)$$

$$\text{k_3rd} = \frac{k_0(T)M}{1 + k_{\text{ratio}}} \times f_c^{\left(\frac{1}{1 + (\log_{10}(k_{\text{ratio}}))^2}\right)} \quad (4)$$

A similar function, called `k_3rd_iupac` here, is used by Wallington et al. (2018) for three-body reactions. It has the same function parameters as `k_3rd` and it is defined as:

$$k_0(T) = k_0^{300} \times \left(\frac{300\text{K}}{T}\right)^n \quad (5)$$

$$k_{\text{inf}}(T) = k_{\text{inf}}^{300} \times \left(\frac{300\text{K}}{T}\right)^m \quad (6)$$

$$k_{\text{ratio}} = \frac{k_0(T)M}{k_{\text{inf}}(T)} \quad (7)$$

$$N = 0.75 - 1.27 \times \log_{10}(f_c) \quad (8)$$

$$\text{k_3rd_iupac} = \frac{k_0(T)M}{1 + k_{\text{ratio}}} \times f_c^{\left(\frac{1}{1 + (\log_{10}(k_{\text{ratio}})/N)^2}\right)} \quad (9)$$

Structure-Activity Relationships (SAR)

Some unmeasured rate coefficients are estimated with structure-activity relationships, using the following parameters and substituent factors:

k for H-abstraction by OH in $\text{cm}^{-3}\text{s}^{-1}$	
<code>k_p</code>	$4.49 \times 10^{-18} \times (T/\text{K})^2 \exp(-320\text{K}/T)$
<code>k_s</code>	$4.50 \times 10^{-18} \times (T/\text{K})^2 \exp(253\text{K}/T)$
<code>k_t</code>	$2.12 \times 10^{-18} \times (T/\text{K})^2 \exp(696\text{K}/T)$
<code>k_ROHRO</code>	$2.1 \times 10^{-18} \times (T/\text{K})^2 \exp(-85\text{K}/T)$
<code>k_CO2H</code>	$0.7 \times k_{\text{CH}_3\text{CO}_2\text{H}+\text{OH}}$
<code>k_ROOHRo</code>	$0.6 \times k_{\text{CH}_3\text{OOH}+\text{OH}}$
<code>f_alk</code>	1.23
<code>f_sOH</code>	3.44
<code>f_tOH</code>	2.68
<code>f_sOOH</code>	8.
<code>f_tOOH</code>	8.
<code>f_ONO2</code>	0.04
<code>f_CH2ONO2</code>	0.20
<code>f_cpan</code>	0.25
<code>f_allyl</code>	3.6
<code>f_CHO</code>	0.55
<code>f_CO2H</code>	1.67
<code>f_CO</code>	0.73
<code>f_O</code>	8.15
<code>f_pCH2OH</code>	1.29
<code>f_tCCH2OH</code>	0.53

k for OH-addition to double bonds in $\text{cm}^{-3}\text{s}^{-1}$	
<code>k_adp</code>	$4.5 \times 10^{-12} \times (T/300\text{K})^{-0.85}$
<code>k_ads</code>	$1/4 \times (1.1 \times 10^{-11} \times \exp(485\text{K}/T) + 1.0 \times 10^{-11} \times \exp(553\text{K}/T))$
<code>k_adt</code>	$1.922 \times 10^{-11} \times \exp(450\text{K}/T) - k_{\text{ads}}$
<code>k_adsecprim</code>	3.0×10^{-11}
<code>k_adtertprim</code>	5.7×10^{-11}
<code>a_PAN</code>	0.56
<code>a_CHO</code>	0.31
<code>a_COCH3</code>	0.76
<code>a_CH2OH</code>	1.7
<code>a_CH2OOH</code>	1.7
<code>a_COH</code>	2.2
<code>a_COOH</code>	2.2
<code>a_CO2H</code>	0.25
<code>a_CH2ONO2</code>	0.64

RO₂ self and cross reactions

The self and cross reactions of organic peroxy radicals are treated according to the permutation reaction formalism as implemented in the MCM (Rickard, 2022), as described by Jenkin et al. (1997). Every organic peroxy radical reacts in a pseudo-first-order reaction with a rate constant that is expressed as $k^{1\text{st}} = 2 \times \sqrt{k_{\text{self}} \times k_{\text{CH}3\text{O}2}} \times [\text{RO}_2]$ where k_{self} = second-order rate coefficient of the self reaction of the organic peroxy radical, $k_{\text{CH}3\text{O}2}$ = second-order rate coefficient of the self reaction of CH₃O₂, and $[\text{RO}_2]$ = sum of the concentrations of all organic peroxy radicals.

Specific notes

G2110: The rate coefficient is: $k_{\text{HO}_2\text{-HO}_2} = (3.0\text{E-}13\text{*EXP}(460./\text{temp})+2.1\text{E-}33\text{*EXP}(920./\text{temp})\text{*cair})*(1.+1.4\text{E-}21\text{*EXP}(2200./\text{temp})\text{*C(ind_H2O)})$.

G2117: Converted to Kc [molec⁻¹ cm³] = $K_p \cdot R \cdot T / N_A$, where R is 82.05736 [cm³ atm K⁻¹ mol⁻¹].

G2118: Assuming fast equilibrium.

G3109: The rate coefficient is: $k_{\text{NO}_3\text{-NO}_2} = k_{\text{3rd}(\text{temp, cair}, 2.4\text{E-}30, 3.0, 1.6\text{E-}12, -0.1, 0.6)}$.

G3110: The rate coefficient is defined as backward reaction divided by equilibrium constant.

G3203: The rate coefficient is: $k_{\text{NO}_2\text{-HO}_2} = k_{\text{3rd}(\text{temp, cair}, 1.9\text{E-}31, 3.4, 4.0\text{E-}12, 0.3, 0.6)}$.

G3206: The rate coefficient is: $k_{\text{HN}_3\text{-OH}} = 1.32\text{E-}14 * \text{EXP}(527/\text{temp}) + 1 / (1 / (7.39\text{E-}32 * \text{EXP}(453/\text{temp})\text{*cair}) + 1 / (9.73\text{E-}17 * \text{EXP}(1910/\text{temp})))$

G3207: The rate coefficient is defined as backward reaction divided by equilibrium constant.

G3227: Backward reaction divided by equilibrium constant from Burkholder et al. (2015).

G3228: Same as for OH + HNO₄.

G4104b: Methyl nitrate yield according to Banic et al. (2003) but reduced by a factor of 10 according to the upper limit derived from measurements by Munger et al. (1999).

G4109: Same temperature dependence as for CH₃CHO+NO₃ assumed.

G4115: The rate coefficient is defined as backward reaction divided by equilibrium constant.

G4116: Same value as for PAN + OH.

G4126: Same as for G4104 but scaled to match the recommended value at 298K.

G4127: Same as for CH₃O₂ + NO₃ in G4105.

G4130a: SAR for H-abstraction by OH.

G4130b: SAR for H-abstraction by OH.

G4132: SAR for H-abstraction by OH.

G4133: Lower limit of the rate constant. Products uncertain but CH₃OH can be excluded because of a likely high energy barrier (L. Vereecken, pers. comm.). CH₂OO production cannot be excluded.

G4134: Estimate based on the decomposition lifetime of 3 s (Olzmann et al., 1997) and a 20 kcal/mol energy barrier (Vereecken and Francisco, 2012).

G4135: Rate constant for CH₂OO + NO₂ (G4138) multiplied by the factor from Ouyang et al. (2013).

G4136: Average of two measurements.

G4137: Upper limit.

G4138: Average of 7.E-12 and 1.5E-12.

G4141: HOCH₂OCHO forms and then decomposes to formic anhydride (Gruzdev et al., 1993) which hydrolyses in the humid atmosphere (Conn et al., 1942).

G4142: High-pressure limit.

G4143: Generic estimate for reaction with alcohols.

G4144: Generic estimate for reaction with RO₂.

G4148: Same value as for NO₂+CH₃O₂.

G4149: Barnes et al. (1985) estimated a decomposition rate equal to that of CH₃O₂NO₂.

G4150: Value for CH₃O₂NO₂ + OH, H-abstraction enhanced by the HO-group by f.sOH.

G4154: Products assumed to be CH₃O₂ + O₂ (could also be HCHO + O₂ + OH).

G4160b: Half of the H-yield is attributed to fast secondary chemistry.

G4160c: The NH + CO channel is also significant but neglected here.

G4161: No studies below 450 K and only the major channel is considered.

G4164: Upper limit. Dominant pathway under atmospheric conditions.

G42001: The product distribution is from Rickard (2022), after substitution of the energized Criegee intermediate, CH₂OO, by its decomposition products and reaction of the stabilized CI with the water dimer.

G42010: Only major channel considered as the end products are essentially the same.

G42013: The rate coefficient is: $k_{\text{CH}_3\text{CO}_3\text{-NO}_2} = k_{\text{3rd}(\text{temp, cair}, 9.7\text{E-}29, 5.6, 9.3\text{E-}12, 1.5, 0.6)}$.

G42018: The rate coefficient is the same as for the CH₃ channel in G4107 (CH₃OOH+OH).

G42021: The rate coefficient is $k_{\text{PAN-M}} = k_{\text{CH}_3\text{CO}_3\text{-NO}_2} / 9.0\text{E-}29 * \text{EXP}(-14000./\text{temp})$, i.e. the rate coefficient is defined as backward reaction divided by equilibrium constant.

G42022a: Quantum yields and products are from Glowacki et al. (2012).

G42022b: Quantum yields and products are from Glowacki et al. (2012).

G42024a: Rate constant is the high-pressure limit as recommended by Atkinson et al. (2006).

G42024b: Rate constant is the high-pressure limit as recommended by Atkinson et al. (2006).

G42047: Orlando et al. (1998) estimated that about 25% of the HOCH₂CH₂O in this reaction is produced with sufficient excess energy that it decomposes promptly. The decomposition products are 2 HCHO + HO₂.

G42051a: Same as for the CH₃O₂ channel in G4107: CH₃OOH+OH.

- G42058b: The aldehydic H is assumed to be like the analogous H of HOCH₂CHO.
- G42074a: Factor of 3 to match the estimate of $k = 1.E-11$ molec/cm³/s by Paulot et al. (2009a).
- G42074b: Factor of 3 to match the estimate of $k = 1.E-11$ molec/cm³/s by Paulot et al. (2009a).
- G42075: NO₃CH₂CO₂H and NO₃CH₂CO₃H neglected.
- G42078: NO₃CH₂CO₂H neglected.
- G42082: Same rate constant as for PAN + OH.
- G42083a: Rate constant is the high-pressure limit as recommended by Atkinson et al. (2006).
- G42083b: Rate constant is the high-pressure limit as recommended by Atkinson et al. (2006).
- G42085a: Uncertainties on the kinetics at pressures < 0.1 bar.
- G42085b: Channel proposed by Hynes and Wine 1991, OH + HCHO + HOCl, could not be confirmed by Tyndall et al. (2001b). There is no alternative mechanism at the moment. Products assumed to be OH + CH₃CO₃ + NO
- G42086b: Assuming HCN is from channel 2h, HCO + H + HCN. HCO is replaced by H + CO.
- G42086c: Assuming exothermic channels 2b and 2d are equally important.
- G42087: HCOCN is produced but replaced here by its likely oxidation products (HCN + CO₂) as studied by Tyndall et al. (2001b). The rate constant for a typical RO₂ + NO reaction is used.
- G42088: NCCH₂OOH is produced but replaced here by its likely oxidation products (HCN + CO₂) as studied by Tyndall et al. (2001b). The rate constant for a typical RO₂ + HO₂ reaction is used.
- G42089a: The minor channel with $k=5.2E-12$ is combined with the major one producing HCOOH.
- G42090: Theoretical keto-enol tautomerization catalyzed by formic acid (Grenfell et al., 2006).
- G42091: Theoretical keto-enol tautomerization catalyzed by formic acid (Grenfell et al., 2006).
- G42092: approximated OH reaction for oxalic acid
- G42093a: SAR for H-abstraction by OH
- G42093b: SAR for H-abstraction by OH, assuming that -CHOHOH has an effect like -CH₂OH
- G42093c: SAR for H-abstraction by OH
- G42093d: SAR for H-abstraction by OH
- G42094a: SAR for H-abstraction by OH
- G42094b: SAR for H-abstraction by OH
- G42095a: SAR for H-abstraction by OH
- G42095b: SAR for H-abstraction by OH
- G42096a: SAR for H-abstraction by OH
- G42096b: SAR for H-abstraction by OH
- G42097a: SAR for H-abstraction by OH
- G42097b: SAR for H-abstraction by OH
- G42098a: SAR for H-abstraction by OH, assuming that -CH₂OOH has the same effect as -CH₂OH
- G42098b: SAR for H-abstraction by OH
- G42098c: SAR for H-abstraction by OH
- G43001a: Branching ratios according to Rickard et al. (1999).
- G43001b: Branching ratios according to Rickard et al. (1999).
- G43004: The value for the generic RO₂ + HO₂ reaction from Atkinson (1997) is used here.
- G43008: The value for the generic RO₂ + HO₂ reaction from Atkinson (1997) is used here.
- G43011: Strong positive deviation of k below 240 K compared to the expression recommended by JPL (Burkholder et al., 2015).
- G43015a: The same value as for G4107 (CH₃OOH + OH) is used, multiplied by the branching ratio of the CH₃O₂ channel.
- G43028: Alkyl nitrate formation neglected. (also not considered in MCM).
- G43037: Alkyl nitrate formation neglected. (also not considered in MCM).
- G43040a: Rate coefficient estimated with SAR (Taraborrelli, 2010).
- G43040b: Rate coefficient estimated with SAR (Taraborrelli, 2010).
- G43044: Alkyl nitrate formation neglected.
- G43045c: Rate coefficient assumed to equal to the one of hydroxyacetone (ACETOL) for this channel.
- G43048: Using the high-pressure limit.
- G43049: The pressure fall-off between 1000 and 100 mbar is only 3% (Kirchner et al., 1999).
- G43050: Value for CH₃O₂NO₂ + OH, H-abstraction enhanced by the CH₃CO-group by f.CO.
- G43051c: Products approximated with C₂H₅CHO + HO₂.
- G43052: Only major H-abstraction channel considered.
- G43059: Products approximated with the major end-product CH₃CHO.
- G43060b: Products approximated with the major end-product CH₃CHO.
- G43061: Products approximated with the likely end-product CH₃CHO.
- G43065: As for HCOCO₃.
- G43070a: Branching ratios estimated with SAR for H-abstraction rate constants by OH.

- G43070b: Branching ratios estimated with SAR for H-abstraction rate constants by OH.
- G43071a: Only this channel considered as the intermediate radical is likely more stable than CHCH(OH)₂.
- G43072: Theoretical keto-enol tautomerization catalyzed by formic acid (Grenfell et al., 2006).
- G43073: Theoretical keto-enol tautomerization catalyzed by formic acid (Grenfell et al., 2006).
- G43074: HCOCOCHO would be produced but undergoes fast photolysis (faster than MGLYOX) and is substituted with its products.
- G43075a: Same value as for methanediol.
- G43075b: Same value as for methanediol.
- G43223: Products simplified
- G43419: KDEC C3DIALO → GLYOX + CO + HO₂
- G43420: KDEC C3DIALO → GLYOX + CO + HO₂
- G43421: Permutation reaction (minor channels removed).
- G44000: The LC₄H₉O₂ composition (nC₄H₉O₂:sC₄H₉O₂ ratio) is assumed to be equal to the ratio of the production rates at 298K: k_p/(k_p+k_s) = 0.1273 and k_s/(k_p+k_s) = 0.8727.
- G44001b: sC₄H₉O₂ products are substituted with 0.636 MEK + HO₂ and 0.364 CH₃CHO + C₂H₅O₂ at 1 bar and 298 K.
- G44003c: The alkyl nitrate yield is the weighted average yield for the two isomers forming from nC₄H₉O₂ and sC₄H₉O₂.
- G44010b: H-abstraction from primary C and substitution of the resulting peroxy radical with its products from the reaction with NO.
- G44011: H-abstraction from primary C and substitution of the resulting peroxy radical with its products from the reaction with NO.
- G44015b: Products assumed to be only from H-abstraction from a secondary C bearing the -OOH group.
- G44016: Products assumed to be only from H-abstraction from a secondary C bearing the -ONO₂ group.
- G44018: LHMVKABO₂ is 0.12 HMVKAO₂ + 0.88 HMVKBO₂.
- G44019: LMEKO₂ represents 0.62 MEKBO₂ + 0.38 MEKAO₂.
- G44021a: The products of MEKAO are substituted with HCHO + CO₂ + HOCH₂CH₂O₂.
- G44023a: Products from H-abstraction from the tertiary carbon bearing the ONO₂ group.
- G44023b: Products from H-abstraction from the secondary carbon bearing the ONO₂ group.
- G44025: Same value as for PAN.
- G44026: Products as in G4415. Only the main channels for each isomer are considered. Weighted average for the isomers.
- G44035: Rate constant replaced with the one of beta hydroxy RO₂.
- G44046b: Using value for secondary nitrate (88% of total).
- G44061a: Using value for secondary nitrate (88% of total).
- G44061b: Using value for secondary nitrate (88% of total).
- G44062a: Simplified products.
- G44062b: Simplified products.
- G44066: Alkyl nitrate formation neglected.
- G44070: Alkyl nitrate formation neglected.
- G44076: Alkyl nitrate formation neglected.
- G44078: Other channel neglected.
- G44081: Alkyl nitrate formation neglected.
- G44082: Other channel neglected.
- G44085: k for CH₃CHCO from Hatakeyama et al. (1985) adjusted.
- G44086: Simplified product distribution.
- G44089: The nitrated RO₂ is replaced by its products upon reaction with NO.
- G44096: Both LBUT1ENO₂ isomers mostly C₂H₅CHO.
- G44097a: Branching ratios according to Rickard et al. (1999). CH₃CHO₂CHO is replaced with its major products CH₃CHO + CO + HO₂.
- G44097b: Branching ratios according to Rickard et al. (1999).
- G44098: The nitrated RO₂ is replaced by its products upon reaction with NO.
- G44103b: MEKCOH replaced by its major oxidation products.
- G44104: Carbonyl nitrate replaced by its major oxidation products.
- G44106: CH₃CHOOA products as from C₃H₆ + O₃ reaction.
- G44107: The nitrated RO₂ is replaced by its products upon reaction with NO.
- G44110: The nitrated RO₂ is replaced by its products upon reaction with NO.
- G44124b: Skipping intermediate steps mostly leading to acetone.
- G44126: Skipping intermediate steps mostly leading to acetone.
- G44127: Only this channel considered as the intermediate radical is likely more stable than CHCH(OH)₂.

- G44128: Theoretical keto-enol tautomerization catalyzed by formic acid (Grenfell et al., 2006).
- G44129: Theoretical keto-enol tautomerization catalyzed by formic acid (Grenfell et al., 2006).
- G44130: Only this channel considered as the intermediate radical is likely more stable than CHCH(OH)_2 .
- G44131: Theoretical keto-enol tautomerization catalyzed by formic acid (Grenfell et al., 2006).
- G44132: Theoretical keto-enol tautomerization catalyzed by formic acid (Grenfell et al., 2006).
- G44133: Only this channel considered as the intermediate radical is likely more stable than CHCH(OH)_2 .
- G44134: Theoretical keto-enol tautomerization catalyzed by formic acid (Grenfell et al., 2006).
- G44135: Theoretical keto-enol tautomerization catalyzed by formic acid (Grenfell et al., 2006).
- G44136: Only this channel considered as the intermediate radical is likely more stable than CHCH(OH)_2 .
- G44137: Theoretical keto-enol tautomerization catalyzed by formic acid (Grenfell et al., 2006).
- G44138: Theoretical keto-enol tautomerization catalyzed by formic acid (Grenfell et al., 2006).
- G44139: Simplified oxidation.
- G44140: Simplified oxidation.
- G44141: Simplified oxidation.
- G44142: Simplified oxidation.
- G44202: Alkyl nitrate formation neglected.
- G44203a: Rate coefficient estimated with SAR (Taraborrelli, 2010).
- G44205: Alkyl nitrate formation neglected.
- G44210: Alkyl nitrate formation neglected.
- G44221: Same k as for $\text{MGLYOX} + \text{OH}$ (Tyndall et al., 1995).
- G44402: KDEC $\text{NC4DCO}_2 \rightarrow \text{MALANHY} + \text{NO}_2$
- G44406c: KDEC $\text{MALDIALCO}_2 \rightarrow 0.6 \text{ MALANHY} + \text{HO}_2 + 0.4 \text{ GLYOX} + 0.4 \text{ CO} + 0.4 \text{ CO}_2$
- G44407: KDEC $\text{MALDIALCO}_2 \rightarrow 0.6 \text{ MALANHY} + \text{HO}_2 + 0.4 \text{ GLYOX} + 0.4 \text{ CO} + 0.4 \text{ CO}_2$
- G44409: KDEC $\text{MALDIALCO}_2 \rightarrow 0.6 \text{ MALANHY} + \text{HO}_2 + 0.4 \text{ GLYOX} + 0.4 \text{ CO} + 0.4 \text{ CO}_2$
- G44410: KDEC $\text{MALDIALCO}_2 \rightarrow 0.6 \text{ MALANHY} + \text{HO}_2 + 0.4 \text{ GLYOX} + 0.4 \text{ CO} + 0.4 \text{ CO}_2$
- G44412: KDEC $\text{BZFUONOOA} \rightarrow 0.5 \text{ BZFUONOO} + 0.5 \text{ CO} + 0.5 \text{ CO}_2 + 0.5 \text{ HCOCH}_2\text{O}_2 + 0.5 \text{ OH}$ and $\text{BZFUONOO} \rightarrow 0.625 \text{ CO14O}_3\text{CO}_2\text{H} + 0.375 \text{ CO14O}_3\text{CHO} + 0.375 \text{ H}_2\text{O}_2$
- G44421: Only major channel.
- G44424: KDEC: $\text{GLYOOA} \rightarrow 0.125 \text{ HCHO} + 0.18 \text{ GLYOO} + 0.82 \text{ HO}_2 + 0.57 \text{ OH} + 1.265 \text{ CO} + 0.25 \text{ CO}_2$ and H_2O substitution $\text{GLYOO} \rightarrow 0.625 \text{ HCOCO}_2\text{H} + 0.375 \text{ GLYOX} + 0.375 \text{ H}_2\text{O}_2$
- G44425: Merged equations.
- G44430: KDEC $\text{MALANHYO} \rightarrow \text{HCOCO}_2\text{H} + 0.375 \text{ GLYOX} + 0.375 \text{ H}_2\text{O}_2$
- G44431: KDEC $\text{MALANHYO} \rightarrow \text{HCOCO}_2\text{H} + 0.375 \text{ GLYOX} + 0.375 \text{ H}_2\text{O}_2$
- G44432: Only major channel. KDEC $\text{MALANHYO} \rightarrow \text{HCOCO}_2\text{H} + 0.375 \text{ GLYOX} + 0.375 \text{ H}_2\text{O}_2$
- G44436: KDEC $\text{NBZFUO} \rightarrow 0.5 \text{ CO14O}_3\text{CHO} + 0.5 \text{ NO}_2 + 0.5 \text{ NBZFUONE} + 0.5 \text{ HO}_2$
- G44437: KDEC $\text{NBZFUO} \rightarrow 0.5 \text{ CO14O}_3\text{CHO} + 0.5 \text{ NO}_2 + 0.5 \text{ NBZFUONE} + 0.5 \text{ HO}_2$
- G44438: KDEC $\text{NBZFUO} \rightarrow 0.5 \text{ CO14O}_3\text{CHO} + 0.5 \text{ NO}_2 + 0.5 \text{ NBZFUONE} + 0.5 \text{ HO}_2$ and RO_2 Only major channel.
- G44439: KDEC $\text{MALDIALCO}_2 \rightarrow 0.6 \text{ MALANHY} + \text{HO}_2 + 0.4 \text{ GLYOX} + 0.4 \text{ CO} + 0.4 \text{ CO}_2$
- G44443: KDEC $\text{MEOCACETO} \rightarrow \text{CH}_3\text{CO}_3 + \text{HCHO}$
- G44444: KDEC $\text{MEOCACETO} \rightarrow \text{CH}_3\text{CO}_3 + \text{HCHO}$
- G44445: KDEC $\text{MEOCACETO} \rightarrow \text{CH}_3\text{CO}_3 + \text{HCHO}$
- G44450: KDEC $\text{BZFUO} \rightarrow \text{CO14O}_3\text{CHO} + \text{HO}_2$
- G44451: KDEC $\text{BZFUO} \rightarrow \text{CO14O}_3\text{CHO} + \text{HO}_2$
- G44452: KDEC $\text{BZFUO} \rightarrow \text{CO14O}_3\text{CHO} + \text{HO}_2$. Only major channel.
- G44457: KDEC $\text{MALDIALO} \rightarrow \text{GLYOX} + \text{GLYOX} + \text{HO}_2$
- G44458: KDEC $\text{MALDIALO} \rightarrow \text{GLYOX} + \text{GLYOX} + \text{HO}_2$
- G44459: KDEC $\text{MALDIALO} \rightarrow \text{GLYOX} + \text{GLYOX} + \text{HO}_2$. Only major channel.
- G44461: $\text{KBPAN} \rightarrow \text{k_PAN_M}$
- G45019d: Delta-1 and delta-2 LIEPOX are not considered and replaced by beta-LIEPOX formed by ISOP-BOOH and ISOPDOOH.
- G45021: SAR estimate within uncertainty range of the experimentally determined rate constant by Solberg et al. (1997), 1.1E-11.
- G45037: SAR estimate within uncertainty range of the experimentally determined rate constant by Solberg et al. (1997), 4.2E-11.
- G45040: Alkyl nitrate formation neglected.
- G45043: Old MCM rate constant 4.16E-11.
- G45047: Alkyl nitrate formation neglected.
- G45055: Alkyl nitrate formation neglected.
- G45071: Alkyl nitrate formation neglected.
- G45074: Formic acid production consistent with results of Bates et al. (2014). Here, the high yields of formic

acid and hydroxycarbonyls at low NO from oxidation of cis-beta-LIEPOX (the most abundant isomer) are approximated with the production of DB1O which undergo both the Dibble double H-transfer to DB2O2 and HOCH2 elimination yielding HVMK and HMAC (ketovinyl alcohol potentially arising from decomposition of the alkoxy radical resulting from the ring opening after H-abstraction). The rate constant is from Paulot et al. (2009b) and adjusted based on Bates et al. (2014) that determined the single rate constants for the cis- and trans- beta isomer.

G45080: Alkyl nitrate formation neglected.

G45092a: C4MDIAL = CM4DIAL in MCM only from aromatics.

G45092b: Only one acyl peroxy radical considered.

G45093: Two aldehydic sites reacting with NO_3 but only one isomer product considered.

G45095: Alkyl nitrate formation neglected.

G45098: Alkyl nitrate formation neglected.

G45100: Alkyl nitrate formation neglected.

G45104a: DB1OOH is a hydroperoxide bearing a vinyl alcohol moiety that upon reaction with OH yields HCOOH (Davis et al., 1998).

G45107: OH production here is to take into account the hydroperoxidic function formed by the shift of the enolic hydrogen and not present in DB2O2. This approximation leads to spurious HO_2 production.

G45108a: Consistent with the results of Bates et al. (2014).

G45108b: Consistent with the results of Bates et al. (2014). Assuming that the enol alkoxy radical partly decomposes yielding a substitute vinyl alcohol.

G45111: Alkyl nitrate formation neglected.

G45114b: Here, formic acid is mechanistically produced by the OH-addition to the vinyl alcohol which, upon RO_2 -to- RO conversion (skipped here), yields the HOCHOH fragment which in turn reacts with O_2 forming $\text{HCOOH} + \text{HO}_2$. Along $\text{CH}_3\text{COCHOHCHO}$ should be produced but not in the mechanism. Only $\text{CH}_3\text{COCHO}_2\text{CHO}$. The rate constant is consistent with predictions by Ganzeveld et al. (2006) for ENOL. OH-addition to the OH-bearing carbon is considered the dominant channel as it is already for the ENOL (Ganzeveld et al., 2006).

G45115: Theoretical keto-enol tautomerization catalyzed by formic acid (Grenfell et al., 2006). The product should be C1ODC3OOHC4OD but it is neglected in the mechanism.

G45116: As for $\text{DB1OOH} + \text{OH}$.

G45117: Additional sinks for DB2OOH are neglected.

G45121b: Nitrate assumed to be major isomer that is mostly similar to products of ISOPDO2-chemistry.

G45128: Rate constant by Liljegren and Stevens (2013). A lumped RO_2 that upon conversion to RO yields 100% 2-methyl-butenedial (C4MDIAL) although Aschmann et al. (2014) quantified a 38% yield of the Z/E mixture.

G45129: As for 3METHYLFURAN + OH but with additional NO_2 production for mass conservation.

G45131: Alkyl nitrate formation neglected.

G45132: Hydroperoxide formation neglected.

G45134b: ZCO2HC23DBCOD formation is neglected. However, it is produced in MCM and in aromatic-related reactions under the name of MC3ODBCO2H.

G45139: LZCPANC23DBCOD is assumed to react like LC5PAN1719.

G45201: Alkyl nitrate formation neglected.

G45207: Alkyl nitrate formation neglected.

G45214: Alkyl nitrate formation neglected.

G45217: Alkyl nitrate formation neglected.

G45225: Alkyl nitrate formation neglected.

G45236: $\text{LMBOABO2} = 0.67 \text{ MBOAO2} + 0.33 \text{ MBOBO2}$

G45247: Alkyl nitrate formation neglected.

G45400: KDEC NC4MDCO2 → MMALANHY + NO_2

G45404: KDEC NTLFUO → ACCOMECHO + NO_2

G45405: KDEC NTLFUO → ACCOMECHO + NO_2

G45406: KDEC NTLFUO → ACCOMECHO

G45409: KBPAN → k_PAN_M(renaming)

G45413: KFPAN → k_CH3CO3_NO2 (renaming)

G45422: KDEC MMALANHYO → CO2H3CO3

G45423: KDEC MMALANHYO → CO2H3CO3

G45424: KDEC MMALANHYO → CO2H3CO3 and Only major channel.

G45429: KBPAN → k_PAN_M (renamed)

G45430a: KDEC C5CO14CO2 → 0.83 MALANHY + 0.83 CH3 + 0.17 MGLYOX + 0.17 HO2 + 0.17 CO + 0.17 CO2

G45431: KDEC C5CO14CO2 → 0.83 MALANHY + 0.83 CH3 + 0.17 MGLYOX + 0.17 HO2 + 0.17 CO + 0.17 CO2

G45432: KFPAN → k_CH3CO3_NO2 (renaming)

G45433: KDEC C5CO14CO2 → 0.83 MALANHY + 0.83 CH3 + 0.17 MGLYOX + 0.17 HO2 + 0.17 CO + 0.17 CO2

G45434: KDEC C5CO14CO2 → 0.83 MALANHY + 0.83 CH3 + 0.17 MGLYOX + 0.17 HO2 + 0.17 CO + 0.17 CO2 and only major channel.

G45436: KDEC C5CO14CO2 → 0.83 MALANHY + 0.83 CH3 + 0.17 MGLYOX + 0.17 HO2 + 0.17 CO + 0.17 CO2

- G45444: KDEC MC3CODBCO₂ → 0.35 GLYOX + 0.35 CH₃ + 0.35 CO + 0.35 CO₂ + 0.65 MMALANHY + 0.65 HO₂
- G45452: KDEC TLFUONOOA → 0.5 CO + 0.5 OH + 0.5 MECOACETO₂ + 0.5 TLFUONOO and H₂O
subs TLFUONOO → 0.625 C₂₄O₃CCO₂H + 0.375 ACCOMECHO + 0.375 H₂O₂
- G45456: KFPAN →_kCH₃CO₃_NO₂ (renaming)
- G45476b: KDEC NTLFUO → ACCOMECHO + NO₂ and reactions with KRO₂HO₂.
- G45477: KDEC NTLFUO → ACCOMECHO + NO₂
- G45478: KDEC NTLFUO → ACCOMECHO + NO₂
- G45479: KDEC NTLFUO → ACCOMECHO + NO₂
- G45486b: KDEC C5DIALO → MALDIAL + CO + HO₂ and reactions with KRO₂HO₂.
- G45487: KDEC C5DIALO → MALDIAL
- G45488: KDEC C5DIALO → MALDIAL
- G45489: KDEC C5DIALO → MALDIAL
- G45491b: Reactions with KRO₂HO₂.
- G45492: MGLYOX + GLYOX + HO₂ from KDEC substitution
- G45493: MGLYOX + GLYOX + HO₂ from KDEC substitution
- G45494: Permutation reaction (minor channels removed).
- G46201: Alkyl nitrate formation neglected.
- G46404b: Reactions with KRO₂HO₂ and KDEC C₆15CO₂O → C₅DICARB + CO + HO₂.
- G46405: KDEC C₆15CO₂O → C₅DICARB + CO + HO₂
- G46406: KDEC C₆15CO₂O → C₅DICARB + CO + HO₂
- G46407: Only major channel.
- G46413b: Reactions with KRO₂HO₂ and KDEC ND-NPHENO → NC₄DCO₂H + HNO₃ + CO + CO + NO₂.
- G46414: KDEC ND-NPHENO → NC₄DCO₂H + HNO₃ + CO + CO + NO₂
- G46415: KDEC ND-NPHENO → NC₄DCO₂H + HNO₃ + CO + CO + NO₂
- G46416: KDEC ND-NPHENO → NC₄DCO₂H + HNO₃ + CO + CO + NO₂
- G46418: KDEC CATECOOA → MALDALCO₂H + HCOCO₂H + HO₂ + OH
- G46426: KFPAN →_kCH₃CO₃_NO₂
- G46430: KDEC GLYOOA → .125 HCHO + .18 GLYOO + .82 HO₂ + .57 OH + 1.265 CO
- G46432b: Reactions with KRO₂HO₂ and KDEC NCATECO → NC₄DCO₂H + HCOCO₂H + HO₂
- G46433: KDEC NCATECO → NC₄DCO₂H + HCOCO₂H + HO₂
- G46434: KDEC NCATECO → NC₄DCO₂H + HCOCO₂H + HO₂
- G46435: KDEC NCATECO → NC₄DCO₂H + HCOCO₂H + HO₂
- G46437b: Reactions with KRO₂HO₂ and KDEC NPHENO → MALDALCO₂H + GLYOX + NO₂
- G46438: KDEC NPHENO → MALDALCO₂H + GLYOX + NO₂
- G46439: KDEC NPHENO → MALDALCO₂H + GLYOX + NO₂
- G46440: KDEC NPHENO → MALDALCO₂H + GLYOX + NO₂
- G46441: Merged equations.
- G46447b: reactions with KRO₂HO₂ and KDEC NNCATECO → NC₄DCO₂H + HCOCO₂H + NO₂
- G46448: KDEC NNCATECO → NC₄DCO₂H + HCOCO₂H + NO₂
- G46449: KDEC NNCATECO → NC₄DCO₂H + HCOCO₂H + NO₂
- G46450: KDEC NNCATECO → NC₄DCO₂H + HCOCO₂H + NO₂
- G46457: Merged equations.
- G46458: Merged equations.
- G46461b: Reactions with KRO₂HO₂ and KDEC PHENO → 0.71 MALDALCO₂H + 0.71 GLYOX + 0.29 PBZQONE + HO₂
- G46462: KDEC PHENO → 0.71 MALDALCO₂H + 0.71 GLYOX + 0.29 PBZQONE + HO₂
- G46463: KDEC PHENO → 0.71 MALDALCO₂H + 0.71 GLYOX + 0.29 PBZQONE + HO₂
- G46464: KDEC PHENO → 0.71 MALDALCO₂H + 0.71 GLYOX + 0.29 PBZQONE + HO₂ and Only major channel.
- G46468: KFPAN →_kCH₃CO₃_NO₂
- G46472b: new channel
- G46476: HOC₆H₄NO₂ is a nitro-phenol
- G46480b: Reactions with KRO₂HO₂ and KDEC PBZQO → C₅CO₂OHCO₃
- G46481: KDEC PBZQO → C₅CO₂OHCO₃
- G46482: KDEC PBZQO → C₅CO₂OHCO₃
- G46483: KDEC PBZQO → C₅CO₂OHCO₃ and Only major channel.
- G46485b: Reactions with KRO₂HO₂ and KDEC DNPHENNO → NC₄DCO₂H + HCOCO₂H + NO₂

G46486: KDEC DNPHENO → NC4DCO2H + HCOCO2H + NO2	G47210: Alkyl nitrate formation neglected.	C5DICARB + 0.2 TLFUONE + 0.2 BZFUONE + 0.2 MALDIAL
G46487: KDEC DNPHENO → NC4DCO2H + HCOCO2H + NO2	G47214: Alkyl nitrate formation neglected.	G47410: Only major channel and KDEC TLBIPERO → 0.6 GLYOX + 0.4 MGLYOX + HO2 + 0.2 ZCODC23DB COD + 0.2 C5DICARB + 0.2 TLFUONE + 0.2 BZFUONE + 0.2 MALDIAL
G46488: KDEC DNPHENO → NC4DCO2H + HCOCO2H + NO2	G47218: Alkyl nitrate formation neglected.	G47412: KDEC MGLOOB → 0.125 CH3CHO + 0.695 CH3CO + 0.57 CO + 0.57 OH + 0.125 HO2 + 0.18 MGLOO + 0.25 CO2
G46490b: Reactions with KRO2HO2 and KDEC BZEMUCO → 0.5 EPXC4DIAL + 0.5 GLYOX + 0.5 HO2 + 0.5 C3DIALO2 + 0.5 C32OH13CO.	G47222: Alkyl nitrate formation neglected.	G47413: Merged.
G46491b: KDEC BZEMUCO → 0.5 EPXC4DIAL + 0.5 GLYOX + 0.5 HO2 + 0.5 C3DIALO2 + 0.5 C32OH13CO.	G47223: ROO6R3OOH produced but no sink for it.	G47418b: Reactions with KRO2HO2 and KDEC CRESO → 0.68 C5CO14OH + 0.68 GLYOX + HO2 + 0.32 PTLQONE.
G46492: KDEC BZEMUCO → 0.5 EPXC4DIAL + 0.5 GLYOX + 0.5 HO2 + 0.5 C3DIALO2 + 0.5 C32OH13CO	G47225: ROO6R4P produced but no sink for it.	G47419: KDEC CRESO → 0.68 C5CO14OH + 0.68 GLYOX + HO2 + 0.32 PTLQONE
G46493: KDEC BZEMUCO → 0.5 EPXC4DIAL + 0.5 GLYOX + 0.5 HO2 + 0.5 C3DIALO2 + 0.5 C32OH13CO and Only major channel.	G47226: ROO6R5P produced but no sink for it	G47420: KDEC CRESO → 0.68 C5CO14OH + 0.68 GLYOX + HO2 + 0.32 PTLQONE
G46499b: Reactions with KRO2HO2 and KDEC NBZQO → C6CO4DB + NO2.	G47400: Merged.	G47421: KDEC CRESO → 0.68 C5CO14OH + 0.68 GLYOX + HO2 + 0.32 PTLQONE and Only major channel.
G46500: KDEC NBZQO → C6CO4DB + NO2	G47402a: KROPRIM*O2 fast reaction C6H5CH2O = BENZAL + HO2.	G47422b: Reactions with KRO2HO2 and KDEC NCRESO → C5CO14OH + GLYOX + NO2
G46501: KDEC NBZQO → C6CO4DB + NO2	G47402b: KROPRIM*O2 fast reaction C6H5CH2O = BENZAL + HO2.	G47423: KDEC NCRESO → C5CO14OH + GLYOX + NO2
G46502: KDEC NBZQO → C6CO4DB + NO2	G47403: KROPRIM*O2 fast reaction C6H5CH2O = BENZAL + HO2.	G47424: KDEC NCRESO → C5CO14OH + GLYOX + NO2
G46505b: New channel.	G47404: KROPRIM*O2 fast reaction C6H5CH2O = BENZAL + HO2. C6H5CH2OH replaced by its oxidation product BENZAL.	G47425: KDEC NCRESO → C5CO14OH + GLYOX + NO2 and Only major channel.
G46515: Only major channel.	G47405: Merged.	G47426: TOL1OHNO2 is a nitro-phenol
G46522b: In analogy to TLBIPERO2 from toluene (Birdsall et al., 2010).	G47406: Merged.	G47429: KDEC MCATECOOA → MC3ODBCO2H + HCOCO2H + HO2 + OH
G46523b: KDEC BZBIPERO → GLYOX + HO2 + 0.5 BZFUONE + 0.5 BZFUONE	G47407b: According to Birdsall et al. (2010), the branching ratio rbipero2_oh is set to 0.4 in order to take into account the OH-recycling and summed yield of butendial and methylbutendial.	G47436: KFPAN → k_CH3CO3_NO2
G46524: KDEC BZBIPERO → GLYOX + HO2 + 0.5 BZFUONE + 0.5 BZFUONE	G47408a: KDEC TLBIPERO → 0.6 GLYOX + 0.4 MGLYOX + HO2 + 0.2 C4MDIAL + 0.2 C5DICARB + 0.2 TLFUONE + 0.2 BZFUONE + 0.2 MALDIAL	G47438: Only major channel.
G46525: KDEC BZBIPERO → GLYOX + HO2 + 0.5 BZFUONE + 0.5 BZFUONE and Only major channel.	G47408b: KDEC TLBIPERO → 0.6 GLYOX + 0.4 MGLYOX + HO2 + 0.2 ZCODC23DB COD + 0.2 C5DICARB + 0.2 TLFUONE + 0.2 BZFUONE + 0.2 MALDIAL	
	G47409: KDEC TLBIPERO → 0.6 GLYOX + 0.4 MGLYOX + HO2 + 0.2 ZCODC23DB COD + 0.2	

- G47439b: Reactions with KRO₂HO₂ and KDEC TLEMU_{CO} → 0.5 C₃DIALO₂ + 0.5 CO₂H₃CHO + 0.5 EPXC4DIAL + 0.5 MGLYOX + 0.5 HO₂
- G47440b: KDEC TLEMU_{CO} → 0.5 C₃DIALO₂ + 0.5 CO₂H₃CHO + 0.5 EPXC4DIAL + 0.5 MGLYOX + 0.5 HO₂
- G47441: KDEC TLEMU_{CO} → 0.5 C₃DIALO₂ + 0.5 CO₂H₃CHO + 0.5 EPXC4DIAL + 0.5 MGLYOX + 0.5 HO₂
- G47442: KDEC TLEMU_{CO} → 0.5 C₃DIALO₂ + 0.5 CO₂H₃CHO + 0.5 EPXC4DIAL + 0.5 MGLYOX + 0.5 HO₂ and Only major channel.
- G47445: KFPAN →_k CH₃CO₃.NO₂
- G47447: Only major channel.
- G47454: New channel.
- G47479: New channel.
- G47482b: Reactions with KRO₂HO₂ and KDEC NPTLQO → C₇CO₄DB + NO₂
- G47483: KDEC NPTLQO → C₇CO₄DB + NO₂
- G47484: KDEC NPTLQO → C₇CO₄DB + NO₂
- G47485: KDEC NPTLQO → C₇CO₄DB + NO₂
- G47486b: Reactions with KRO₂HO₂ and KDEC PTLQO → C₆CO₂OHCO₃
- G47487: KDEC PTLQO → C₆CO₂OHCO₃
- G47488: KDEC PTLQO → C₆CO₂OHCO₃
- G47489: Only major channel. KDEC PTLQO → C₆CO₂OHCO₃.
- G47494: New channel.
- G47497b: Reactions with KRO₂HO₂ and KDEC MN-CATECO → NC₄MDCO₂H + HCOCO₂H + NO₂
- G47498: KDEC MN-CATECO → NC₄MDCO₂H + HCOCO₂H + NO₂
- G47499: KDEC MN-CATECO → NC₄MDCO₂H + HCOCO₂H + NO₂
- G47501b: Reactions with KRO₂HO₂ and KDEC MN-CATECO → NC₄MDCO₂H + HCOCO₂H + HO₂
- G47502: KDEC MN-CATECO → NC₄MDCO₂H + HCOCO₂H + HO₂
- G47503: KDEC MN-CATECO → NC₄MDCO₂H + HCOCO₂H + HO₂
- G47504: KDEC MN-CATECO → NC₄MDCO₂H + HCOCO₂H + HO₂
- G47509b: Reactions with KRO₂HO₂ and KDEC ND-NCRESO → NC₄MDCO₂H + HNO₃ + CO + CO + NO₂
- G47510: KDEC ND-NCRESO → NC₄MDCO₂H + HNO₃ + CO + CO + NO₂
- G47511: KDEC ND-NCRESO → NC₄MDCO₂H + HNO₃ + CO + CO + NO₂
- G47512: KDEC ND-NCRESO → NC₄MDCO₂H + HNO₃ + CO + CO + NO₂
- G47513b: Reactions with KRO₂HO₂ and KDEC DNCRESO → NC₄MDCO₂H + HCOCO₂H + NO₂
- G47514: KDEC DNCRESO → NC₄MDCO₂H + HCOCO₂H + NO₂
- G47515: KDEC DNCRESO → NC₄MDCO₂H + HCOCO₂H + NO₂
- G47516: KDEC DNCRESO → NC₄MDCO₂H + HCOCO₂H + NO₂
- G48202: Alkyl nitrate formation neglected.
- G48205: Alkyl nitrate formation neglected.
- G48210: Alkyl nitrate formation neglected.
- G48212: Alkyl nitrate formation neglected.
- G48216: Alkyl nitrate formation neglected.
- G48222: Alkyl nitrate formation neglected.
- G48400a: Same products as for toluene. Assuming a 1:1:1 proportion in xylenes emissions the analogous toluene product is produced with a rate constant equal to (1.36E-11*0.24 + 2.31E-11*0.29 + 1.43E-11*0.155)/3, where k and coefficients are for the single isomers ortho, meta and para from MCM.
- G48400b: Same products as for toluene. Assuming a 1:1:1 proportion in xylenes emissions the analogous toluene product is produced with a rate constant equal to (1.36E-11*0.05 + 2.31E-11*0.04 + 1.43E-11*0.10)/3, where k and coefficients are for the single isomers ortho, meta and para from MCM.
- G48400c: Same products as for toluene. Assuming a 1:1:1 proportion in xylenes emissions the analogous toluene product is produced with a rate constant equal to (1.36E-11*0.16 + 2.31E-11*0.17 + 1.43E-11*0.12)/3, where k and coefficients are for the single isomers ortho, meta and para from MCM.
- G48400d: Same products as for toluene. Assuming a 1:1:1 proportion in xylenes emissions the analogous toluene product is produced with a rate constant equal to (1.36E-11*0.55 + 2.31E-11*0.50 + 1.43E-11*0.625)/3, where k and coefficients are for the single isomers ortho, meta and para from MCM.
- G48401: Same products as for toluene. The rate constant is the average of m, p, o k=(4.10E-16+2.60E-16+5.00E-16)/3 = 3.9E-16.
- G48402: merged under same rate constant
- G48403: Same products as for toluene
- G48405: KDEC CH₂OOB → 0.24 CH₂OO + 0.40 CO + 0.36 HO₂ + 0.36 CO + 0.36 OH and H₂O + PH-CHOO → 0.625 PHCOOH + 0.375 BENZAL + 0.375 H₂O₂ + 0.2 CO₂
- G48408: KDEC NSTYRENEO → NO₂ + HCHO + BENZAL

- G48409: KDEC NSTYRENEO → NO₂ + HCHO + BENZAL
- G48410: KDEC NSTYRENEO → NO₂ + HCHO + BENZAL
- G48412b: KDEC STYRENO → HO₂ + HCHO + BENZAL and reactions with KRO₂HO₂.
- G48413: KDEC STYRENO → HO₂ + HCHO + BENZAL
- G48414: KDEC STYRENO → HO₂ + HCHO + BENZAL
- G48415: KDEC STYRENO → HO₂ + HCHO + BENZAL
- G49207: Alkyl nitrate formation neglected.
- G49238: Alkyl nitrate formation neglected.
- G49246: Only this channel considered as the intermediate radical is likely more stable than CHCH(OH)₂. Instead of the (lacking) carbonyl a product of further degradation is assumed.
- G49247: Theoretical keto-enol tautomerization catalyzed by formic acid (Grenfell et al., 2006).
- G49248: Theoretical keto-enol tautomerization catalyzed by formic acid (Grenfell et al., 2006).
- G49400a: Same products as for toluene. Assuming a 1:1:1 proportion in xylenes emissions the analogous toluene product is produced with a rate constant equal to (3.27E-11*0.21 + 3.25E-11*0.30 + 5.67E-11*0.14)/3, where k and coefficients are for the single isomers 1,2,3-, 1,3,4- and 1,3,5- from MCM.
- G49400b: Same products as for toluene. Assuming a 1:1:1 proportion in xylenes emissions the analogous toluene product is produced with a rate constant equal to (3.27E-11*0.06 + 3.25E-11*0.06 + 5.67E-11*0.03)/3, where k and coefficients are for the single isomers 1,2,3-, 1,3,4- and 1,3,5- from MCM.
- G49400c: Same products as for toluene. Assuming a 1:1:1 proportion in xylenes emissions the analogous toluene product is produced with a rate constant equal to (3.27E-11*0.03 + 3.25E-11*0.03 + 5.67E-11*.04)/3, where k and coefficients are for the single isomers 1,2,3-, 1,3,4- and 1,3,5- from MCM.
- G49400d: Same products as for toluene. Assuming a 1:1:1 proportion in xylenes emissions the analogous toluene product is produced with a rate constant equal to (3.27E-11*0.70 + 3.25E-11*0.61 + 5.67E-11*0.79)/3, where k and coefficients are for the single isomers 1,2,3-, 1,3,4- and 1,3,5- from MCM.
- G49401: Same products as for toluene. The rate constant is the average of m, p, o $k = (1.90+1.80+0.88)E-15/3=1.52E-15$.
- G40200: Products from Vereecken et al. (2007). LAP-INABO₂ = 0.65 APINAO₂ + 0.35 APINBO₂
- G40203: Weighted average for isomers A and B, $k = 0.33*9.20E-14+0.67*8.80E-13$.
- G40204: Weighted average for isomers A and B, $k = 0.35*1.83E-11+0.65*3.28E-11$.
- G40205: Weighted average for isomers A and B, $k = 0.35*5.50E-12+0.65*3.64E-12$.
- G40206: SAR-estimated rate constant, $(k_{ads}+k_{adt})*acoch3 = 6.46E-11$ where $k_{ads} = 3.0E-11$, $k_{adt} = 5.5E-11$, $acoch3 = 0.76$
- G40207: Alkyl nitrate formation neglected.
- G40211: Products from Rickard (2022).
- G40212: Products from Rickard (2022).
- G40232: Products from Capouet et al. (2008).
- G40242: Alkyl nitrate formation neglected.
- G40246: Products from Rickard (2022).
- G40248: Alkyl nitrate formation neglected.
- G40252a: Products from Vereecken and Peeters (2012).
- G40252b: Products from Vereecken and Peeters (2012).
- G40259: ROO₆R₁OOH is produced but no sink for it.
- G40262: RO₆R₁OOH is produced but no sink for it.
- G40266: Rate constant modified according to MCM protocol.
- G40267a: Products from Nguyen et al. (2009).
- G40268: Products from Rickard (2022).
- G40270: Alkyl nitrate neglected.
- G40274: As for RO₆R₁NO₃ in G4085.
- G40276: Only this channel considered as the intermediate radical is likely more stable than CHCH(OH)₂.
- G40277: Theoretical keto-enol tautomerization catalyzed by formic acid (Grenfell et al., 2006).
- G40278: Theoretical keto-enol tautomerization catalyzed by formic acid (Grenfell et al., 2006).
- G40282a: Products from Vereecken and Peeters (2012).
- G40282b: Products from Vereecken and Peeters (2012).
- G40283a: Products from Nguyen et al. (2009).
- G40284: Products from Rickard (2022).
- G40285a: Products from Vereecken and Peeters (2012).
- G40285b: Products from Vereecken and Peeters (2012).
- G40286a: Products from Nguyen et al. (2009).
- G40287: Products from Rickard (2022).
- G40400: DIET35TOL(from MCM) as representative of higher aromatics
- G40401: Same products as for toluene.
- G6103: The rate coefficient is defined as backward reaction divided by equilibrium constant.
- G6204: At low temperatures, there may be a minor reaction channel leading to O₃+HCl. See Finkbeiner et al. (1995) for details. It is neglected here.

G6402: The initial products are probably HCl and CH₂OOH (Atkinson et al., 2006). It is assumed that CH₂OOH dissociates into HCHO and OH.

G6409: It is assumed that the reaction liberates all Cl atoms in the form of HCl.

G643DT: H-abstraction channel producing HCl. Organic peroxy radicals (crudely) approximated with the ones from reaction with OH.

G644DT: Cl-addition channel. Chlorinated organic peroxy radicals (crudely) approximated with the ones from reaction with OH.

G652DT: Product distribution could be different.

G654DT: By using LMEKO2 (lumped RO₂ from reaction with OH) the assumed yield of the isomer MEKBO₂ is 0.62 instead of 0.79 as recommended by IUPAC.

G7302: The rate coefficient is: $k_{BrO_N02} = k_{3rd(temp, cair, 5.2E-31, 3.2, 6.9E-12, 2.9, 0.6)}$.

G7303: The rate coefficient is defined as backward reaction (Atkinson et al., 2007) divided by equilibrium constant (Orlando and Tyndall, 1996).

G7404: It is assumed that the reaction liberates all Br atoms in the form of HBr.

G7407: It is assumed that the reaction liberates all Br atoms. The fate of the carbon atom is currently not considered.

G7408: It is assumed that the reaction liberates all Br atoms. The fate of the carbon atom is currently not considered.

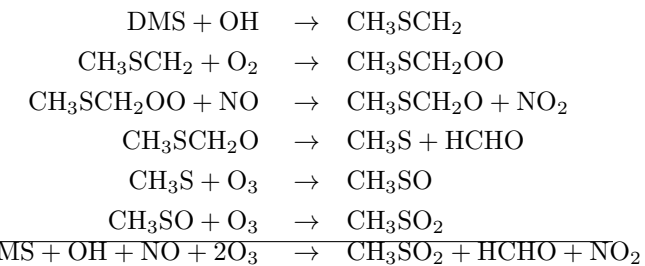
G7605: Same value as for G7408: CH₂Br₂+OH assumed. It is assumed that the reaction liberates all Br and Cl atoms. The fate of the carbon atom is currently not considered.

G7606: Same value as for G7408: CH₂Br₂+OH assumed. It is assumed that the reaction liberates all Br and Cl atoms. The fate of the carbon atom is currently not considered.

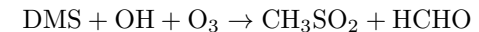
G7607: It is assumed that the reaction liberates all Br and Cl atoms. The fate of the carbon atom is currently not considered.

G9400a: For the abstraction path, the assumed reaction sequence (omitting H₂O and O₂ as products) according

to Yin et al. (1990) is:



Neglecting the effect on O₃ and NO_x, the remaining reaction is:



G9400b: For the addition path, the rate coefficient is: $k_{DMS_OH} = 1.0E-39 * \text{EXP}(5820./\text{temp}) * C(\text{ind_O2}) / (1.+5.0E-30 * \text{EXP}(6280./\text{temp}) * C(\text{ind_O2}))$.

G9402: Products and yields are not from Hynes and Wine (1996).

G9408: Average of 3.9E-11 and 3.42E-11.

Table 2: Photolysis reactions

#	labels	reaction	rate coefficient	reference
J (gas)				
J1000a	UpStTrGJ	O ₂ + hν → O(³ P) + O(³ P)	jx(ip_O2)	Sander et al. (2014)
J1001a	UpStTrGJ	O ₃ + hν → O(¹ D) + O ₂	jx(ip_01D)	Sander et al. (2014)
J1001b	UpStTrGJ	O ₃ + hν → O(³ P) + O ₂	jx(ip_03P)	Sander et al. (2014)
J2101	UpStTrGJ	H ₂ O ₂ + hν → 2 OH	jx(ip_H2O2)	Sander et al. (2014)
J3101	UpStTrGJN	NO ₂ + hν → NO + O(³ P)	jx(ip_NO2)	Sander et al. (2014)
J3103a	UpStTrGJN	NO ₃ + hν → NO ₂ + O(³ P)	jx(ip_NO20)	Sander et al. (2014)
J3103b	UpStTrGJN	NO ₃ + hν → NO + O ₂	jx(ip_NO02)	Sander et al. (2014)
J3104	StTrGJN	N ₂ O ₅ + hν → NO ₂ + NO ₃	jx(ip_N205)	Sander et al. (2014)
J3200	TrGJN	HONO + hν → NO + OH	jx(ip_HONO)	Sander et al. (2014)
J3201	StTrGJN	HNO ₃ + hν → NO ₂ + OH	jx(ip_HN03)	Sander et al. (2014)
J3202	StTrGJN	HNO ₄ + hν → .667 NO ₂ + .667 HO ₂ + .333 NO ₃ + .333 OH	jx(ip_HN04)	Sander et al. (2014)
J41000	StTrGJ	CH ₃ OOH + hν → CH ₃ O + OH	jx(ip_CH300H)	Sander et al. (2014)
J41001a	StTrGJ	HCHO + hν → H ₂ + CO	jx(ip_COH2)	Sander et al. (2014)
J41001b	StTrGJ	HCHO + hν → H + CO + HO ₂	jx(ip_CHOH)	Sander et al. (2014)
J41004	StTrGJN	CH ₃ ONO + hν → CH ₃ O + NO	jx(ip_CH30NO)	Sander et al. (2014)
J41005	StTrGJN	CH ₃ ONO ₂ + hν → CH ₃ O + NO ₂	jx(ip_CH3N03)	Sander et al. (2014)
J41006	StTrGJN	CH ₃ O ₂ NO ₂ + hν → .667 NO ₂ + .667 CH ₃ O ₂ + .333 NO ₃ + .333 CH ₃ O	jx(ip_CH302N02)	Sander et al. (2014)*
J41007	StTrGJ	HOCH ₂ OOH + hν → HCOOH + OH + HO ₂	jx(ip_CH300H)	Sander et al. (2014)
J41008	StTrGJ	CH ₃ O ₂ + hν → HCHO + OH	jx(ip_CH302)	Sander et al. (2014)
J41009	StTrGJ	HCOOH + hν → CO + HO ₂ + OH	jx(ip_HCOOH)	Sander et al. (2014)
J41010	StTrGJN	HOCH ₂ O ₂ NO ₂ + hν → .667 NO ₂ + .667 HOCH ₂ O ₂ + .333 NO ₃ + .333 HCOOH + .333 HO ₂	jx(ip_CH302N02)	Sander et al. (2014)
J42000	TrGJC	C ₂ H ₅ OOH + hν → CH ₃ CHO + HO ₂ + OH	jx(ip_CH300H)	von Kuhlmann (2001)
J42001a	TrGJC	CH ₃ CHO + hν → CH ₃ + HO ₂ + CO	jx(ip_CH3CHO)	Sander et al. (2014)
J42001b	TrGJC	CH ₃ CHO + hν → CH ₂ CHOH	jx(ip_CH3CHO2VINY)	Clubb et al. (2012)
J42002	TrGJC	CH ₃ C(O)OOH + hν → CH ₃ + OH + CO ₂	jx(ip_CH3C03H)	Sander et al. (2014)
J42004	TrGJCN	PAN + hν → .7 CH ₃ C(O) + .7 NO ₂ + .3 CH ₃ + .3 CO ₂ + .3 NO ₃	jx(ip_PAN)	Sander et al. (2014)*
J42005a	TrGJC	HOCH ₂ CHO + hν → HCHO + 2 HO ₂ + CO	jx(ip_HOCH2CHO)*0.83	Sander et al. (2014)*
J42005b	TrGJC	HOCH ₂ CHO + hν → OH + HCOCH ₂ O ₂	jx(ip_HOCH2CHO)*0.07	Sander et al. (2014)*
J42005c	TrGJC	HOCH ₂ CHO + hν → CH ₃ OH + CO	jx(ip_HOCH2CHO)*0.10	Sander et al. (2014)*
J42006	TrGJC	HOCH ₂ CO ₃ H + hν → HCHO + HO ₂ + OH + CO ₂	jx(ip_CH300H)	Rickard (2022)

Table 2: Photolysis reactions (... continued)

#	labels	reaction	rate coefficient	reference
J42007	TrGJCN	PHAN + hν → .7 HOCH2CO + .7 NO ₂ + .3 HCHO + .3 HO ₂ + .3 CO ₂ + .3 NO ₃	jx(ip_PAN)	see note*
J42008	TrGJC	GLYOX + hν → 2 CO + 2 HO ₂	jx(ip_GLYOX)	Sander et al. (2014)
J42009	TrGJC	HCOCO ₂ H + hν → 2 HO ₂ + CO + CO ₂	jx(ip_MGLYOX)	Rickard (2022)
J42010	TrGJC	HCOCO ₃ H + hν → HO ₂ + CO + OH + CO ₂	jx(ip_CH3OOH)+jx(ip_HOCH2CHO)	Rickard (2022)
J42011	TrGJC	HYETHO2H + hν → HOCH ₂ CH ₂ O + OH	jx(ip_CH3OOH)	Rickard (2022)
J42012	TrGJCN	ETHOHN03 + hν → HO ₂ + 2 HCHO + NO ₂	j_IC3H7N03	Rickard (2022)
J42013	TrGJC	HOOCH2CO3H + hν → OH + HCHO + CO ₂ + OH	2.*jx(ip_CH3OOH)	Sander et al. (2019)
J42014	TrGC	HOOCH2CO2H + hν → OH + HCHO + HO ₂ + CO ₂	jx(ip_CH3OOH)	Sander et al. (2019)
J42015	TrGC	CH ₂ CO + hν → .4 CO ₂ + .8 H + .34 CO + .34 OH + .34 HO ₂ + .16 HCHO + .16 O(³ P) + .1 HCOOH + CO	j_ketene*0.36	Sander et al. (2019)
J42016	TrGC	CH3CHOHOOH + hν → CH ₃ + HCOOH + OH	jx(ip_CH3OOH)	Sander et al. (2019)
J42017	TrGJCN	NO ₃ CH2CHO + hν → HO ₂ + CO + HCHO + NO ₂	(jx(ip_C2H5N03)+jx(ip_CH3CHO)) * (jx(ip_NOA)+1E-10)/(0.59*j_IC3H7N03+jx(ip_CH3COCH3)+1E-10)	Sander et al. (2019)*
J42018	TrGJC	HOOCH2CHO + hν → OH + HCHO + CO + HO ₂	jx(ip_CH3OOH)+jx(ip_HOCH2CHO)	Sander et al. (2019)
J42019	TrGJCN	C ₂ H ₅ ONO ₂ + hν → CH ₃ CHO + HO ₂ + NO ₂	jx(ip_C2H5N03)	Sander et al. (2019)
J42020	TrGJCN	NO ₃ CH2CHO + hν → .7 NO ₃ CH2CO ₃ + .7 NO ₂ + .3 HCHO + .3 NO ₂ + .3 CO ₂ + .3 NO ₃	jx(ip_PAN)	Sander et al. (2019)*
J42021	StTrGJCN	C ₂ H ₅ O ₂ NO ₂ + hν → .667 NO ₂ + .667 C ₂ H ₅ O ₂ + .333 NO ₃ + .333 CH ₃ CHO + .333 HO ₂	jx(ip_CH3O2N02)	Sander et al. (2019)*
J42022	TrGJC	HOOCCOOH + hν → CO ₂ + .72 HCOOH + .28 CO + .28 H ₂ O	jx(ip_HOOCCOOH)	Yamamoto and Back (1985)
J43000	TrGJC	iC ₃ H ₇ OOH + hν → CH ₃ COCH ₃ + HO ₂ + OH	jx(ip_CH3OOH)	von Kuhlmann (2001)
J43001	TrGJC	CH ₃ COCH ₃ + hν → CH ₃ C(O) + CH ₃	jx(ip_CH3COCH3)	Sander et al. (2014)
J43002	TrGJC	CH ₃ COCH ₂ OH + hν → .5 CH ₃ C(O) + .5 HCHO + .5 HO ₂ + .5 HOCH2CO + .5 CH ₃	j_ACETOL	Sander et al. (2014)*
J43003	TrGJC	MGLYOX + hν → CH ₃ C(O) + CO + HO ₂	jx(ip_MGLYOX)	Sander et al. (2014)
J43004	TrGJC	CH ₃ COCH ₂ O ₂ H + hν → CH ₃ C(O) + HCHO + OH	jx(ip_CH3OOH)+j_ACETOL	Rickard (2022)
J43005	TrGJC	HOCH2COCH ₂ OOH + hν → HOCH2CO + HCHO + OH	jx(ip_CH3OOH)+j_ACETOL	Sander et al. (2019)
J43006	TrGJCN	iC ₃ H ₇ ONO ₂ + hν → CH ₃ COCH ₃ + NO ₂ + HO ₂	j_IC3H7N03	von Kuhlmann et al. (2003)*
J43007	TrGJCN	NOA + hν → CH ₃ C(O) + HCHO + NO ₂	jx(ip_NOA)	Barnes et al. (1993)
J43009	TrGJC	HYPROPO2H + hν → CH ₃ CHO + HCHO + HO ₂ + OH	jx(ip_CH3OOH)	Rickard (2022)
J43010	TrGJCN	PR2O2HNO3 + hν → NOA + HO ₂ + OH	jx(ip_CH3OOH)	Rickard (2022)

Table 2: Photolysis reactions (... continued)

#	labels	reaction	rate coefficient	reference
J43011	TrGJC	$\text{HOCH}_2\text{COCHO} + h\nu \rightarrow \text{HOCH}_2\text{CO} + \text{CO} + \text{HO}_2$	$jx(ip_MGLYOX)$	Rickard (2022)
J43012	TrGJC	$\text{HCOCOCH}_2\text{OOH} + h\nu \rightarrow \text{HCOCO} + \text{HCHO} + \text{OH}$	$jx(ip_CH3OOH) + j_ACETOL$	Sander et al. (2019)
J43013	TrGJC	$\text{HCOCOCH}_2\text{OOH} + h\nu \rightarrow \text{HOOCH}_2\text{CO}_3 + \text{CO} + \text{HO}_2$	$jx(ip_MGLYOX)$	Sander et al. (2019)
J43014	TrGJTerC	$\text{HCOCH}_2\text{CHO} + h\nu \rightarrow \text{HCOCH}_2\text{O}_2 + \text{HO}_2 + \text{CO}$	$jx(ip_HOCH2CHO)*2.$	Rickard (2022)
J43015	TrGJTerC	$\text{HCOCH}_2\text{CO}_2\text{H} + h\nu \rightarrow \text{HCOCH}_2\text{O}_2 + \text{CO}_2 + \text{HO}_2$	$jx(ip_HOCH2CHO)$	Rickard (2022)
J43016	TrGJTerC	$\text{HOC}_2\text{H}_4\text{CO}_3\text{H} + h\nu \rightarrow \text{HOCH}_2\text{CH}_2\text{O}_2 + \text{CO}_2 + \text{OH}$	$jx(ip_CH3OOH)$	Rickard (2022)
J43017	TrGJC	$\text{HCOCOCHO} + h\nu \rightarrow \text{HCOCO} + \text{HO}_2 + \text{CO}$	$2.*jx(ip_MGLYOX)$	Sander et al. (2019)
J43018	TrGJC	$\text{CH}_3\text{COCO}_2\text{H} + h\nu \rightarrow .32 \text{CH}_3\text{CHO} + .16 \text{CH}_2\text{CHOH} + .54 \text{CO}_2 + .38 \text{CH}_3\text{C(O)} + .38 \text{HO}_2 + .38 \text{CO}_2 + .07 \text{CH}_3\text{COOH} + .07 \text{CO} + .05 \text{CH}_3\text{C(O)} + .05 \text{CO} + .05 \text{OH}$	$jx(ip_CH3COCO2H)$	Sander et al. (2019)*
J43019	TrGC	$\text{CH}_3\text{COCO}_3\text{H} + h\nu \rightarrow \text{CH}_3\text{C(O)} + \text{OH} + \text{CO}_2$	$jx(ip_MGLYOX) + jx(ip_CH3OOH)$	Sander et al. (2019)
J43020	TrGC	$\text{CH}_3\text{CHCO} + h\nu \rightarrow \text{C}_2\text{H}_4 + \text{CO}$	$j_ketene*0.36*2.$	Sander et al. (2019)
J43021	TrGCN	$\text{PROPOLNO}_3 + h\nu \rightarrow \text{HOCH}_2\text{CHO} + \text{HCHO} + \text{HO}_2 + \text{NO}_2$	$j_IC3H7N03$	Sander et al. (2019)
J43022	TrGCN	$\text{CH}_3\text{COCH}_2\text{OONO}_2 + h\nu \rightarrow \text{CH}_3\text{C(O)} + \text{HCHO} + \text{NO}_3$	$jx(ip_CH3O2N02) + jx(ip_CH3COCH3)$	Sander et al. (2019)
J43023	TrGJC	$\text{C}_3\text{H}_7\text{OOH} + h\nu \rightarrow \text{C}_2\text{H}_5\text{CHO} + \text{HO}_2 + \text{OH}$	$jx(ip_CH3OOH)$	von Kuhlmann (2001)
J43024	TrGJCN	$\text{C}_3\text{H}_7\text{ONO}_2 + h\nu \rightarrow \text{C}_2\text{H}_5\text{CHO} + \text{NO}_2 + \text{HO}_2$	$0.59*j_IC3H7N03$	see note*
J43025a	TrGJC	$\text{C}_2\text{H}_5\text{CHO} + h\nu \rightarrow \text{C}_2\text{H}_5\text{O}_2 + \text{HO}_2 + \text{CO}$	$jx(ip_C2H5CH02HCO)$	see note*
J43025b	TrGJC	$\text{C}_2\text{H}_5\text{CHO} + h\nu \rightarrow \text{CH}_2\text{CHCH}_2\text{OH}$	$jx(ip_C2H5CH02ENOL)$	Andrews et al. (2012), Sander et al. (2019)*
J43026	TrGJCN	$\text{PPN} + h\nu \rightarrow .7 \text{C}_2\text{H}_5\text{CO}_3 + .7 \text{NO}_2 + .3 \text{C}_2\text{H}_5\text{O}_2 + .3 \text{CO}_2 + .3 \text{NO}_3$	$jx(ip_PAN)$	Sander et al. (2014)
J43027	TrGJC	$\text{C}_2\text{H}_5\text{CO}_3\text{H} + h\nu \rightarrow \text{C}_2\text{H}_5\text{O}_2 + \text{CO}_2 + \text{OH}$	$jx(ip_CH3OOH)$	von Kuhlmann (2001)
J43028a	TrGJC	$\text{HCOCOCH}_2\text{OOH} + h\nu \rightarrow \text{HOOCH}_2\text{CO}_3 + \text{CO} + \text{HO}_2$	$jx(ip_MGLYOX)$	Sander et al. (2019)
J43028b	TrGJC	$\text{HCOCOCH}_2\text{OOH} + h\nu \rightarrow \text{HCOCO} + \text{HCHO} + \text{OH}$	$jx(ip_HOCH2CHO) + jx(ip_CH3OOH)$	Sander et al. (2019)
J43200	TrGJTerC	$\text{HCOCH}_2\text{CO}_3\text{H} + h\nu \rightarrow \text{HCOCH}_2\text{O}_2 + \text{CO}_2 + \text{OH}$	$jx(ip_HOCH2CHO) + jx(ip_CH3OOH)$	Rickard (2022)
J43400	TrGJAroC	$\text{C}_3\text{DIALOOH} + h\nu \rightarrow \text{GLYOX} + \text{CO} + \text{HO}_2 + \text{OH}$	$jx(ip_HOCH2CHO)*2. + jx(ip_CH3OOH)$	Rickard (2022)*
J43401	TrGJAroC	$\text{C}_3\text{2OH13CO} + h\nu \rightarrow \text{GLYOX} + \text{HO}_2 + \text{HO}_2 + \text{CO}$	$jx(ip_HOCH2CHO)*2.$	Rickard (2022)
J43402	TrGJAroC	$\text{HCOCOHCO}_3\text{H} + h\nu \rightarrow \text{GLYOX} + \text{HO}_2 + \text{CO}_2 + \text{OH}$	$jx(ip_CH3OOH)$	Rickard (2022)
J44000a	TrGJC	$\text{LC}_4\text{H}_9\text{OOH} + h\nu \rightarrow \text{OH} + \text{C}_3\text{H}_7\text{CHO} + \text{HO}_2$	$jx(ip_CH3OOH)*(k_p/(k_p+k_s))$	Rickard (2022), Sander et al. (2019)
J44000b	TrGJC	$\text{LC}_4\text{H}_9\text{OOH} + h\nu \rightarrow \text{OH} + .636 \text{MEK} + .636 \text{HO}_2 + .364 \text{CH}_3\text{CHO} + .364 \text{C}_2\text{H}_5\text{O}_2$	$jx(ip_CH3OOH)*(k_s/(k_p+k_s))$	Rickard (2022), Sander et al. (2019)
J44001	TrGJC	$\text{MVK} + h\nu \rightarrow .5 \text{C}_3\text{H}_6 + .5 \text{CH}_3\text{C(O)} + .5 \text{HCHO} + \text{CO} + .5 \text{HO}_2$	$jx(ip_MVK)$	Sander et al. (2014)

Table 2: Photolysis reactions (... continued)

#	labels	reaction	rate coefficient	reference
J44002	TrGJC	MEK + $h\nu \rightarrow \text{CH}_3\text{C(O)} + \text{C}_2\text{H}_5\text{O}_2$	$0.42 * jx(ip_{\text{CHOH}})$	von Kuhlmann et al. (2003)
J44003	TrGJC	LMEKOOH + $h\nu \rightarrow .62 \text{CH}_3\text{C(O)} + .62 \text{CH}_3\text{CHO} + .38 \text{HCHO} + .38 \text{CO}_2 + .38 \text{HOCH}_2\text{CH}_2\text{O}_2 + \text{OH}$	$jx(ip_{\text{CH3OOH}}) + 0.42 * jx(ip_{\text{CHOH}})$	Sander et al. (2019)
J44004	TrGJC	BIACET + $h\nu \rightarrow 2 \text{CH}_3\text{C(O)}$	$2.15 * jx(ip_{\text{MGLYOX}})$	see note*
J44005a	TrGJCN	LC4H9NO3 + $h\nu \rightarrow \text{NO}_2 + \text{C}_3\text{H}_7\text{CHO} + \text{HO}_2$	$j_{\text{IC3H7NO3}} * (k_p / (k_p + k_s))$	see note*
J44005b	TrGJCN	LC4H9NO3 + $h\nu \rightarrow \text{NO}_2 + \text{MEK} + \text{HO}_2$	$j_{\text{IC3H7NO3}} * (k_s / (k_p + k_s))$	see note*
J44006	TrGJCN	MPAN + $h\nu \rightarrow .7 \text{MACO3} + .7 \text{NO}_2 + .3 \text{MACO2} + .3 \text{NO}_3$	$jx(ip_{\text{PAN}})$	see note*
J44007a	TrGJC	CO2H3CO3H + $h\nu \rightarrow \text{MGLYOX} + \text{HO}_2 + \text{OH} + \text{CO}_2$	$jx(ip_{\text{CH3OOH}})$	Rickard (2022)
J44007b	TrGJC	CO2H3CO3H + $h\nu \rightarrow \text{CH}_3\text{C(O)} + \text{HO}_2 + \text{HCOCO}_3\text{H}$	j_{ACETOL}	Rickard (2022)
J44008	TrGJC	MACR + $h\nu \rightarrow .5 \text{MACO3} + .5 \text{CH}_3\text{C(O)} + .5 \text{HCHO} + .5 \text{CO} + \text{HO}_2$	$jx(ip_{\text{MACR}})$	Sander et al. (2014)
J44009	TrGJC	MACROOH + $h\nu \rightarrow \text{MACRO} + \text{OH}$	$jx(ip_{\text{CH3OOH}}) + 2.77 * jx(ip_{\text{HOCH2CHO}})$	Sander et al. (2019)*
J44010	TrGJC	MACROH + $h\nu \rightarrow \text{CH}_3\text{COCH}_2\text{OH} + \text{CO} + \text{HO}_2 + \text{HO}_2$	$2.77 * jx(ip_{\text{HOCH2CHO}})$	see note*
J44011	TrGJC	MACO3H + $h\nu \rightarrow \text{MACO2} + \text{OH}$	$jx(ip_{\text{CH3OOH}})$	Sander et al. (2019)
J44012	TrGJC	LHMVKABOOH + $h\nu \rightarrow .12 \text{MGLYOX} + .12 \text{HO}_2 + .88 \text{CH}_3\text{C(O)} + .88 \text{HOCH}_2\text{CHO} + .12 \text{HCHO} + \text{OH}$	$jx(ip_{\text{CH3OOH}}) + j_{\text{ACETOL}}$	Sander et al. (2019)
J44013	TrGJC	CO2H3CHO + $h\nu \rightarrow \text{MGLYOX} + \text{CO} + \text{HO}_2 + \text{HO}_2$	$jx(ip_{\text{HOCH2CHO}}) + j_{\text{ACETOL}}$	Sander et al. (2019)
J44014	TrGJC	HO12CO3C4 + $h\nu \rightarrow \text{CH}_3\text{C(O)} + \text{HOCH}_2\text{CHO} + \text{HO}_2$	j_{ACETOL}	Rickard (2022)
J44015	TrGJC	BIACETOH + $h\nu \rightarrow \text{CH}_3\text{C(O)} + \text{HOCH}_2\text{CO}$	$2.15 * jx(ip_{\text{MGLYOX}})$	see note*
J44016	TrGC	HCOCC ₃ CO + $h\nu \rightarrow .5 \text{OH} + .5 \text{CH}_3\text{CHO} + \text{CO} + .5 \text{CH}_3\text{CHCO} + .5 \text{CO}$	j_{ketene}	Sander et al. (2019)
J44017a	TrGC	$\text{CH}_3\text{COCHCO} + h\nu \rightarrow .0192 \text{CH}_3\text{COCO}_2\text{H} + .1848 \text{H}_2\text{O}_2 + .2208 \text{MGLYOX} + .36 \text{OH} + .36 \text{CO} + .56 \text{CH}_3\text{C(O)} + .2 \text{CH}_3\text{CHO} + .2 \text{CO}_2 + .2 \text{HCHO} + .2 \text{HO}_2 + \text{CO}$	$j_{\text{ketene}} * 0.5$	Sander et al. (2019), Rickard (2022)*
J44017b	TrGC	$\text{CH}_3\text{COCHCO} + h\nu \rightarrow \text{CH}_3\text{CHCO} + \text{CO}$	$j_{\text{ketene}} * 0.5$	Sander et al. (2019)
J44018a	TrGJC	$\text{CH}_3\text{COCOCHO} + h\nu \rightarrow \text{CH}_3\text{C(O)} + 2 \text{CO} + \text{HO}_2$	$jx(ip_{\text{MGLYOX}})$	Sander et al. (2019)
J44018b	TrGJC	$\text{CH}_3\text{COCOCHO} + h\nu \rightarrow \text{HCOCO} + \text{CH}_3\text{C(O)}$	$2.15 * jx(ip_{\text{MGLYOX}})$	Sander et al. (2019)
J44019	TrGJC	$\text{CH}_3\text{COCOCO}_2\text{H} + h\nu \rightarrow \text{CH}_3\text{C(O)} + \text{CO} + \text{CO}_2 + \text{HO}_2$	$3.15 * jx(ip_{\text{MGLYOX}})$	Sander et al. (2019)
J44020a	TrGJTerC	$\text{CH}_3\text{COCOCH}_2\text{OOH} + h\nu \rightarrow \text{CH}_3\text{C(O)} + \text{OH} + \text{HCHO} + \text{CO}$	$jx(ip_{\text{CH3OOH}}) + j_{\text{ACETOL}}$	Rickard (2022)
J44020b	TrGJTerC	$\text{CH}_3\text{COCOCH}_2\text{OOH} + h\nu \rightarrow \text{CH}_3\text{C(O)} + \text{HCOCO}$	$2.15 * jx(ip_{\text{MGLYOX}})$	Rickard (2022)
J44021	TrGJTerC	$\text{C44OOH} + h\nu \rightarrow \text{HCOCH}_2\text{CHO} + \text{CO}_2 + \text{HO}_2 + \text{OH}$	$jx(ip_{\text{CH3OOH}})$	Rickard (2022)
J44022	TrGJTerC	$\text{C413COOOH} + h\nu \rightarrow \text{HCOCH}_2\text{CO}_3 + \text{HCHO} + \text{OH}$	$jx(ip_{\text{CH3OOH}}) + jx(ip_{\text{HOCH2CHO}}) + j_{\text{ACETOL}}$	Rickard (2022)

Table 2: Photolysis reactions (... continued)

#	labels	reaction	rate coefficient	reference
J44023a	TrGJTerC	C4CODIAL + hν → HCOCOCH ₂ O ₂ + HO ₂ + CO	jx(ip_HOCH2CHO)	Rickard (2022)
J44023b	TrGJTerC	C4CODIAL + hν → HCOCH ₂ CO ₃ + HO ₂ + CO	jx(ip_MGLYOX)	Rickard (2022)
J44024	TrGJTerC	C312COCO ₃ H + hν → HCOCOCH ₂ O ₂ + CO ₂ + OH	jx(ip_CH300H)+jx(ip_MGLYOX)	Rickard (2022)
J44025	TrGJCN	LMEKNO ₃ + hν → .62 CH ₃ C(O) + .62 CH ₃ CHO + .38 HCHO + .38 CO ₂ + .38 HOCH ₂ CH ₂ O ₂ + NO ₂	jx(ip_MEKN03)	Barnes et al. (1993), Sander et al. (2019)*
J44026	TrGJCN	MVKNO ₃ + hν → CH ₃ C(O) + HOCH ₂ CHO + NO ₂	jx(ip_MEKN03)	Barnes et al. (1993), Sander et al. (2019)*
J44027	TrGJCN	MACRNO ₃ + hν → CH ₃ COCH ₂ OH + CO + HO ₂ + NO ₂	(2.84*j_IC3H7N03+jx(ip_CH3CHO)) *(jx(ip_MEKN03)+1E-10)/(j_IC3H7N03+0.42*jx(ip_CHOH)+1E-10)	Müller et al. (2014), Sander et al. (2019)*
J44028	TrGJCN	TC4H ₉ NO ₃ + hν → CH ₃ COCH ₃ + CH ₃ + NO ₂	2.84*j_IC3H7N03	Sander et al. (2019)
J44029	TrGJC	TC ₄ H ₉ OOH + hν → CH ₃ COCH ₃ + CH ₃ + OH	jx(ip_CH300H)	Sander et al. (2019)
J44030	TrGJCN	IBUTOLBNO ₃ + hν → CH ₃ COCH ₃ + HCHO + HO ₂ + NO ₂	2.84*j_IC3H7N03	Sander et al. (2019)
J44031	TrGJC	IBUTOLBOOH + hν → CH ₃ COCH ₃ + HCHO + HO ₂ + OH	jx(ip_CH300H)	Sander et al. (2019)
J44032	TrGJC	LBUT1ENO ₀ H + hν → C ₂ H ₅ CHO + HCHO + HO ₂ + OH	jx(ip_CH300H)	Sander et al. (2019)
J44033	TrGJCN	LBUT1ENNO ₃ + hν → C ₂ H ₅ CHO + HCHO + HO ₂ + NO ₂	j_IC3H7N03	Sander et al. (2019)
J44034	TrGJC	BUT2OLOOH + hν → 2 CH ₃ CHO + HO ₂ + OH	jx(ip_CH300H)	Sander et al. (2019)
J44035	TrGJCN	BUT2OLNO ₃ + hν → 2 CH ₃ CHO + HO ₂ + NO ₂	j_IC3H7N03	Sander et al. (2019)
J44036	TrGJC	BUT2OLO + hν → CH ₃ C(O) + HOCH ₂ CO	j_ACETOL	Sander et al. (2019)
J44037a	TrGJC	C ₃ H ₇ CHO + hν → C ₃ H ₇ O ₂ + CO + HO ₂	jx(ip_C3H7CH02HCO)	Sander et al. (2019)
J44037b	TrGJC	C ₃ H ₇ CHO + hν → C ₂ H ₄ + CH ₂ CHOH	jx(ip_C3H7CH02VINY)	Sander et al. (2019)*
J44038	TrGJC	IPRCHO + hν → iC ₃ H ₇ O ₂ + CO + HO ₂	jx(ip_IPRCHO2HCO)	Sander et al. (2019)
J44039	TrGJCN	IC4H ₉ NO ₃ + hν → IPRCHO + NO ₂	j_IC3H7N03	Sander et al. (2019)
J44040	TrGJC	IC ₄ H ₉ OOH + hν → IPRCHO + HO ₂ + OH	jx(ip_CH300H)	Sander et al. (2019)
J44041	TrGJC	PERIBUACID + hν → iC ₃ H ₇ O ₂ + CO ₂ + OH	jx(ip_CH300H)	Sander et al. (2019)
J44042	TrGJCN	PIP _N + hν → .7 IPRCO ₃ + .7 NO ₂ + .3 iC ₃ H ₇ O ₂ + .3 CO ₂ + .3 NO ₃	jx(ip_PAN)	Sander et al. (2019), Sander et al. (2014)
J44043	TrGJC	HVMK + hν → MGLYOX + CO + 2 OH	jx(ip_PeDIONE24)	Sander et al. (2019), Nakanishi et al. (1977), Messaadie et al. (2015), Yoon et al. (1999)*
J44044	TrGJC	HMAC + hν → HCOCCH ₃ CO + 2 OH	jx(ip_PeDIONE24)	Sander et al. (2019), Nakanishi et al. (1977), Messaadie et al. (2015), Yoon et al. (1999)*

Table 2: Photolysis reactions (... continued)

#	labels	reaction	rate coefficient	reference
J44045a	TrGJC	CO2C3CHO + hν → CH ₃ COCH ₂ O ₂ + HO ₂ + CO	jx(ip_C2H5CH02HCO)	Rickard (2022)
J44045b	TrGJC	CO2C3CHO + hν → HVMK	jx(ip_C2H5CH02ENOL)	Andrews et al. (2012), Sander et al. (2019)
J44046a	TrGJC	IBUTDIAL + hν → CH ₃ CHO + CO + HO ₂ + CO ₂ + H ₂ O	jx(ip_C2H5CH02HCO)*2.	see note*
J44046b	TrGJC	IBUTDIAL + hν → HMAC	jx(ip_C2H5CH02ENOL)*2.	Andrews et al. (2012), Sander et al. (2019)
J44200	TrGJTerC	IBUTALOH + hν → CH ₃ COCH ₃ + HO ₂ + HO ₂ + CO	j_ACETOL	Rickard (2022)
J44201	TrGJTerC	IPRHOCO3H + hν → CH ₃ COCH ₃ + HO ₂ + CO ₂ + OH	jx(ip_CH300H)	Rickard (2022)
J44400a	TrGJAroC	MALDIALOOH + hν → C32OH13CO + CO + OH + HO ₂	jx(ip_HOCH2CHO)*2.	Rickard (2022)
J44400b	TrGJAroC	MALDIALOOH + hν → GLYOX + GLYOX + HO ₂ + OH	jx(ip_CH300H)	Rickard (2022)*
J44401	TrGJAroC	BZFUOOH + hν → CO14O3CHO + HO ₂ + OH	jx(ip_CH300H)	Rickard (2022)*
J44402	TrGJAroC	HOCOC4DIAL + hν → HCOCOHCO ₃ + HO ₂ + CO	jx(ip_MGLYOX)+jx(ip_HOCH2CHO)	Rickard (2022)
J44403	TrGJAroCN	NBZFUOOH + hν → .5 CO14O3CHO + .5 NO ₂ + .5 NBZFUONE + .5 HO ₂ + OH	jx(ip_CH300H)	Rickard (2022)*
J44404a	TrGJAroC	MALDALCO3H + hν → HCOCO ₃ H + HO ₂ + CO + HO ₂ + CO	jx(ip_MACR)	Rickard (2022)
J44404b	TrGJAroC	MALDALCO3H + hν → .6 MALANHY + HO ₂ + .4 GLYOX + .4 CO + .4 CO ₂ + OH	jx(ip_CH300H)	Rickard (2022)*
J44405	TrGJAroC	EPXDLCO2H + hν → C3DIALO ₂ + CO ₂ + HO ₂	2.77*jx(ip_HOCH2CHO)	Rickard (2022)
J44406	TrGJAroC	MALDIAL + hν → .4 BZFUONE + .6 MALDIALCO ₃ + .6 HO ₂	jx(ip_NO2)*0.14	Rickard (2022)
J44407	TrGJAroC	MALANHYOOH + hν → HCOCOHCO ₃ + CO ₂ + OH	jx(ip_CH300H)	Rickard (2022)*
J44408	TrGJAroC	EPXDLCO3H + hν → C3DIALO ₂ + OH + CO ₂	jx(ip_CH300H)+2.77*jx(ip_HOCH2CHO)	Rickard (2022)
J44409	TrGJAroC	CO2C4DIAL + hν → CO + CO + HO ₂ + HO ₂ + CO + CO	jx(ip_MGLYOX)*2.	Rickard (2022)
J44410	TrGJAroC	MALDALCO2H + hν → HCOCO ₂ H + HO ₂ + CO + HO ₂ + CO	jx(ip_MACR)	Rickard (2022)
J44411	TrGJAroC	EPXC4DIAL + hν → C3DIALO ₂ + CO + HO ₂	2.77*jx(ip_HOCH2CHO)*2.	Rickard (2022)
J44412	TrGJAroC	CO14O3CHO + hν → HO ₂ + CO + HCOCH ₂ O ₂ + CO ₂	jx(ip_MGLYOX)	Rickard (2022)
J44414	TrGJAroC	MECOACEOOH + hν → CH ₃ C(O) + HCHO + CO ₂ + OH	jx(ip_CH300H)	Rickard (2022)*
J45002	TrGJC	LISOPACOOH + hν → LISOPACO + OH	jx(ip_CH300H)	Rickard (2022)
J45003	TrGJCN	LISOPACNO ₃ + hν → LISOPACO + NO ₂	0.59*j_IC3H7N03	see note*
J45004	TrGJC	ISOPBOOH + hν → MVK + HCHO + HO ₂ + OH	jx(ip_CH300H)	Rickard (2022)
J45005	TrGJCN	ISOPBNO ₃ + hν → MVK + HCHO + HO ₂ + NO ₂	2.84*j_IC3H7N03	see note*
J45006	TrGJC	ISOPDOOH + hν → MACR + HCHO + HO ₂ + OH	jx(ip_CH300H)	Rickard (2022)
J45007	TrGJCN	ISOPDNO ₃ + hν → MACR + HCHO + HO ₂ + NO ₂	j_IC3H7N03	see note*
J45008	TrGJCN	NISOPOOH + hν → NC4CHO + HO ₂ + OH	jx(ip_CH300H)	Rickard (2022)

Table 2: Photolysis reactions (... continued)

#	labels	reaction	rate coefficient	reference
J45009	TrGJCN	$\text{NC4CHO} + h\nu \rightarrow \text{LHC4ACCO}_3 + \text{NO}_2$	$(.59*j_{\text{IC3H7N03}}+j_x(\text{ip_MACR})) * (j_x(\text{ip_MEKN03})+1\text{E}-10) / (j_{\text{IC3H7N03}}+0.42*j_x(\text{ip_CHOH})+1\text{E}-10)$	Müller et al. (2014), Sander et al. (2019)*
J45010	TrGJCN	$\text{LNISOOH} + h\nu \rightarrow \text{NOA} + \text{OH} + .5 \text{HOCHCHO} + .5 \text{CO} + .5 \text{HO}_2 + .5 \text{CO}_2$	$j_x(\text{ip_CH300H})$	Taraborrelli et al. (2009), Sander et al. (2019)
J45011	TrGJC	$\text{LHC4ACCHO} + h\nu \rightarrow .5 \text{LHC4ACCO}_3 + .5 \text{HO}_2 + .5 \text{CO} + .5 \text{OH} + .25 \text{MACRO}_2 + .25 \text{LHMVKABO}_2$	$j_x(\text{ip_MACR})$	Sander et al. (2019)
J45012	TrGJC	$\text{LC578OOH} + h\nu \rightarrow .25 \text{CH}_3\text{COCH}_2\text{OH} + .75 \text{MGLYOX} + .25 \text{HOCHCHO} + .75 \text{HOCH}_2\text{CHO} + .75 \text{HO}_2 + \text{OH}$	$j_x(\text{ip_CH300H}) + 2.77*j_x(\text{ip_HOCH2CHO})$	Sander et al. (2019)
J45013	TrGJC	$\text{LHC4ACCO}_3\text{H} + h\nu \rightarrow \text{OH} + .5 \text{MACRO}_2 + .5 \text{LHMVKABO}_2 + \text{OH} + \text{CO}_2$	j_{HPALD}	Sander et al. (2019)
J45014	TrGJCN	$\text{LC5PAN1719} + h\nu \rightarrow .7 \text{LHC4ACCO}_3 + .7 \text{NO}_2 + .15 \text{MACRO}_2 + .15 \text{LHMVKABO}_2 + .3 \text{CO}_2 + .3 \text{NO}_3$	$j_x(\text{ip_PAN})$	Sander et al. (2019)
J45015	TrGJC	$\text{HCOC5} + h\nu \rightarrow .65 \text{CH}_3 + .65 \text{CO} + .65 \text{HCHO} + .35 \text{OH} + .35 \text{CH}_3\text{COCH}_2\text{O}_2 + \text{HOCH2CO}$	$0.5*j_x(\text{ip_MVK})$	Sander et al. (2019)*
J45016	TrGJC	$\text{C59OOH} + h\nu \rightarrow \text{CH}_3\text{COCH}_2\text{OH} + \text{HOCH2CO} + \text{OH}$	$j_{\text{ACETOL}}+j_x(\text{ip_CH300H})$	Sander et al. (2019)
J45017	TrGJTerC	$\text{C511OOH} + h\nu \rightarrow \text{CH}_3\text{C(O)} + \text{HCOCH2CHO} + \text{OH}$	$j_x(\text{ip_CH300H})+j_x(\text{ip_HOCH2CHO})$	Rickard (2022)
J45018a	TrGJTerC	$\text{CO23C4CHO} + h\nu \rightarrow \text{CH}_3\text{COCOCH}_2\text{O}_2 + \text{HO}_2 + \text{CO}$	$j_x(\text{ip_HOCH2CHO})$	Rickard (2022)
J45018b	TrGJTerC	$\text{CO23C4CHO} + h\nu \rightarrow \text{CH}_3\text{C(O)} + \text{HCOCH2CO}_3$	$2.15*j_x(\text{ip_MGLYOX})$	Rickard (2022)
J45019	TrGJTerC	$\text{CO23C4CO3H} + h\nu \rightarrow \text{CH}_3\text{COCOCH}_2\text{O}_2 + \text{CO}_2 + \text{OH}$	$j_x(\text{ip_CH300H})+j_x(\text{ip_HOCH2CHO})$	Rickard (2022)
J45020	TrGJTerC	$\text{C512OOH} + h\nu \rightarrow \text{C513O}_2 + \text{OH}$	$j_x(\text{ip_CH300H})+j_x(\text{ip_HOCH2CHO})$	Rickard (2022)
J45021	TrGJTerC	$\text{CO13C4CHO} + h\nu \rightarrow \text{CHOC3COO}_2 + \text{CO} + \text{HO}_2$	$j_x(\text{ip_HOCH2CHO})*2.$	Rickard (2022)
J45022	TrGJTerC	$\text{C513OOH} + h\nu \rightarrow \text{GLYOX} + \text{HOC}_2\text{H}_4\text{CO}_3 + \text{OH}$	$j_x(\text{ip_CH300H})+j_x(\text{ip_HOCH2CHO})$	Rickard (2022)
J45023	TrGJTerC	$\text{C513CO} + h\nu \rightarrow \text{HOC}_2\text{H}_4\text{CO}_3 + \text{HO}_2 + \text{CO} + \text{CO}$	$j_x(\text{ip_MGLYOX})+2.15*j_x(\text{ip_MGLYOX})$	Rickard (2022)
J45024	TrGJTerC	$\text{C514OOH} + h\nu \rightarrow \text{CO13C4CHO} + \text{HO}_2 + \text{OH}$	$j_x(\text{ip_CH300H})+j_x(\text{ip_HOCH2CHO})*2.$	Rickard (2022)
J45025	TrGJTerCN	$\text{C514NO}_3 + h\nu \rightarrow \text{CO13C4CHO} + \text{HO}_2 + \text{NO}_2$	$j_{\text{IC3H7N03}}+j_x(\text{ip_HOCH2CHO})*2.$	Rickard (2022)
J45026a	TrGJC	$\text{LZCODC23DBCOOH} + h\nu \rightarrow \text{OH} + \text{CO} + \text{HVMK} + \text{OH}$	$j_{\text{HPALD}}*0.6*0.5$	Sander et al. (2019), Jenkin et al. (2015), Peeters et al. (2014)
J45026b	TrGJC	$\text{LZCODC23DBCOOH} + h\nu \rightarrow \text{OH} + \text{CO} + \text{CH}_3\text{C(O)} + \text{HOCH}_2\text{CHO}$	$j_{\text{HPALD}}*0.6*0.5$	Sander et al. (2019), Jenkin et al. (2015), Peeters et al. (2014)
J45026c	TrGJC	$\text{LZCODC23DBCOOH} + h\nu \rightarrow \text{OH} + \text{CO} + \text{HMAC} + \text{OH}$	$j_{\text{HPALD}}*0.4*0.5$	Sander et al. (2019), Jenkin et al. (2015), Peeters et al. (2014)

Table 2: Photolysis reactions (... continued)

#	labels	reaction	rate coefficient	reference
J45026d	TrGJC	LZC0DC23DBCOOH + h ν → OH + CO + CO + j_HPALD*0.4*0.5 CH ₃ COCH ₂ OH + HO ₂	j_HPALD*0.4*0.5	Sander et al. (2019), Jenkin et al. (2015), Peeters et al. (2014)
J45027	TrGJC	LZC03HC23DBCOD + h ν → .62 EZCH3CO ₂ CHCHO + .38 j_HPALD EZCHOCCH3CHO ₂ + OH + CO ₂	j_HPALD	Sander et al. (2019)
J45028a	TrGJC	C1OOHC2OOHC4OD + h ν → CH ₃ COCH ₂ O ₂ H + OH + 2 CO + HO ₂	2.77*jx(ip_HOCH2CHO)	Sander et al. (2019)
J45028b	TrGJC	C1OOHC2OOHC4OD + h ν → .5 CH ₃ COCH ₂ O ₂ H + .5 HOCHCHO + .5 CO ₂ H3CHO + .5 HCHO + 1.5 OH	2.*jx(ip_CH300H)	Sander et al. (2019)
J45029	TrGC	DB1OOH + h ν → DB1O ₂ + OH	jx(ip_CH300H)	Sander et al. (2019)
J45030	TrGC	DB2OOH + h ν → .48 CH ₃ COCH ₂ OH + .52 HOCH ₂ CHO + .52 MGLYOX + .48 GLYOX + HO ₂ + OH	jx(ip_CH300H)	Sander et al. (2019)
J45031a	TrGJC	C1ODC2OOHC4OD + h ν → MGLYOX + HOCHCHO + OH	jx(ip_CH300H)	Sander et al. (2019)
J45031b	TrGJC	C1ODC2OOHC4OD + h ν → CO ₂ H3CHO + CO + HO ₂ + OH	2.*2.77*jx(ip_HOCH2CHO)	Sander et al. (2019)
J45032	TrGJC	C4MDIAL + h ν → .5 CH ₃ COCHCO + .5 HCOCCH ₃ CO + CO + HO ₂ + OH	jx(ip_NO2)*0.1*0.5	Sander et al. (2019)*
J45033	TrGCN	DB1NO ₃ + h ν → DB1O ₂ + NO ₂	j_IC3H7N03	Sander et al. (2019)
J45034	TrGJTerC	CHOC3COOOH + h ν → CHOC3COO ₂ + CO ₂ + OH	jx(ip_CH300H)+jx(ip_HOCH2CHO) +j_ACETOL	Rickard (2022)
J45200a	TrGJTerC	LMBOABOOH + h ν → HOCH ₂ CHO + CH ₃ COCH ₃ + HO ₂ + OH	jx(ip_CH300H)*.67	Rickard (2022), Sander et al. (2019)
J45200b	TrGJTerC	LMBOABOOH + h ν → IBUTALOH + HCHO + HO ₂ + OH	jx(ip_CH300H)*.33	Rickard (2022), Sander et al. (2019)
J45201	TrGJTerC	MBOACO + h ν → HCHO + HO ₂ + IPRHO ₃	j_ACETOL	Rickard (2022)
J45202	TrGJTerC	MBOOCOCO + h ν → CO + HO ₂ + IPRHO ₃	jx(ip_MGLYOX)	Rickard (2022)
J45203a	TrGJTerCN	LNMBOABOOH + h ν → NO ₃ CH ₂ CHO + CH ₃ COCH ₃ + HO ₂ + OH	jx(ip_CH300H)*.65	Rickard (2022), Sander et al. (2019)
J45203b	TrGJTerCN	LNMBOABOOH + h ν → IBUTALOH + HCHO + NO ₂ + OH	jx(ip_CH300H)*.35	Rickard (2022), Sander et al. (2019)
J45204	TrGJTerCN	NC4OHCO ₃ H + h ν → IBUTALOH + CO ₂ + NO ₂ + OH	jx(ip_CH300H)	Rickard (2022)
J45400	TrGJAroC	C54CO + h ν → HO ₂ + CO + CO + CO + CH ₃ C(O)	jx(ip_MGLYOX)+2.15*jx(ip_MGLYOX)*2.	Rickard (2022)
J45401	TrGJAroC	C5134CO ₂ OH + h ν → CH ₃ COCOCHO + HO ₂ + CO + HO ₂	jx(ip_HOCH2CHO)+2.15*jx(ip_MGLYOX)	Rickard (2022)
J45402	TrGJAroC	C5DIALOOH + h ν → MALDIAL + CO + HO ₂ + OH	jx(ip_CH300H)+jx(ip_MACR)	Rickard (2022)*

Table 2: Photolysis reactions (... continued)

#	labels	reaction	rate coefficient	reference
J45406	TrGJAroC	C5CO14OH + hν → CH ₃ C(O) + HCOCO ₂ H + HO ₂ + CO	jx(ip_MVK)	Rickard (2022)
J45407	TrGJAroC	C5DICARB + hν → .6 C5CO14O ₂ + .6 HO ₂ + .4 TLFUONE	jx(ip_NO2)*0.2	Rickard (2022)*
J45408	TrGJAroC	MC3ODBCO2H + hν → CH ₃ COCO ₂ H + HO ₂ + CO + HO ₂ + CO	jx(ip_MACR)	Rickard (2022)
J45409	TrGJAroC	ACCOMECHO + hν → MECOACETO ₂ + HO ₂ + CO	jx(ip_HOCH2CHO)	Rickard (2022)
J45410	TrGJAroC	MMALNHYOOH + hν → CO ₂ H ₃ CO ₃ + CO ₂ + OH	jx(ip_CH300H)	Rickard (2022)*
J45411	TrGJAroC	C5DICAROOH + hν → MGLYOX + GLYOX + HO ₂ + OH	jx(ip_CH300H)+jx(ip_HOCH2CHO) +j_ACETOL	Rickard (2022)*
J45412	TrGJAroCN	NTLFUOOH + hν → ACCOMECHO + NO ₂ + OH	jx(ip_CH300H)	Rickard (2022)*
J45414	TrGJAroC	C5CO14OOH + hν → .83 MALANHY + .83 CH ₃ + .17 MGLYOX + .17 HO ₂ + .17 CO + .17 CO ₂ + OH	jx(ip_CH300H)	Rickard (2022)*
J45415	TrGJAroC	TLFUOOH + hν → ACCOMECHO + HO ₂ + OH	jx(ip_CH300H)	Rickard (2022)*
J45417	TrGJAroC	ACCOMEKO3H + hν → MECOACETO ₂ + CO ₂ + OH	jx(ip_CH300H)	Rickard (2022)
J45418	TrGJAroC	C5DIALCO + hν → MALDIALCO ₃ + CO + HO ₂	jx(ip_MGLYOX)+jx(ip_MACR)	Rickard (2022)
J46200	TrGJTernCN	C614NO ₃ + hν → CO ₂ 3C4CHO + HCHO + HO ₂ + NO ₂	2.15*jx(ip_MGLYOX)	Rickard (2022)
J46201	TrGJTernC	C614OOH + hν → CO ₂ 3C4CHO + HCHO + HO ₂ + OH	jx(ip_CH300H)+2.15*jx(ip_MGLYOX)	Rickard (2022)
J46202	TrGJTernC	CO ₂ 3C5CHO + hν → CO ₂ 3C4CO ₃ + CO + HO ₂	jx(ip_MGLYOX)	Rickard (2022)
J46203	TrGJTernC	CO ₂ 3C6OOH + hν → CO ₂ 3C4CO ₃ + HCHO + OH	jx(ip_CH300H)+2.15*jx(ip_MGLYOX)	Rickard (2022)
J46400	TrGJAroC	PHENOOH + hν → .71 MALDALCO ₂ H + .71 GLYOX + .29 PBZQONE + HO ₂ + OH	jx(ip_CH300H)	Rickard (2022)*
J46401	TrGJAroC	C6CO4DB + hν → C4CO2DBCO ₃ + HO ₂ + CO	jx(ip_MGLYOX)*2.	Rickard (2022)
J46402	TrGJAroC	C5CO2DCO3H + hν → CH ₃ C(O) + HCOCOCHO + CO ₂ + OH	jx(ip_CH300H)+jx(ip_MGLYOX)	Rickard (2022)
J46403	TrGJAroCN	NDNPHENOOH + hν → NC4DCO ₂ H + HNO ₃ + CO + CO + NO ₂ + OH	jx(ip_CH300H)	Rickard (2022)*
J46404	TrGJAroCN	BZBIPERNO ₃ + hν → GLYOX + HO ₂ + .5 BZFUONE + .5 BZFUONE + NO ₂	j_IC3H7N03	Rickard (2022)*
J46405	TrGJAroCN	HOC6H4NO ₂ + hν → HONO + CPDKETENE	jx(ip_HOC6H4N02)	Chen et al. (2011)*
J46406	TrGJAroC	CPDKETENE + hν → CO ₂ + CO + 2 HO ₂ + MALDIAL	j_ketene	see note*
J46407	TrGJAroC	C5COOHCO3H + hν → HOCOC4DIAL + HO ₂ + CO + CO ₂ + OH	jx(ip_CH300H)	Rickard (2022)
J46408	TrGJAroC	BZEPOXMUC + hν → .5 C5DIALO ₂ + 1.5 HO ₂ + 1.5 CO + .5 MALDIAL	4.E3*jx(ip_MVK)*0.1	Rickard (2022)
J46409	TrGJAroCN	NPHEN1OOH + hν → NPHEN1O + OH	jx(ip_CH300H)	Rickard (2022)
J46410	TrGJAroC	BZEMUCCO + hν → HCOCOHCO ₃ + C3DIALO ₂	jx(ip_HOCH2CHO)*2.+j_ACETOL	Rickard (2022)
J46411	TrGJAroC	BZEMUCCO2H + hν → C5DIALO ₂ + CO ₂ + HO ₂	jx(ip_MACR)	Rickard (2022)

Table 2: Photolysis reactions (... continued)

#	labels	reaction	rate coefficient	reference
J46412	TrGJAroCN	NNCATECOOH + hν → NC4DCO2H + HCOCO ₂ H + NO ₂ + OH	jx(ip_CH300H)	Rickard (2022)*
J46413	TrGJAroC	C615CO2OOH + hν → C5DICARB + CO + HO ₂ + OH	jx(ip_MVK)+jx(ip_CH300H)	Rickard (2022)
J46414	TrGJAroCN	NPHENOOH + hν → MALDALCO2H + GLYOX + OH + NO ₂	j_IC3H7NO3 + jx(ip_CH300H)	Rickard (2022)
J46415	TrGJAroCN	NCATECOOH + hν → NC4DCO2H + HCOCO ₂ H + HO ₂ + OH	jx(ip_CH300H)	Rickard (2022)*
J46416	TrGJAroC	PBZQOOH + hν → C5CO2OHCO ₃ + OH	jx(ip_CH300H)	Rickard (2022)*
J46417	TrGJAroC	BZOBIPEROH + hν → MALDIALCO ₃ + GLYOX + HO ₂	j_ACETOL	Rickard (2022)
J46418	TrGJAroC	BZBIPEROOH + hν → GLYOX + HO ₂ + .5 BZFUONE + .5 BZFUONE + OH	jx(ip_CH300H)	Rickard (2022)*
J46419	TrGJAroCN	NBZQOOH + hν → C6CO4DB + NO ₂ + OH	jx(ip_CH300H)	Rickard (2022)*
J46420	TrGJAroC	CATEC1OOH + hν → CATEC1O + OH	jx(ip_CH300H)	Rickard (2022)
J46421	TrGJAroC	C6125CO + hν → C5CO14O ₂ + CO + HO ₂	jx(ip_MGLYOX)+jx(ip_MVK)	Rickard (2022)
J46422	TrGJAroCN	DNPHENOOH + hν → NC4DCO2H + HCOCO ₂ H + NO ₂ + OH	jx(ip_CH300H)	Rickard (2022)*
J46423	TrGJAroC	BZEMUCCO3H + hν → C5DIALO ₂ + CO ₂ + OH	jx(ip_CH300H)+jx(ip_MACR)	Rickard (2022)
J46424	TrGJAroC	C6H5OOH + hν → C6H5O + OH	jx(ip_CH300H)	Rickard (2022)
J46425	TrGJAroC	BZEMUCOOH + hν → .5 EPXC4DIAL + .5 GLYOX + .5 HO ₂ + .5 C3DIALO ₂ + .5 C32OH13CO + OH	jx(ip_CH300H)+jx(ip_HOCH2CHO)*2.	Rickard (2022)*
J46427	TrGJAroCN	BZEMUCNO ₃ + hν → EPXC4DIAL + NO ₂ + GLYOX + HO ₂	2.77*jx(ip_HOCH2CHO)	Rickard (2022)
J46428	TrGJAroCN	DNPHEN + hν → HONO + NCPDKETENE	jx(ip_HOC6H4NO ₂)	Sander et al. (2019)
J46429	TrGJAroCN	NCPDKETENE + hν → CO ₂ + CO + 2 HO ₂ + NC4DCO2H	j_ketene	see note*
J47200	TrGJTerC	CO235C6CHO + hν → CHOC3COCO ₃ + CH ₃ C(O)	2.15*jx(ip_MGLYOX)	Rickard (2022)
J47201	TrGJTerC	C235C6CO3H + hν → CO235C6O ₂ + CO ₂ + OH	jx(ip_CH300H)+2.15*jx(ip_MGLYOX)	Rickard (2022)
J47202	TrGJTerC	C716OOH + hν → CO13C4CHO + CH ₃ C(O) + OH	jx(ip_CH300H)+jx(ip_HOCH2CHO)	Rickard (2022)
J47203	TrGJTerC	C721OOH + hν → C722O ₂ + OH	jx(ip_CH300H)	Rickard (2022)
J47204	TrGJTerC	C722OOH + hν → CH ₃ COCH ₃ + C44O ₂ + OH	jx(ip_CH300H)	Rickard (2022)
J47400	TrGJAroC	TLEPOXMUC + hν → .5 C615CO2O ₂ + HO ₂ + CO + .5 EPXC4DIAL + .5 CH ₃ C(O)	4.E3*jx(ip_MVK)*0.1	Rickard (2022)
J47401	TrGJAroC	C6H5CH2OOH + hν → BENZAL + HO ₂ + OH	jx(ip_CH300H)	Rickard (2022)*
J47402	TrGJAroCN	C6H5CH2NO ₃ + hν → BENZAL + HO ₂ + NO ₂	0.59*j_IC3H7NO3	Rickard (2022)*
J47403	TrGJAroC	BENZAL + hν → HO ₂ + CO + C6H5O ₂	jx(ip_BENZAL)	Wallington et al. (2018)
J47404	TrGJAroC	TLBIPEROOH + hν → .6 GLYOX + .4 MGLYOX + HO ₂ + .2 C4MDIAL + .2 C5DICARB + .2 TLFUONE + .2 BZFUONE + .2 MALDIAL + OH	jx(ip_CH300H)	Rickard (2022)*

Table 2: Photolysis reactions (... continued)

#	labels	reaction	rate coefficient	reference
J47405	TrGJAroCN	TLBIPERNO3 + hν → .6 GLYOX + .4 MGLYOX + HO ₂ + .2 C4MDIAL + .2 C5DICARB + .2 TLFUONE + .2 BZFUONE + .2 MALDIAL + NO ₂	j_IC3H7N03	Rickard (2022)*
J47406	TrGJAroC	TLOBIPEROH + hν → C5CO14O2 + GLYOX + HO ₂	j_ACETOL	Rickard (2022)
J47407	TrGJAroC	CRESOOH + hν → .68 C5CO14OH + .68 GLYOX + HO ₂ + .32 PTLQONE + OH	jx(ip_CH300H)	Rickard (2022)*
J47408a	TrGJAroCN	NCRESOOH + hν → .68 C5CO14OH + .68 GLYOX + HO ₂ + .32 PTLQONE + OH + NO ₂	j_IC3H7N03	Rickard (2022)*
J47408b	TrGJAroCN	NCRESOOH + hν → C5CO14OH + GLYOX + NO ₂ + OH	jx(ip_CH300H)	Rickard (2022)*
J47409	TrGJAroCN	TOL1OHNO2 + hν → HONO + MCPDKETENE	jx(ip_HOPh3Me2N02)	see note*
J47410	TrGJAroC	TLEMUCCO2H + hν → C615CO2O2 + CO ₂ + HO ₂	jx(ip_MACR)	Rickard (2022)
J47411	TrGJAroC	TLEMUCCO3H + hν → C615CO2O2 + CO ₂ + OH	jx(ip_CH300H)+jx(ip_MACR)	Rickard (2022)
J47412	TrGJAroC	TLEMUCOOH + hν → .5 C3DIALO2 + .5 CO2H3CHO + .5 EPXC4DIAL + .5 MGLYOX + .5 HO ₂ + OH	jx(ip_CH300H)+2.77*jx(ip_HOCH2CHO)+j_ACETOL	Rickard (2022)*
J47413	TrGJAroCN	TLEMUCNO3 + hν → EPXC4DIAL + NO ₂ + CH ₃ C(O) + CO + HO ₂	2.77*jx(ip_HOCH2CHO)+j_ACETOL	Rickard (2022)
J47414	TrGJAroC	TLEMUCCO + hν → CH ₃ C(O) + EPXC4DIAL + CO + HO ₂	2.77*jx(ip_HOCH2CHO)+2.15*jx(ip_MGLYOX)	Rickard (2022)
J47415	TrGJAroC	C6H5CO3H + hν → C6H5O2 + CO ₂ + OH	jx(ip_CH300H)	Rickard (2022)
J47416	TrGJAroC	OXYL1OOH + hν → TOL1O + OH	jx(ip_CH300H)	Rickard (2022)
J47417	TrGJAroCN	MNCATECH + hν → HONO + MCPDKETENE	jx(ip_HOPh3Me2N02)	see note*
J47418	TrGJAroC	MCPDKETENE + hν → CO ₂ + CO + 2 HO ₂ + C4MDIAL	j_ketene	see note*
J47419	TrGJAroCN	DNCRES + hν → HONO + MNCPDKETENE	jx(ip_HOPh3Me2N02)	see note*
J47420	TrGJAroCN	MNCPDKETENE + hν → CO ₂ + CO + 2 HO ₂ + NC4MDCO2HN	j_ketene	see note*
J47421	TrGJAroC	MCATEC1OOH + hν → MCATEC1O + OH	jx(ip_CH300H)	Rickard (2022)
J47422	TrGJAroCN	NPTLQOOH + hν → C7CO4DB + NO ₂ + OH	jx(ip_CH300H)	Rickard (2022)*
J47423	TrGJAroC	PTLQOOH + hν → C6CO2OHCO3 + OH	jx(ip_CH300H)	Rickard (2022)*
J47424	TrGJAroCN	NCRES1OOH + hν → NCRES1O + OH	jx(ip_CH300H)	Rickard (2022)
J47425	TrGJAroCN	MNNCATCOOH + hν → NC4MDCO2HN + HCOCO ₂ H + NO ₂ + OH	jx(ip_CH300H)	Rickard (2022)*
J47426	TrGJAroCN	MNCATECOOH + hν → NC4MDCO2HN + HCOCO ₂ H + HO ₂ + OH	jx(ip_CH300H)	Rickard (2022)*
J47427	TrGJAroC	C7CO4DB + hν → C5CO2DBCO3 + HO ₂ + CO	jx(ip_MGLYOX)*2.	Rickard (2022)

Table 2: Photolysis reactions (... continued)

#	labels	reaction	rate coefficient	reference
J47428	TrGJAroCN	NDNCRESOOH + hν → NC4MDCO2HN + HNO ₃ + CO + CO + NO ₂ + OH	jx(ip_CH300H)	Rickard (2022)*
J47429	TrGJAroCN	DNCRESOOH + hν → NC4MDCO2HN + HCOCO ₂ H + NO ₂ + OH	jx(ip_CH300H)	Rickard (2022)*
J47430	TrGJAroC	C6COOHCO3H + hν → C5134CO2OH + HO ₂ + CO + CO ₂ + OH	jx(ip_CH300H)	Rickard (2022)
J48200	TrGJTerC	C86OOH + hν → C511O ₂ + CH ₃ COCH ₃ + OH	jx(ip_CH300H)+ jx(ip_HOCH2CHO)	Rickard (2022)
J48201	TrGJTerC	C812OOH + hν → C813O ₂ + OH	jx(ip_CH300H)	Rickard (2022)
J48202	TrGJTerC	C813OOH + hν → CH ₃ COCH ₃ + C512O ₂ + OH	jx(ip_CH300H)+jx(ip_MGLYOX)	Rickard (2022)
J48203	TrGJTerC	C721CHO + hν → C721O ₂ + CO + HO ₂	jx(ip_HOCH2CHO)	Rickard (2022)
J48204	TrGJTerC	C721CO3H + hν → C721O ₂ + CO ₂ + OH	jx(ip_CH300H)	Rickard (2022)
J48205	TrGJTerC	C8BCOOH + hν → C89O ₂ + OH	jx(ip_CH300H)	Rickard (2022)
J48206	TrGJTerC	C89OOH + hν → C810O ₂ + OH	jx(ip_CH300H)+jx(ip_HOCH2CHO)	Rickard (2022)
J48207	TrGJTerCN	C89NO ₃ + hν → C810O ₂ + NO ₂	jx(ip_CH300H)+jx(ip_HOCH2CHO)	Rickard (2022)
J48208	TrGJTerC	C810OOH + hν → CH ₃ COCH ₃ + C514O ₂ + OH	jx(ip_CH300H)+jx(ip_HOCH2CHO)	Rickard (2022)
J48209	TrGJTerCN	C810NO ₃ + hν → CH ₃ COCH ₃ + C514O ₂ + NO ₂	2.84*j_IC3H7N03+jx(ip_HOCH2CHO)	Rickard (2022)
J48210	TrGJTerCN	C8BCNO ₃ + hν → C89O ₂ + NO ₂	j_IC3H7N03	Rickard (2022)
J48211	TrGJTerC	C85OOH + hν → C86O ₂ + OH	jx(ip_CH300H)+j_ACETOL	Rickard (2022)
J48400	TrGJAroC	STYRENOOH + hν → HO ₂ + HCHO + BENZAL + OH	jx(ip_CH300H)	Rickard (2022)*
J49200	TrGJTerC	C96OOH + hν → C97O ₂ + OH	jx(ip_CH300H)+j_ACETOL	Rickard (2022)
J49201	TrGJTerC	C97OOH + hν → C98O ₂ + OH	jx(ip_CH300H)+j_ACETOL	Rickard (2022)
J49202	TrGJTerC	C98OOH + hν → C614O ₂ + CH ₃ COCH ₃ + OH	(jx(ip_CH300H)+2.15*jx(ip_MGLYOX))	Rickard (2022)
J49203a	TrGJTerC	NORPINAL + hν → C85O ₂ + CO + HO ₂	jx(ip_PINAL2HCO)	Rickard (2022), Sander et al. (2019)
J49203b	TrGJTerC	NORPINAL + hν → NORPINENOL	jx(ip_PINAL2ENOL)	Sander et al. (2019), Andrews et al. (2012)
J49204	TrGJTerC	C85CO3H + hν → C85O ₂ + CO ₂ + OH	jx(ip_CH300H)+j_ACETOL	Rickard (2022)
J49205	TrGJTerC	C89CO2H + hν → .8 C811CO3 + .2 C89O ₂ + .2 CO ₂ + HO ₂	jx(ip_HOCH2CHO)	Rickard (2022)
J49206	TrGJTerC	C89CO3H + hν → .8 C811CO3 + .2 C89O ₂ + .2 CO ₂ + OH	jx(ip_CH300H)+jx(ip_HOCH2CHO)	Rickard (2022)
J49207	TrGJTerC	C811CO3H + hν → C811O ₂ + CO ₂ + OH	jx(ip_CH300H)	Rickard (2022)
J49208	TrGJTerC	NOPINDOOH + hν → C89CO3 + OH	jx(ip_CH300H)	Rickard (2022)
J40200	TrGJTerC	LAPINABOOH + hν → PINAL + HO ₂ + OH	jx(ip_CH300H)	Rickard (2022)
J40201	TrGJTerC	MENTHEN6ONE + hν → RO6R1O ₂ + OH	jx(ip_CH300H)	Vereecken et al. (2007)
J40202	TrGJTerC	2OHMENTHEN6ONE + hν → 10 LCARBON + OH	jx(ip_CH300H)	Vereecken et al. (2007)

Table 2: Photolysis reactions (... continued)

#	labels	reaction	rate coefficient	reference
J40203a	TrGJTerC	PINAL + hν → C96O2 + CO + HO2	jx(ip_PINAL2HCO)	Rickard (2022)
J40203b	TrGJTerC	PINAL + hν → PINEOL	jx(ip_PINAL2ENOL)	Sander et al. (2019), Andrews et al. (2012)*
J40204	TrGJTerC	PERPINONIC + hν → C96O2 + CO2 + OH	jx(ip_CH300H)+j_ACETOL	Rickard (2022)
J40205	TrGJTerC	PINALOOH + hν → C106O2 + OH	jx(ip_CH300H)+jx(ip_HOCH2CHO)	Rickard (2022)
J40206	TrGJTerCN	PINALNO3 + hν → C106O2 + NO2	j_IC3H7N03+jx(ip_HOCH2CHO)	Rickard (2022)
J40207	TrGJTerC	C106OOH + hν → C716O2 + CH3COCH3 + OH	jx(ip_CH300H)+jx(ip_HOCH2CHO)	Rickard (2022)
J40208	TrGJTerCN	C106NO3 + hν → C716O2 + CH3COCH3 + NO2	j_IC3H7N03+ jx(ip_HOCH2CHO)	Rickard (2022)
J40209	TrGJTerC	C109OOH + hν → C89CO3 + HCHO + OH	jx(ip_CH300H)+jx(ip_HOCH2CHO)	Rickard (2022)
J40210	TrGJTerC	C109CO + hν → C89CO3 + CO + HO2	jx(ip_MGLYOX)+jx(ip_HOCH2CHO)	Rickard (2022)
J40211	TrGJTerCN	LNAPINABOOH + hν → PINAL + NO2 + OH	jx(ip_CH300H)	Rickard (2022)
J40212	TrGJTerC	BPINAOOH + hν → NOPINONE + HCHO + HO2 + OH	jx(ip_CH300H)	Rickard (2022)
J40213	TrGJTerCN	LNPINABOOH + hν → NOPINONE + HCHO + NO2 + OH	jx(ip_CH300H)	Rickard (2022)
J40214	TrGJTerCN	ROO6R1NO3 + hν → ROO6R3O2 + CH3COCH3 + NO2	2.84*j_IC3H7N03+jx(ip_CH300H)	Sander et al. (2019)
J40215	TrGJTerCN	RO6R1NO3 + hν → 9 LCARBON + HCHO + HO2 + NO2	2.84*j_IC3H7N03	Sander et al. (2019)
J6000	StTrGJC1	Cl2 + hν → Cl + Cl	jx(ip_C12)	Sander et al. (2014)
J6100	StTrGJC1	Cl2O2 + hν → 2 Cl	jx(ip_C12O2)	Sander et al. (2014)
J6101	StTrGJC1	OClO + hν → ClO + O(3P)	jx(ip_OClO)	Sander et al. (2014)
J61MS	StTrGJC1	ClO + hν → Cl + O(3P)	jx(ip_ClO)	Atkinson et al. (2007)
J62MS	StTrGJC1	Cl2O + hν → Cl + ClO	jx(ip_C12O)	Atkinson et al. (2007)
J63MS	StTrGJC1	Cl2O3 + hν → ClO + ClO2	jx(ip_C12O3)	Atkinson et al. (2007)
J6201	StTrGJC1	HOCl + hν → OH + Cl	jx(ip_HOCl)	Sander et al. (2014)
J6300	TrGJC1N	ClNO2 + hν → Cl + NO2	jx(ip_C1N02)	Sander et al. (2014)
J6301a	StTrGJC1N	ClNO3 + hν → Cl + NO3	jx(ip_C1N03)	Sander et al. (2014)
J6301b	StTrGJC1N	ClNO3 + hν → ClO + NO2	jx(ip_C1ON02)	Sander et al. (2014)
J64MS	TrGJC1N	ClNO + hν → Cl + NO	jx(ip_C1NO)	Atkinson et al. (2007)
J65MS	TrGJC1N	ClONO + hν → Cl + NO2	jx(ip_C1ONO)	Atkinson et al. (2007)
J7000	StTrGJBr	Br2 + hν → Br + Br	jx(ip_Br2)	Sander et al. (2014)
J7100	StTrGJBr	BrO + hν → Br + O(3P)	jx(ip_BrO)	Sander et al. (2014)
J7200	StTrGJBr	HOBBr + hν → Br + OH	jx(ip_HOBr)	Sander et al. (2014)
J7300	TrGJBrN	BrNO2 + hν → Br + NO2	jx(ip_BrN02)	Sander et al. (2014)
J7301	StTrGJBrN	BrNO3 + hν → .85 Br + .85 NO3 + .15 BrO + .15 NO2	jx(ip_BrN03)	Sander et al. (2014)*
J7401	TrGJBr	CH2Br2 + hν → LCARBON + 2 Br	jx(ip_CH2Br2)	Sander et al. (2014)
J7402	TrGJBr	CHBr3 + hν → LCARBON + 3 Br	jx(ip_CHBr3)	Sander et al. (2014)
J7600	StTrGJBrCl	BrCl + hν → Br + Cl	jx(ip_BrCl)	Sander et al. (2014)

Table 2: Photolysis reactions (... continued)

#	labels	reaction	rate coefficient	reference
J7602	TrGJBrCl	$\text{CH}_2\text{ClBr} + h\nu \rightarrow \text{LCARBON} + \text{Br} + \text{Cl}$	jx(ip_CH2ClBr)	Sander et al. (2014)
J7603	TrGJBrCl	$\text{CHCl}_2\text{Br} + h\nu \rightarrow \text{LCARBON} + \text{Br} + 2 \text{Cl}$	jx(ip_CHC12Br)	Sander et al. (2014)
J7604	TrGJBrCl	$\text{CHClBr}_2 + h\nu \rightarrow \text{LCARBON} + 2 \text{Br} + \text{Cl}$	jx(ip_CHC1Br2)	Sander et al. (2014)
PH (aqueous)				
PH2100_a01	TrAa01ScJ	$\text{H}_2\text{O}_2(\text{aq}) + h\nu \rightarrow 2 \text{OH}(\text{aq})$	2.33*xaer(01)*jx(ip_H2O2)	see note*
PH3200_a01	TrAa01JN	$\text{NO}_3^-(\text{aq}) + h\nu \rightarrow \text{NO}_2(\text{aq}) + \text{OH}(\text{aq}) + \text{OH}^-(\text{aq})$	xaer(01)*jx(ip_NO2) * 1.4E-4	see note*
PH4100_a01	TrAa01ScJ	$\text{HOCH}_2\text{OOH}(\text{aq}) + h\nu \rightarrow \text{HCOOH}(\text{aq}) + \text{OH}(\text{aq}) + \text{HO}_2(\text{aq})$	2.33*xaer(01)*jx(ip_CH3OOH)	Sander et al. (2014)
PH4101_a01	TrAa01ScJ	$\text{CH}_3\text{OOH}(\text{aq}) + h\nu \rightarrow \text{HCHO}(\text{aq}) + \text{OH}(\text{aq}) + \text{HO}_2(\text{aq})$	2.33*xaer(01)*jx(ip_CH3OOH)	Sander et al. (2014)
PH4200_a01	TrAa01ScJC	$\text{C}_2\text{H}_5\text{OOH}(\text{aq}) + h\nu \rightarrow \text{CH}_3\text{CHO}(\text{aq}) + \text{HO}_2(\text{aq}) + \text{OH}(\text{aq})$	2.33*xaer(01)*jx(ip_CH3OOH)	von Kuhlmann (2001)*
PH4201_a01	TrAa01ScJC	$\text{HOOCH}_2\text{CO}_2\text{H}(\text{aq}) + h\nu \rightarrow \text{HCHO}(\text{aq}) + \text{CO}_2(\text{aq}) + \text{HO}_2(\text{aq}) + \text{OH}(\text{aq})$	2.33*xaer(01)*jx(ip_CH3OOH)	Rickard (2022)*
PH4202_a01	TrAa01ScJC	$\text{CH}_2\text{OOHCO}_2^-(\text{aq}) + h\nu \rightarrow \text{CHOHOOCOO}_2^-(\text{aq}) + \text{OH}(\text{aq})$	2.33*xaer(01)*jx(ip_CH3OOH)	see note*
PH4203_a01	TrAa01ScJC	$\text{CH}_3\text{C}(\text{O})\text{OOH}(\text{aq}) + h\nu \rightarrow \text{CH}_3\text{OO}(\text{aq}) + \text{CO}_2(\text{aq}) + \text{OH}(\text{aq})$	2.33*xaer(01)*jx(ip_CH3C03H)	Sander et al. (2014)
PH4204_a01	TrAa01ScJC	$\text{HOCH}_2\text{CO}_3\text{H}(\text{aq}) + h\nu \rightarrow \text{HCHO}(\text{aq}) + \text{OH}(\text{aq}) + \text{HO}_2(\text{aq}) + \text{CO}_2(\text{aq})$	2.33*xaer(01)*jx(ip_CH3OOH)	Rickard (2022)
PH4205_a01	TrAa01ScJC	$\text{CH}_3\text{CHO}(\text{aq}) + h\nu \rightarrow \text{CH}_3\text{OO}(\text{aq}) + \text{HO}_2(\text{aq}) + \text{CO}(\text{aq})$	2.33*xaer(01)*jx(ip_CH3CHO)	Sander et al. (2014)
PH4206_a01	TrAa01ScJC	$\text{CH}_2\text{OOHCHO}(\text{aq}) + h\nu \rightarrow \text{OH}(\text{aq}) + \text{HCHO}(\text{aq}) + \text{CO}(\text{aq}) + \text{HO}_2(\text{aq})$	2.33*xaer(01)*(jx(ip_CH3OOH) + jx(ip_HOCH2CHO))	Sander et al. (2019)
PH4207a_a01	TrAa01ScJC	$\text{CH}_2\text{OHCHO}(\text{aq}) + h\nu \rightarrow \text{HCHO}(\text{aq}) + 2 \text{HO}_2(\text{aq}) + \text{CO}(\text{aq})$	2.33*xaer(01)*jx(ip_HOCH2CHO) *0.83	Sander et al. (2014)*
PH4207b_a01	TrAa01ScJC	$\text{CH}_2\text{OHCHO}(\text{aq}) + h\nu \rightarrow \text{OH}(\text{aq}) + .6 \text{ HCHO}(\text{aq}) + .6 \text{ CO}(\text{aq}) + .6 \text{ HO}_2(\text{aq}) + .2 \text{ GLYOX}(\text{aq}) + .2 \text{ CH}_2\text{OHCHO}(\text{aq})$	2.33*xaer(01)*jx(ip_HOCH2CHO) *0.07	Sander et al. (2014)*
PH4207c_a01	TrAa01ScJC	$\text{CH}_2\text{OHCHO}(\text{aq}) + h\nu \rightarrow \text{CH}_3\text{OH}(\text{aq}) + \text{CO}(\text{aq})$	2.33*xaer(01)*jx(ip_HOCH2CHO) *0.10	Sander et al. (2014)*
PH4208_a01	TrAa01ScJC	$\text{CHOCOOH}(\text{aq}) + h\nu \rightarrow 2 \text{HO}_2(\text{aq}) + \text{CO}(\text{aq}) + \text{CO}_2(\text{aq})$	2.33*xaer(01)*jx(ip_MGLYOX)	Rickard (2022)
PH4209_a01	TrAa01ScJC	$\text{GLYOX}(\text{aq}) + h\nu \rightarrow 2 \text{CO}(\text{aq}) + 2 \text{HO}_2(\text{aq})$	2.33*xaer(01)*jx(ip_GLYOX)	Sander et al. (2014)
PH4210a_a01	TrAa01ScJC	$\text{HOOCCOOH}(\text{aq}) + h\nu \rightarrow \text{CO}_2(\text{aq}) + \text{HCOOH}(\text{aq})$	2.33*xaer(01)*0.72*jx(ip_HOOCCOOH)	Yamamoto and Back (1985)
PH4210b_a01	TrAa01ScJC	$\text{HOOCCOOH}(\text{aq}) + h\nu \rightarrow \text{CO}_2(\text{aq}) + \text{CO}(\text{aq}) + \text{H}_2\text{O}(\text{aq})$	2.33*xaer(01)*0.28*jx(ip_HOOCCOOH)	Yamamoto and Back (1985)
PH4211_a01	TrAa01ScJC	$\text{CHOCHOHOH}(\text{aq}) + h\nu \rightarrow \text{HCOOH}(\text{aq}) + 2 \text{HO}_2(\text{aq}) + \text{CO}(\text{aq})$	2.33*xaer(01)*jx(ip_HOCH2CHO)	Sander et al. (2014)*
PH4300_a01	TrAa01ScJC	$\text{CH}_3\text{COCH}_2\text{O}_2\text{H}(\text{aq}) + h\nu \rightarrow \text{CH}_3\text{COOO}(\text{aq}) + \text{HCHO}(\text{aq}) + \text{OH}(\text{aq})$	2.33*xaer(01)*(jx(ip_CH3OOH) + 0.65*0.11*jx(ip_CHOH))	see note*
PH4301_a01	TrAa01ScJC	$\text{iC}_3\text{H}_7\text{OOH}(\text{aq}) + h\nu \rightarrow \text{CH}_3\text{COCH}_3(\text{aq}) + \text{HO}_2(\text{aq}) + \text{OH}(\text{aq})$	2.33*xaer(01)*jx(ip_CH3OOH)	see note*

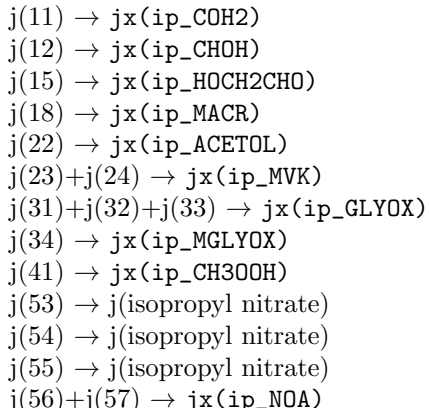
Table 2: Photolysis reactions (... continued)

#	labels	reaction	rate coefficient	reference
PH4302_a01	TrAa01ScJC	$\text{CH}_3\text{COCH}_2\text{OH}(\text{aq}) + h\nu \rightarrow .5 \text{ OH}(\text{aq}) + .5 \text{ HCHO}(\text{aq}) + .5 \text{ CO}(\text{aq}) + .5 \text{ HCHO}(\text{aq}) + .5 \text{ HO}_2(\text{aq}) + .5 \text{ CH}_2\text{OHCO}_3(\text{aq}) + .5 \text{ CH}_3\text{OO}(\text{aq})$	$2.33*\text{xaer}(01)*0.65*0.11*\text{jx(ip_CHOH)}$	Sander et al. (2014)*
PH4303_a01	TrAa01ScJC	$\text{CH}_3\text{C(O)CHO}(\text{aq}) + h\nu \rightarrow \text{OH}(\text{aq}) + \text{HCHO}(\text{aq}) + \text{CO}(\text{aq}) + \text{CO}(\text{aq}) + \text{HO}_2(\text{aq})$	$2.33*\text{xaer}(01)*\text{jx(ip_MGLYOX)}$	Sander et al. (2014)*
PH11200_a01	TrAa01JFe	$\text{FeOH}^{2+}(\text{aq}) + h\nu \rightarrow \text{Fe}^{2+}(\text{aq}) + \text{OH}(\text{aq})$	$\text{xaer}(01)*4.51\text{E}-3*0.312$	Herrmann et al. (2000)
PH11201_a01	TrAa01JFe	$\text{Fe}(\text{OH})_2^+(\text{aq}) + h\nu \rightarrow \text{Fe}^{2+}(\text{aq}) + \text{OH}(\text{aq}) + \text{OH}^-(\text{aq})$	$\text{xaer}(01)*5.77\text{E}-3*0.255$	Herrmann et al. (2000)
PH11800_a01	TrAa01JFeS	$\text{FeSO}_4^+(\text{aq}) + h\nu \rightarrow \text{Fe}^{2+}(\text{aq}) + \text{SO}_4^-(\text{aq})$	$\text{xaer}(01)*6.43\text{E}-3*7.9\text{E}-3$	Herrmann et al. (2000)

General notes

j -values are calculated with an external module (e.g., JVAL) and then supplied to the MECCA chemistry.

Values that originate from the Master Chemical Mechanism (MCM) by Rickard (2022) are translated according in the following way:



Specific notes

J41006: product distribution as for HNO₄

- J42004: Quantum yields from Burkholder et al. (2015).
- J42005a: Quantum yields from Burkholder et al. (2015).
- J42005b: Quantum yields from Burkholder et al. (2015).
- J42005c: Quantum yields from Burkholder et al. (2015).
- J42007: It is assumed that $J(\text{PHAN})$ is the same as $J(\text{PAN})$.
- J42017: Enhancement of j according to Müller et al. (2014).
- J42020: It is assumed that $j(\text{NO}_3\text{CH}_2\text{CHO})$ is the same as $j(\text{PAN})$.
- J42021: In analogy to what is assumed for $\text{CH}_3\text{O}_2\text{NO}_2$ photolysis as in (Sander et al., 2014).
- J43002: Following von Kuhlmann et al. (2003), we use $j(\text{CH}_3\text{COCH}_2\text{OH}) = 0.11*\text{jx(ip_CHOH)}$. As an additional factor, the quantum yield of 0.65 is taken from Orlando et al. (1999a).
- J43006: Following von Kuhlmann et al. (2003), we use $J(\text{iC}_3\text{H}_7\text{ONO}_2) = 3.7*\text{jx(ip_PAN)}$.
- J43018: One third of the acetaldehyde channel is considered to be CH_2CHOH according to Hjorth (2002) EUPHORE Report.
- J43024: Assuming $J(\text{C}_3\text{H}_7\text{ONO}_2) = 0.59 \times J(\text{iC}_3\text{H}_7\text{ONO}_2)$, consistent with the photolysis rate coefficients used in the MCM (Rickard, 2022).
- J43025a: Photolysis frequencies very similar to the ones of CH_3CHO .
- J43025b: Photolysis frequencies very similar to the ones of CH_3CHO .
- J43400: KDEC C3DIALO $\rightarrow \text{GLYOX} + \text{CO} + \text{HO}_2$
- J44004: It is assumed that $J(\text{BIACET})$ is 2.15 times larger than $J(\text{MGLYOX})$, consistent with the photolysis rate coefficients used in the MCM (Rickard, 2022).
- J44005a: It is assumed that $J(\text{LC4H}_9\text{NO}_3)$ is the same as $J(\text{iC}_3\text{H}_7\text{ONO}_2)$.
- J44005b: It is assumed that $J(\text{LC4H}_9\text{NO}_3)$ is the same as $J(\text{iC}_3\text{H}_7\text{ONO}_2)$.
- J44006: It is assumed that $J(\text{MPAN})$ is the same as $J(\text{PAN})$.
- J44009: It is assumed that $J(\text{MACROOH})$ is 2.77 times larger than $J(\text{HOCH}_2\text{CHO})$, consistent with the photolysis rate coefficients used in the MCM (Rickard, 2022).
- J44010: It is assumed that $J(\text{MACROH})$ is 2.77 times larger than $J(\text{HOCH}_2\text{CHO})$, consistent with the photolysis rate coefficients used in the MCM (Rickard, 2022).

- J44015: It is assumed that J(BIACETO_H) is 2.15 times larger than J(MGLYO_X), consistent with the photolysis rate coefficients used in the MCM (Rickard, 2022).
- J44017a: CO-channel yielding CH₃COCH which upon reaction with O₂ produces an excited Criegee Intermediate assumed to be similar to MGLOOA in MCM. MGLOOA is produced also in other reactions and is substituted by its decomposition products. Furthermore, the stabilized Criegge Intermediate is assumed to solely react with water.
- J44025: J values only for the secondary nitrate.
- J44026: Like for LMEKNO₃ photolysis
- J44027: 2.84*J_IC3H7NO₃ like for other tertiary alkyl nitrates (see J4505). Enhancement of J according to Müller et al. (2014).
- J44037b: Channel which produces just vinyl alcohol and not a larger enol via keto-enol photo-tautomerization.
- J44043: The resulting vinyl peroxy radical is assumed to mostly form with HO₂ a labile hydroperoxide (see ketene formation). The products are further simplified.
- J44044: 1,5-H-shift for the resulting vinyl peroxy radical assumed to be dominant.
- J44046a: Simplified oxidation.
- J44400b: KDEC MALDIALO → GLYO_X + GLYO_X + HO₂
- J44401: KDEC BZFUO → CO₁₄O₃CHO + HO₂
- J44403: KDEC NBZFUO → 0.5 CO₁₄O₃CHO + 0.5 NO₂ + 0.5 NBZFUONE + 0.5 HO₂
- J44404b: KDEC MALDIALCO₂ → 0.6 MALANHY + HO₂ + 0.4 GLYO_X + 0.4 CO
- J44407: KDEC MALANHYO → HCOCOHC_O₃
- J44414: KDEC MECOACETO → CH₃CO₃ + HCHO
- J45003: It is assumed that J(LISOPACNO₃) = 0.59 × J(iC₃H₇ONO₂), consistent with the photolysis rate coefficients used in the MCM (Rickard, 2022).
- J45005: It is assumed that J(ISOPBNO₃) = 2.84 × J(iC₃H₇ONO₂), consistent with the photolysis rate coefficients used in the MCM (Rickard, 2022).
- J45007: It is assumed that J(ISOPDNO₃) is the same as J(iC₃H₇ONO₂).
- J45009: 0.59*J_IC3H7NO₃ like for other primary alkyl nitrates (see J4503). Enhancement of J according to Müller et al. (2014).
- J45015: Consistent with the MCM (Rickard, 2022), we assume that J(HCOC₅) is half as large as J(MVK). With exception of HOCH₂CO the products of MACO₂ decomposition without CO₂.
- J45032: approximation with 4-oxo-pentenal photolysis combining results of Thüner et al(2004) and Xiang et al(2007)
- J45402: KDEC C₅DIALO → MALDIAL + CO + HO₂
- J45407: KDEC TLFUONE → 0.6 C₅CO₁₄O₂ + 0.6 HO₂ + 0.4 TLFUONE
- J45410: KDEC MMALANHYO → CO₂H₃CO₃
- J45411: KDEC C₅DICARBO → MGLYO_X + GLYO_X + HO₂
- J45412: KDEC NTLFUO → ACCOMECHO + NO₂
- J45414: KDEC C₅CO₁₄CO₂ → 0.83 MALANHY + 0.83 CH₃ + .17 MGLYO_X + .17 HO₂ + .17 CO + .17 CO₂
- J45415: KDEC TLFUO → ACCOMECHO + HO₂
- J46400: KDEC PHENO → 0.71 MALDALCO₂H + 0.71 GLYO_X + 0.29 PBZQONE + HO₂
- J46403: KDEC NDNPHENO → NC₄DCO₂H + HNO₃ + CO + CO + NO₂
- J46404: KDEC BZBIPERO → GLYO_X + HO₂ + 0.5 BZFUONE + 0.5 BZFUONE
- J46405: new channel created for nitrophenol decomposition
- J46406: new channel created for nitrophenol decomposition
- J46412: KDEC NNCATECO → NC₄DCO₂H + HCOCO₂H + NO₂
- J46415: KDEC NCATECO → NC₄DCO₂H + HCOCO₂H + HO₂
- J46416: KDEC PBZQO → C₅CO₂OHCO₃
- J46418: KDEC BZBIPERO → GLYO_X + HO₂ + 0.5 BZFUONE + 0.5 BZFUONE
- J46419: KDEC NBZQO → C₆CO₄DB + NO₂
- J46422: KDEC DNPHEN_O → NC₄DCO₂H + HCOCO₂H + NO₂
- J46425: KDEC BZEMUCO → 0.5 EPXC4DIAL + .5 GLYO_X + .5 HO₂ + .5 C₃DIALO₂ + .5 C₃2OH₁₃CO
- J46429: new channel
- J47401: KROPRIM*O₂ fast reaction C₆H₅CH₂O = BENZAL + HO₂
- J47402: KROPRIM*O₂ fast reaction C₆H₅CH₂O = BENZAL + HO₂
- J47404: KDEC TLBIPERO → 0.6 GLYO_X + 0.4 MGLYO_X + HO₂ + 0.2 C₄MDIAL + 0.2 C₅DICARB + 0.2 TLFUONE + 0.2 BZFUONE + 0.2 MALDIAL
- J47405: KDEC TLBIPERO → 0.6 GLYO_X + 0.4 MGLYO_X + HO₂ + 0.2 C₄MDIAL + 0.2 C₅DICARB + 0.2 TLFUONE + 0.2 BZFUONE + 0.2 MALDIAL
- J47407: KDEC CRESO → 0.68 C₅CO₁₄OH + 0.68 GLYO_X + HO₂ + 0.32 PTLQONE
- J47408a: KDEC CRESO → 0.68 C₅CO₁₄OH + 0.68 GLYO_X + HO₂ + 0.32 PTLQONE
- J47408b: KDEC NCRESO → C₅CO₁₄OH + GLYO_X + NO₂

J47409: Using J for 3-methyl-2-nitrophenol.	J48400: KDEC STYRENO → HO ₂ + HCHO + BEN-ZAL	PH4207b_a01: Quantum yields from Burkholder et al. (2015). HCOCH ₂ O ₂ decomposes directly to .6 HCHO + .6 CO + .6 HO ₂ + .2 GLYOX + .2 HOCH ₂ CHO
J47412: KDEC TLEMUCO → 0.5 C3DIALO2 + 0.5 CO ₂ H3CHO + 0.5 EPXC4DIAL + 0.5 MGLYOX + 0.5 HO ₂	J40203b: Substituted vinyl alcohol in analogy to CH ₃ CHO photolysis.	PH4207c_a01: Quantum yields from Burkholder et al. (2015).
J47417: Using J for 3-methyl-2-nitrophenol.	J7301: The quantum yields are recommended by Burkholder et al. (2015) for $\lambda > 300\text{nm}$ and used here for the entire spectrum.	PH4211_a01: Assumed in analogy to the main channel for $j(\text{HOCH}_2\text{CHO})$.
J47418: new channel	PH2100_a01: 2.33 times the gas-phase value	PH4300_a01: 2.33* k from the gas-phase reaction, CH ₃ CO directly reacts with O ₂ to form CH ₃ CO ₃
J47419: Using J for 3-methyl-2-nitrophenol.	PH3200_a01: Scaled to J(NO ₂) so that its lifetime is about 10.5 days, as suggested by Zellner et al. (1990).	PH4301_a01: 2.33 * k from the gas-phase reaction,
J47420: new channel	PH4200_a01: CH ₃ CHOHO ₂ is assumed to directly decompose into CH ₃ CHO + HO ₂	PH4302_a01: Following von Kuhlmann et al. (2003), we use $j(\text{CH}_3\text{COCH}_2\text{OH}) = 0.11*jx(ip_{\text{CHOH}})$. As an additional factor, the quantum yield of 0.65 is taken from Orlando et al. (1999a). CH ₃ CO reacts with O ₂ to form OH + HCHO + CO. HOCH ₂ CO reacts with O ₂ to form HOCH ₂ CO ₃
J47422: KDEC NPTLQO → C7CO4DB + NO ₂	PH4201_a01: COOHOO is not formed but directly dissociates into CO ₂ + HO ₂	PH4303_a01: CH ₃ CO reacts with O ₂ to form OH + HCHO + CO
J47423: KDEC PTLQO → C6CO2OHCO ₃	PH4202_a01: assumed to be the same as C ₂ H ₅ OOH + hν	
J47425: KDEC MNNCATECO → NC4MDCO ₂ H + HCOCO ₂ H + NO ₂	PH4207a_a01: Quantum yields from Burkholder et al. (2015).	
J47426: KDEC MNCCATECO → NC4MDCO ₂ H + HCOCO ₂ H + HO ₂		
J47428: KDEC NDNCRESO → NC4MDCO ₂ H + HNO ₃ + CO + CO + NO ₂		
J47429: KDEC DNCRESO → NC4MDCO ₂ H + HCOCO ₂ H + NO ₂		

Table 3: Reversible (Henry's law) equilibria and irreversible ("heterogenous") uptake

#	labels	reaction	rate coefficient	reference
H10000f_a01	TrAa01Sc	O ₂ → O ₂ (aq)	k_exf(01, ind_O2)	see general notes*
H10000b_a01	TrAa01Sc	O ₂ (aq) → O ₂	k_exb(01, ind_O2)	see general notes*
H10001f_a01	TrAa01MblScScm	O ₃ → O ₃ (aq)	k_exf(01, ind_O3)	see general notes*
H10001b_a01	TrAa01MblScScm	O ₃ (aq) → O ₃	k_exb(01, ind_O3)	see general notes*
H21000f_a01	TrAa01Sc	OH → OH(aq)	k_exf(01, ind_OH)	see general notes*
H21000b_a01	TrAa01Sc	OH(aq) → OH	k_exb(01, ind_OH)	see general notes*
H21001f_a01	TrAa01Sc	HO ₂ → HO ₂ (aq)	k_exf(01, ind_HO2)	see general notes*
H21001b_a01	TrAa01Sc	HO ₂ (aq) → HO ₂	k_exb(01, ind_HO2)	see general notes*
H21002f_a01	TrAa01MblScScm	H ₂ O ₂ → H ₂ O ₂ (aq)	k_exf(01, ind_H2O2)	see general notes*
H21002b_a01	TrAa01MblScScm	H ₂ O ₂ (aq) → H ₂ O ₂	k_exb(01, ind_H2O2)	see general notes*
H31000f_a01	TrAa01ScN	NO → NO(aq)	k_exf(01, ind_NO)	see general notes*
H31000b_a01	TrAa01ScN	NO(aq) → NO	k_exb(01, ind_NO)	see general notes*
H31001f_a01	TrAa01ScN	NO ₂ → NO ₂ (aq)	k_exf(01, ind_NO2)	see general notes*
H31001b_a01	TrAa01ScN	NO ₂ (aq) → NO ₂	k_exb(01, ind_NO2)	see general notes*
H31002f_a01	TrAa01ScN	NO ₃ → NO ₃ (aq)	k_exf(01, ind_NO3)	see general notes*
H31002b_a01	TrAa01ScN	NO ₃ (aq) → NO ₃	k_exb(01, ind_NO3)	see general notes*
H32000f_a01	TrAa01MblScScmN	NH ₃ → NH ₃ (aq)	k_exf(01, ind_NH3)	see general notes*
H32000b_a01	TrAa01MblScScmN	NH ₃ (aq) → NH ₃	k_exb(01, ind_NH3)	see general notes*
H32002f_a01	TrAa01ScN	HONO → HONO(aq)	k_exf(01, ind_HONO)	see general notes*
H32002b_a01	TrAa01ScN	HONO(aq) → HONO	k_exb(01, ind_HONO)	see general notes*
H32003f_a01	TrAa01MblScScmN	HNO ₃ → HNO ₃ (aq)	k_exf(01, ind_HN03)	see general notes*
H32003b_a01	TrAa01MblScScmN	HNO ₃ (aq) → HNO ₃	k_exb(01, ind_HN03)	see general notes*
H32004f_a01	TrAa01ScN	HNO ₄ → HNO ₄ (aq)	k_exf(01, ind_HN04)	see general notes*
H32004b_a01	TrAa01ScN	HNO ₄ (aq) → HNO ₄	k_exb(01, ind_HN04)	see general notes*
H320MSf_a01	TrAa01MblScScmN	N ₂ O ₅ → N ₂ O ₅ (aq)	k_exf(01, ind_N205)	see general notes*
H320MSb_a01	TrAa01MblScScmN	N ₂ O ₅ (aq) → N ₂ O ₅	k_exb(01, ind_N205)	see general notes*
H41000f_a01	TrAa01MblScScm	CO ₂ → CO ₂ (aq)	k_exf(01, ind_CO2)	see general notes*
H41000b_a01	TrAa01MblScScm	CO ₂ (aq) → CO ₂	k_exb(01, ind_CO2)	see general notes*
H41001f_a01	TrAa01ScScm	HCHO → HCHO(aq)	k_exf(01, ind_HCHO)	see general notes*
H41001b_a01	TrAa01ScScm	HCHO(aq) → HCHO	k_exb(01, ind_HCHO)	see general notes*
H41002f_a01	TrAa01Sc	CH ₃ O ₂ → CH ₃ OO(aq)	k_exf(01, ind_CH302)	see general notes*
H41002b_a01	TrAa01Sc	CH ₃ OO(aq) → CH ₃ O ₂	k_exb(01, ind_CH302)	see general notes*
H41003f_a01	TrAa01ScScm	HCOOH → HCOOH(aq)	k_exf(01, ind_HC00H)	see general notes*
H41003b_a01	TrAa01ScScm	HCOOH(aq) → HCOOH	k_exb(01, ind_HC00H)	see general notes*

Table 3: Reversible (Henry's law) equilibria and irreversible ("heterogenous") uptake

#	labels	reaction	rate coefficient	reference
H41004f_a01	TrAa01ScScm	$\text{CH}_3\text{OOH} \rightarrow \text{CH}_3\text{OOH(aq)}$	k_exf(01, ind_CH3OOH)	see general notes*
H41004b_a01	TrAa01ScScm	$\text{CH}_3\text{OOH(aq)} \rightarrow \text{CH}_3\text{OOH}$	k_exb(01, ind_CH3OOH)	see general notes*
H41005f_a01	TrAa01Sc	$\text{CH}_3\text{OH} \rightarrow \text{CH}_3\text{OH(aq)}$	k_exf(01, ind_CH3OH)	see general notes*
H41005b_a01	TrAa01Sc	$\text{CH}_3\text{OH(aq)} \rightarrow \text{CH}_3\text{OH}$	k_exb(01, ind_CH3OH)	see general notes*
H41006f_a01	TrAa01Sc	$\text{HOCH}_2\text{OH} \rightarrow \text{HOCH}_2\text{OH(aq)}$	k_exf(01, ind_HOCH2OH)	see general notes*
H41006b_a01	TrAa01Sc	$\text{HOCH}_2\text{OH(aq)} \rightarrow \text{HOCH}_2\text{OH}$	k_exb(01, ind_HOCH2OH)	see general notes*
H41007f_a01	TrAa01Sc	$\text{HOCH}_2\text{OOH} \rightarrow \text{HOCH}_2\text{OOH(aq)}$	k_exf(01, ind_HOCH2OOH)	see general notes*
H41007b_a01	TrAa01Sc	$\text{HOCH}_2\text{OOH(aq)} \rightarrow \text{HOCH}_2\text{OOH}$	k_exb(01, ind_HOCH2OOH)	see general notes*
H41008f_a01	TrAa01Sc	$\text{CO} \rightarrow \text{CO(aq)}$	k_exf(01, ind_CO)	see general notes*
H41008b_a01	TrAa01Sc	$\text{CO(aq)} \rightarrow \text{CO}$	k_exb(01, ind_CO)	see general notes*
H41009MSf_a01	TrAa01ScN	$\text{CH}_3\text{ONO}_2 \rightarrow \text{CH}_3\text{ONO}_2(\text{aq})$	k_exf(01, ind_CH3ONO3)	see general notes*
H41010MSb_a01	TrAa01ScN	$\text{CH}_3\text{ONO}_2(\text{aq}) \rightarrow \text{CH}_3\text{ONO}_2$	k_exb(01, ind_CH3ONO3)	see general notes*
H42000f_a01	TrAa01ScScmC	$\text{CH}_3\text{COOH} \rightarrow \text{CH}_3\text{COOH(aq)}$	k_exf(01, ind_CH3CO2H)	see general notes*
H42000b_a01	TrAa01ScScmC	$\text{CH}_3\text{COOH(aq)} \rightarrow \text{CH}_3\text{COOH}$	k_exb(01, ind_CH3CO2H)	see general notes*
H42001f_a01	TrAa01ScC	$\text{CH}_3\text{CHO} \rightarrow \text{CH}_3\text{CHO(aq)}$	k_exf(01, ind_CH3CHO)	see general notes*
H42001b_a01	TrAa01ScC	$\text{CH}_3\text{CHO(aq)} \rightarrow \text{CH}_3\text{CHO}$	k_exb(01, ind_CH3CHO)	see general notes*
H42002f_a01	TrAa01ScCN	$\text{PAN} \rightarrow \text{PAN(aq)}$	k_exf(01, ind_PAN)	see general notes*
H42002b_a01	TrAa01ScCN	$\text{PAN(aq)} \rightarrow \text{PAN}$	k_exb(01, ind_PAN)	see general notes*
H42003f_a01	TrAa01ScC	$\text{C}_2\text{H}_5\text{OH} \rightarrow \text{CH}_3\text{CH}_2\text{OH(aq)}$	k_exf(01, ind_C2H5OH)	see general notes*
H42003b_a01	TrAa01ScC	$\text{CH}_3\text{CH}_2\text{OH(aq)} \rightarrow \text{C}_2\text{H}_5\text{OH}$	k_exb(01, ind_C2H5OH)	see general notes*
H42004f_a01	TrAa01ScC	$\text{ETHGLY} \rightarrow \text{ETHGLY(aq)}$	k_exf(01, ind_ETHGLY)	see general notes*
H42004b_a01	TrAa01ScC	$\text{ETHGLY(aq)} \rightarrow \text{ETHGLY}$	k_exb(01, ind_ETHGLY)	see general notes*
H42006f_a01	TrAa01ScC	$\text{CH}_3\text{C(O)OO} \rightarrow \text{CH}_3\text{COOO(aq)}$	k_exf(01, ind_CH3C03)	see general notes*
H42006b_a01	TrAa01ScC	$\text{CH}_3\text{COOO(aq)} \rightarrow \text{CH}_3\text{C(O)OO}$	k_exb(01, ind_CH3C03)	see general notes*
H42007f_a01	TrAa01ScC	$\text{HOCH}_2\text{CHO} \rightarrow \text{CH}_2\text{OHCHO(aq)}$	k_exf(01, ind_HOCH2CHO)	see general notes*
H42007b_a01	TrAa01ScC	$\text{CH}_2\text{OHCHO(aq)} \rightarrow \text{HOCH}_2\text{CHO}$	k_exb(01, ind_HOCH2CHO)	see general notes*
H42008f_a01	TrAa01ScC	$\text{GLYOX} \rightarrow \text{GLYOX(aq)}$	k_exf(01, ind_GLYOX)	see general notes*
H42008b_a01	TrAa01ScC	$\text{GLYOX(aq)} \rightarrow \text{GLYOX}$	k_exb(01, ind_GLYOX)	see general notes*
H42009f_a01	TrAa01ScC	$\text{CH}_3\text{C(O)OOH} \rightarrow \text{CH}_3\text{C(O)OOH(aq)}$	k_exf(01, ind_CH3C03H)	see general notes*
H42009b_a01	TrAa01ScC	$\text{CH}_3\text{C(O)OOH(aq)} \rightarrow \text{CH}_3\text{C(O)OOH}$	k_exb(01, ind_CH3C03H)	see general notes*
H42010f_a01	TrAa01ScC	$\text{HOCH}_2\text{CO}_3\text{H} \rightarrow \text{HOCH}_2\text{CO}_3\text{H(aq)}$	k_exf(01, ind_HOCH2CO3H)	see general notes*
H42010b_a01	TrAa01ScC	$\text{HOCH}_2\text{CO}_3\text{H(aq)} \rightarrow \text{HOCH}_2\text{CO}_3\text{H}$	k_exb(01, ind_HOCH2CO3H)	see general notes*
H42011f_a01	TrAa01ScC	$\text{C}_2\text{H}_5\text{OOH} \rightarrow \text{C}_2\text{H}_5\text{OOH(aq)}$	k_exf(01, ind_C2H5OOH)	see general notes*
H42011b_a01	TrAa01ScC	$\text{C}_2\text{H}_5\text{OOH(aq)} \rightarrow \text{C}_2\text{H}_5\text{OOH}$	k_exb(01, ind_C2H5OOH)	see general notes*
H42012f_a01	TrAa01ScC	$\text{HOOCOOH} \rightarrow \text{HOOCOOH(aq)}$	k_exf(01, ind_HOOCOOH)	see general notes*

Table 3: Reversible (Henry's law) equilibria and irreversible ("heterogenous") uptake

#	labels	reaction	rate coefficient	reference
H42012b_a01	TrAa01ScC	$\text{HOOCOOH}(\text{aq}) \rightarrow \text{HOOCOOH}$	$k_{\text{exb}}(01, \text{ind_HOOCOOH})$	see general notes*
H42013f_a01	TrAa01ScC	$\text{HOOCCH}_2\text{CO}_2\text{H} \rightarrow \text{HOOCCH}_2\text{CO}_2\text{H}(\text{aq})$	$k_{\text{exf}}(01, \text{ind_HOOCCH}_2\text{CO}_2\text{H})$	see general notes*
H42013b_a01	TrAa01ScC	$\text{HOOCCH}_2\text{CO}_2\text{H}(\text{aq}) \rightarrow \text{HOOCCH}_2\text{CO}_2\text{H}$	$k_{\text{exb}}(01, \text{ind_HOOCCH}_2\text{CO}_2\text{H})$	see general notes*
H42014f_a01	TrAa01ScC	$\text{HOCH}_2\text{CO}_2\text{H} \rightarrow \text{HOCH}_2\text{CO}_2\text{H}(\text{aq})$	$k_{\text{exf}}(01, \text{ind_HOCH}_2\text{CO}_2\text{H})$	see general notes*
H42014b_a01	TrAa01ScC	$\text{HOCH}_2\text{CO}_2\text{H}(\text{aq}) \rightarrow \text{HOCH}_2\text{CO}_2\text{H}$	$k_{\text{exb}}(01, \text{ind_HOCH}_2\text{CO}_2\text{H})$	see general notes*
H42015f_a01	TrAa01ScC	$\text{HCOCO}_2\text{H} \rightarrow \text{CHOCOOH}(\text{aq})$	$k_{\text{exf}}(01, \text{ind_HCOCO}_2\text{H})$	see general notes*
H42015b_a01	TrAa01ScC	$\text{CHOCOOH}(\text{aq}) \rightarrow \text{HCOCO}_2\text{H}$	$k_{\text{exb}}(01, \text{ind_HCOCO}_2\text{H})$	see general notes*
H42017f_a01	TrAa01ScCN	$\text{C}_2\text{H}_5\text{ONO}_2 \rightarrow \text{C}_2\text{H}_5\text{ONO}_2(\text{aq})$	$k_{\text{exf}}(01, \text{ind_C2H5N03})$	see general notes*
H42017b_a01	TrAa01ScCN	$\text{C}_2\text{H}_5\text{ONO}_2(\text{aq}) \rightarrow \text{C}_2\text{H}_5\text{ONO}_2$	$k_{\text{exb}}(01, \text{ind_C2H5N03})$	see general notes*
H42018f_a01	TrAa01ScCN	$\text{CH}_3\text{CN} \rightarrow \text{CH}_3\text{CN}(\text{aq})$	$k_{\text{exf}}(01, \text{ind_CH3CN})$	see general notes*
H42018b_a01	TrAa01ScCN	$\text{CH}_3\text{CN}(\text{aq}) \rightarrow \text{CH}_3\text{CN}$	$k_{\text{exb}}(01, \text{ind_CH3CN})$	see general notes*
H42019f_a01	TrAa01ScC	$\text{HOCH}_2\text{CHOHOH} \rightarrow \text{CH}_2\text{OHCHOHOH}(\text{aq})$	$k_{\text{exf}}(01, \text{ind_HOCH}_2\text{CHOHOH})$	see general notes*
H42019b_a01	TrAa01ScC	$\text{CH}_2\text{OHCHOHOH}(\text{aq}) \rightarrow \text{HOCH}_2\text{CHOHOH}$	$k_{\text{exb}}(01, \text{ind_HOCH}_2\text{CHOHOH})$	see general notes*
H42020f_a01	TrAa01ScC	$\text{CH}_3\text{CHOHOH} \rightarrow \text{CH}_3\text{CHOHOH}(\text{aq})$	$k_{\text{exf}}(01, \text{ind_CH3CHOHOH})$	see general notes*
H42020b_a01	TrAa01ScC	$\text{CH}_3\text{CHOHOH}(\text{aq}) \rightarrow \text{CH}_3\text{CHOHOH}$	$k_{\text{exb}}(01, \text{ind_CH3CHOHOH})$	see general notes*
H42021f_a01	TrAa01ScC	$\text{CHOHOHCOOH} \rightarrow \text{CHOHOHCOOH}(\text{aq})$	$k_{\text{exf}}(01, \text{ind_CHOHOHCOOH})$	see general notes*
H42021b_a01	TrAa01ScC	$\text{CHOHOHCOOH}(\text{aq}) \rightarrow \text{CHOHOHCOOH}$	$k_{\text{exb}}(01, \text{ind_CHOHOHCOOH})$	see general notes*
H42022f_a01	TrAa01ScC	$\text{CHOHOHCHOHOH} \rightarrow \text{CHOHOHCHOHOH}(\text{aq})$	$k_{\text{exf}}(01, \text{ind_CHOHOHCHOHOH})$	see general notes*
H42022b_a01	TrAa01ScC	$\text{CHOHOHCHOHOH}(\text{aq}) \rightarrow \text{CHOHOHCHOHOH}$	$k_{\text{exb}}(01, \text{ind_CHOHOHCHOHOH})$	see general notes*
H42023f_a01	TrAa01ScC	$\text{HOOCH}_2\text{CHO} \rightarrow \text{CH}_2\text{OOHCHO}(\text{aq})$	$k_{\text{exf}}(01, \text{ind_HOOCH}_2\text{CHO})$	see general notes*
H42023b_a01	TrAa01ScC	$\text{CH}_2\text{OOHCHO}(\text{aq}) \rightarrow \text{HOOCH}_2\text{CHO}$	$k_{\text{exb}}(01, \text{ind_HOOCH}_2\text{CHO})$	see general notes*
H42024f_a01	TrAa01ScC	$\text{CHOCHOHOH} \rightarrow \text{CHOCHOHOH}(\text{aq})$	$k_{\text{exf}}(01, \text{ind_CHOCHOHOH})$	see general notes*
H42024b_a01	TrAa01ScC	$\text{CHOCHOHOH}(\text{aq}) \rightarrow \text{CHOCHOHOH}$	$k_{\text{exb}}(01, \text{ind_CHOCHOHOH})$	see general notes*
H42025f_a01	TrAa01ScC	$\text{HOOCH}_2\text{CHOHOH} \rightarrow \text{HOOCH}_2\text{CHOHOH}(\text{aq})$	$k_{\text{exf}}(01, \text{ind_HOOCH}_2\text{CHOHOH})$	see general notes*
H42025b_a01	TrAa01ScC	$\text{HOOCH}_2\text{CHOHOH}(\text{aq}) \rightarrow \text{HOOCH}_2\text{CHOHOH}$	$k_{\text{exb}}(01, \text{ind_HOOCH}_2\text{CHOHOH})$	see general notes*
H42026f_a01	TrAa01ScC	$\text{CH}_2\text{CO} \rightarrow \text{CH}_2\text{CO}(\text{aq})$	$k_{\text{exf}}(01, \text{ind_CH}_2\text{CO})$	see general notes*
H42026b_a01	TrAa01ScC	$\text{CH}_2\text{CO}(\text{aq}) \rightarrow \text{CH}_2\text{CO}$	$k_{\text{exb}}(01, \text{ind_CH}_2\text{CO})$	see general notes*
H42027f_a01	TrAa01ScC	$\text{CH}_3\text{CHOHOOH} \rightarrow \text{CH}_3\text{CHOHOOH}(\text{aq})$	$k_{\text{exf}}(01, \text{ind_CH}_3\text{CHOHOOH})$	see general notes*
H42027b_a01	TrAa01ScC	$\text{CH}_3\text{CHOHOOH}(\text{aq}) \rightarrow \text{CH}_3\text{CHOHOOH}$	$k_{\text{exb}}(01, \text{ind_CH}_3\text{CHOHOOH})$	see general notes*
H42028f_a01	TrAa01ScCN	$\text{ETHOHN03} \rightarrow \text{ETHOHN03}(\text{aq})$	$k_{\text{exf}}(01, \text{ind_ETHOHN03})$	see general notes*
H42028b_a01	TrAa01ScCN	$\text{ETHOHN03}(\text{aq}) \rightarrow \text{ETHOHN03}$	$k_{\text{exb}}(01, \text{ind_ETHOHN03})$	see general notes*

Table 3: Reversible (Henry's law) equilibria and irreversible ("heterogenous") uptake

#	labels	reaction	rate coefficient	reference
H42029f_a01	TrAa01ScC	$\text{HCOCO}_3\text{H} \rightarrow \text{HCOCO}_3\text{H(aq)}$	k_exf(01, ind_HCOCO3H)	see general notes*
H42029b_a01	TrAa01ScC	$\text{HCOCO}_3\text{H(aq)} \rightarrow \text{HCOCO}_3\text{H}$	k_exb(01, ind_HCOCO3H)	see general notes*
H42030f_a01	TrAa01ScC	$\text{HOOCH}_2\text{CO}_3\text{H} \rightarrow \text{HOOCH}_2\text{CO}_3\text{H(aq)}$	k_exf(01, ind_HOOCH2CO3H)	see general notes*
H42030b_a01	TrAa01ScC	$\text{HOOCH}_2\text{CO}_3\text{H(aq)} \rightarrow \text{HOOCH}_2\text{CO}_3\text{H}$	k_exb(01, ind_HOOCH2CO3H)	see general notes*
H42031f_a01	TrAa01ScC	$\text{HYETHO}_2\text{H} \rightarrow \text{HYETHO}_2\text{H(aq)}$	k_exf(01, ind_HYETHO2H)	see general notes*
H42031b_a01	TrAa01ScC	$\text{HYETHO}_2\text{H(aq)} \rightarrow \text{HYETHO}_2\text{H}$	k_exb(01, ind_HYETHO2H)	see general notes*
H42032f_a01	TrAa01ScCN	$\text{PHAN} \rightarrow \text{PHAN(aq)}$	k_exf(01, ind_PHAN)	see general notes*
H42032b_a01	TrAa01ScCN	$\text{PHAN(aq)} \rightarrow \text{PHAN}$	k_exb(01, ind_PHAN)	see general notes*
H43000f_a01	TrAa01ScC	$\text{CH}_3\text{COCH}_3 \rightarrow \text{CH}_3\text{COCH}_3\text{(aq)}$	k_exf(01, ind_CH3COCH3)	see general notes*
H43000b_a01	TrAa01ScC	$\text{CH}_3\text{COCH}_3\text{(aq)} \rightarrow \text{CH}_3\text{COCH}_3$	k_exb(01, ind_CH3COCH3)	see general notes*
H43001f_a01	TrAa01ScC	$\text{MGLYOX} \rightarrow \text{CH}_3\text{C(O)CHO(aq)}$	k_exf(01, ind_MGLYOX)	see general notes*
H43001b_a01	TrAa01ScC	$\text{CH}_3\text{C(O)CHO(aq)} \rightarrow \text{MGLYOX}$	k_exb(01, ind_MGLYOX)	see general notes*
H43002f_a01	TrAa01ScC	$\text{CH}_3\text{COCO}_2\text{H} \rightarrow \text{CH}_3\text{COCOOH(aq)}$	k_exf(01, ind_CH3COCO2H)	see general notes*
H43002b_a01	TrAa01ScC	$\text{CH}_3\text{COCOOH(aq)} \rightarrow \text{CH}_3\text{COCO}_2\text{H}$	k_exb(01, ind_CH3COCO2H)	see general notes*
H43003f_a01	TrAa01ScC	$\text{CH}_3\text{COCHOHOH} \rightarrow \text{CH}_3\text{COCHOHOH(aq)}$	k_exf(01, ind_CH3COCHOHOH)	see general notes*
H43003b_a01	TrAa01ScC	$\text{CH}_3\text{COCHOHOH(aq)} \rightarrow \text{CH}_3\text{COCHOHOH}$	k_exb(01, ind_CH3COCHOHOH)	see general notes*
H43005f_a01	TrAa01ScC	$\text{IPROPOL} \rightarrow \text{IPROPOL(aq)}$	k_exf(01, ind_IPROPOL)	see general notes*
H43005b_a01	TrAa01ScC	$\text{IPROPOL(aq)} \rightarrow \text{IPROPOL}$	k_exb(01, ind_IPROPOL)	see general notes*
H43006f_a01	TrAa01ScC	$\text{CH}_3\text{COCH}_2\text{O}_2\text{H} \rightarrow \text{CH}_3\text{COCH}_2\text{O}_2\text{H(aq)}$	k_exf(01, ind_HYPERACET)	see general notes*
H43006b_a01	TrAa01ScC	$\text{CH}_3\text{COCH}_2\text{O}_2\text{H(aq)} \rightarrow \text{CH}_3\text{COCH}_2\text{O}_2\text{H}$	k_exb(01, ind_HYPERACET)	see general notes*
H43007f_a01	TrAa01ScC	$\text{iC}_3\text{H}_7\text{OOH} \rightarrow \text{iC}_3\text{H}_7\text{OOH(aq)}$	k_exf(01, ind_IC3H7OOH)	see general notes*
H43007b_a01	TrAa01ScC	$\text{iC}_3\text{H}_7\text{OOH(aq)} \rightarrow \text{iC}_3\text{H}_7\text{OOH}$	k_exb(01, ind_IC3H7OOH)	see general notes*
H43008f_a01	TrAa01ScC	$\text{HCOCOCH}_2\text{OOH} \rightarrow \text{HCOCOCH}_2\text{OOH(aq)}$	k_exf(01, ind_ALCOCH2OOH)	see general notes*
H43008b_a01	TrAa01ScC	$\text{HCOCOCH}_2\text{OOH(aq)} \rightarrow \text{HCOCOCH}_2\text{OOH}$	k_exb(01, ind_ALCOCH2OOH)	see general notes*
H43009f_a01	TrAa01ScC	$\text{C}_3\text{OH}_1\text{H}_1\text{CO} \rightarrow \text{C}_3\text{OH}_1\text{H}_1\text{CO(aq)}$	k_exf(01, ind_C3OH1H1CO)	see general notes*
H43009b_a01	TrAa01ScC	$\text{C}_3\text{OH}_1\text{H}_1\text{CO(aq)} \rightarrow \text{C}_3\text{OH}_1\text{H}_1\text{CO}$	k_exb(01, ind_C3OH1H1CO)	see general notes*
H43010f_a01	TrAa01ScC	$\text{HCOCOCHO} \rightarrow \text{HCOCOCHO(aq)}$	k_exf(01, ind_C33CO)	see general notes*
H43010b_a01	TrAa01ScC	$\text{HCOCOCHO(aq)} \rightarrow \text{HCOCOCHO}$	k_exb(01, ind_C33CO)	see general notes*
H43011f_a01	TrAa01ScC	$\text{C}_3\text{DIALOOH} \rightarrow \text{C}_3\text{DIALOOH(aq)}$	k_exf(01, ind_C3DIALOOH)	see general notes*
H43011b_a01	TrAa01ScC	$\text{C}_3\text{DIALOOH(aq)} \rightarrow \text{C}_3\text{DIALOOH}$	k_exb(01, ind_C3DIALOOH)	see general notes*
H43012f_a01	TrAa01ScCN	$\text{C}_3\text{PAN1} \rightarrow \text{C}_3\text{PAN1(aq)}$	k_exf(01, ind_C3PAN1)	see general notes*
H43012b_a01	TrAa01ScCN	$\text{C}_3\text{PAN1(aq)} \rightarrow \text{C}_3\text{PAN1}$	k_exb(01, ind_C3PAN1)	see general notes*
H43013f_a01	TrAa01ScCN	$\text{C}_3\text{PAN2} \rightarrow \text{C}_3\text{PAN2(aq)}$	k_exf(01, ind_C3PAN2)	see general notes*
H43013b_a01	TrAa01ScCN	$\text{C}_3\text{PAN2(aq)} \rightarrow \text{C}_3\text{PAN2}$	k_exb(01, ind_C3PAN2)	see general notes*
H43014f_a01	TrAa01ScC	$\text{CH}_3\text{CHCO} \rightarrow \text{CH}_3\text{CHCO(aq)}$	k_exf(01, ind_CH3CHCO)	see general notes*

Table 3: Reversible (Henry's law) equilibria and irreversible ("heterogenous") uptake

#	labels	reaction	rate coefficient	reference
H43014b_a01	TrAa01ScC	$\text{CH}_3\text{CHCO(aq)} \rightarrow \text{CH}_3\text{CHCO}$	$k_{\text{exb}}(01, \text{ind_CH3CHCO})$	see general notes*
H43015f_a01	TrAa01ScCN	$\text{CH}_3\text{COCH}_2\text{OONO}_2$ $\text{CH}_3\text{COCH}_2\text{OONO}_2(\text{aq})$	$\rightarrow k_{\text{exf}}(01, \text{ind_CH3COCH2O2N02})$	see general notes*
H43015b_a01	TrAa01ScCN	$\text{CH}_3\text{COCH}_2\text{OONO}_2(\text{aq})$ $\text{CH}_3\text{COCH}_2\text{OONO}_2$	$\rightarrow k_{\text{exb}}(01, \text{ind_CH3COCH2O2N02})$	see general notes*
H43016f_a01	TrAa01ScC	$\text{CH}_3\text{COCO}_3\text{H} \rightarrow \text{CH}_3\text{COCO}_3\text{H(aq)}$	$k_{\text{exf}}(01, \text{ind_CH3COCO3H})$	see general notes*
H43016b_a01	TrAa01ScC	$\text{CH}_3\text{COCO}_3\text{H(aq)} \rightarrow \text{CH}_3\text{COCO}_3\text{H}$	$k_{\text{exb}}(01, \text{ind_CH3COCO3H})$	see general notes*
H43017f_a01	TrAa01ScC	$\text{HCOCH}_2\text{CHO} \rightarrow \text{HCOCH}_2\text{CHO(aq)}$	$k_{\text{exf}}(01, \text{ind_HCOCH2CHO})$	see general notes*
H43017b_a01	TrAa01ScC	$\text{HCOCH}_2\text{CHO(aq)} \rightarrow \text{HCOCH}_2\text{CHO}$	$k_{\text{exb}}(01, \text{ind_HCOCH2CHO})$	see general notes*
H43018f_a01	TrAa01ScC	$\text{HCOCH}_2\text{CO2H} \rightarrow \text{HCOCH}_2\text{CO2H(aq)}$	$k_{\text{exf}}(01, \text{ind_HCOCH2CO2H})$	see general notes*
H43018b_a01	TrAa01ScC	$\text{HCOCH}_2\text{CO2H(aq)} \rightarrow \text{HCOCH}_2\text{CO2H}$	$k_{\text{exb}}(01, \text{ind_HCOCH2CO2H})$	see general notes*
H43019f_a01	TrAa01ScC	$\text{HCOCH}_2\text{CO3H} \rightarrow \text{HCOCH}_2\text{CO3H(aq)}$	$k_{\text{exf}}(01, \text{ind_HCOCH2CO3H})$	see general notes*
H43019b_a01	TrAa01ScC	$\text{HCOCH}_2\text{CO3H(aq)} \rightarrow \text{HCOCH}_2\text{CO3H}$	$k_{\text{exb}}(01, \text{ind_HCOCH2CO3H})$	see general notes*
H43020f_a01	TrAa01ScC	$\text{HCOCOCH}_2\text{OOH} \rightarrow \text{HCOCOCH}_2\text{OOH(aq)}$	$k_{\text{exf}}(01, \text{ind_HCOCOCH2OOH})$	see general notes*
H43020b_a01	TrAa01ScC	$\text{HCOCOCH}_2\text{OOH(aq)} \rightarrow \text{HCOCOCH}_2\text{OOH}$	$k_{\text{exb}}(01, \text{ind_HCOCOCH2OOH})$	see general notes*
H43021f_a01	TrAa01ScC	$\text{HCOCOHC03H} \rightarrow \text{HCOCOHC03H(aq)}$	$k_{\text{exf}}(01, \text{ind_HCOCOHC03H})$	see general notes*
H43021b_a01	TrAa01ScC	$\text{HCOCOHC03H(aq)} \rightarrow \text{HCOCOHC03H}$	$k_{\text{exb}}(01, \text{ind_HCOCOHC03H})$	see general notes*
H43022f_a01	TrAa01ScCN	$\text{HCOCOHPAN} \rightarrow \text{HCOCOHPAN(aq)}$	$k_{\text{exf}}(01, \text{ind_HCOCOHPAN})$	see general notes*
H43022b_a01	TrAa01ScCN	$\text{HCOCOHPAN(aq)} \rightarrow \text{HCOCOHPAN}$	$k_{\text{exb}}(01, \text{ind_HCOCOHPAN})$	see general notes*
H43023f_a01	TrAa01ScC	$\text{HOC2H4CO2H} \rightarrow \text{HOC2H4CO2H(aq)}$	$k_{\text{exf}}(01, \text{ind_HOC2H4CO2H})$	see general notes*
H43023b_a01	TrAa01ScC	$\text{HOC2H4CO2H(aq)} \rightarrow \text{HOC2H4CO2H}$	$k_{\text{exb}}(01, \text{ind_HOC2H4CO2H})$	see general notes*
H43024f_a01	TrAa01ScC	$\text{HOC2H4CO3H} \rightarrow \text{HOC2H4CO3H(aq)}$	$k_{\text{exf}}(01, \text{ind_HOC2H4CO3H})$	see general notes*
H43024b_a01	TrAa01ScC	$\text{HOC2H4CO3H(aq)} \rightarrow \text{HOC2H4CO3H}$	$k_{\text{exb}}(01, \text{ind_HOC2H4CO3H})$	see general notes*
H43025f_a01	TrAa01ScC	$\text{HOCH}_2\text{COCH}_2\text{OOH}$ $\text{HOCH}_2\text{COCH}_2\text{OOH(aq)}$	$\rightarrow k_{\text{exf}}(01, \text{ind_HOCH2COCH2OOH})$	see general notes*
H43025b_a01	TrAa01ScC	$\text{HOCH}_2\text{COCH}_2\text{OOH(aq)}$ $\text{HOCH}_2\text{COCH}_2\text{OOH}$	$\rightarrow k_{\text{exb}}(01, \text{ind_HOCH2COCH2OOH})$	see general notes*
H43026f_a01	TrAa01ScC	$\text{HOCH}_2\text{COCHO} \rightarrow \text{HOCH}_2\text{COCHO(aq)}$	$k_{\text{exf}}(01, \text{ind_HOCH2COCHO})$	see general notes*
H43026b_a01	TrAa01ScC	$\text{HOCH}_2\text{COCHO(aq)} \rightarrow \text{HOCH}_2\text{COCHO}$	$k_{\text{exb}}(01, \text{ind_HOCH2COCHO})$	see general notes*
H43027f_a01	TrAa01ScC	$\text{HYPROPO2H} \rightarrow \text{HYPROPO2H(aq)}$	$k_{\text{exf}}(01, \text{ind_HYPROPO2H})$	see general notes*
H43027b_a01	TrAa01ScC	$\text{HYPROPO2H(aq)} \rightarrow \text{HYPROPO2H}$	$k_{\text{exb}}(01, \text{ind_HYPROPO2H})$	see general notes*
H43028f_a01	TrAa01ScC	$\text{METACETHO} \rightarrow \text{METACETHO(aq)}$	$k_{\text{exf}}(01, \text{ind_METACETHO})$	see general notes*
H43028b_a01	TrAa01ScC	$\text{METACETHO(aq)} \rightarrow \text{METACETHO}$	$k_{\text{exb}}(01, \text{ind_METACETHO})$	see general notes*
H43029f_a01	TrAa01ScCN	$\text{NOA} \rightarrow \text{NOA(aq)}$	$k_{\text{exf}}(01, \text{ind_NOA})$	see general notes*
H43029b_a01	TrAa01ScCN	$\text{NOA(aq)} \rightarrow \text{NOA}$	$k_{\text{exb}}(01, \text{ind_NOA})$	see general notes*

Table 3: Reversible (Henry's law) equilibria and irreversible ("heterogenous") uptake

#	labels	reaction	rate coefficient	reference
H43030f_a01	TrAa01ScCN	$\text{PR2O2HNO3} \rightarrow \text{PR2O2HNO3(aq)}$	k_exf(01, ind_PR202HN03)	see general notes*
H43030b_a01	TrAa01ScCN	$\text{PR2O2HNO3(aq)} \rightarrow \text{PR2O2HNO3}$	k_exb(01, ind_PR202HN03)	see general notes*
H43031f_a01	TrAa01ScCN	$\text{PROPOLNO3} \rightarrow \text{PROPOLNO3(aq)}$	k_exf(01, ind_PROPOLN03)	see general notes*
H43031b_a01	TrAa01ScCN	$\text{PROPOLNO3(aq)} \rightarrow \text{PROPOLNO3}$	k_exb(01, ind_PROPOLN03)	see general notes*
H43032f_a01	TrAa01ScC	$\text{CH}_3\text{COCH}_2\text{OH} \rightarrow \text{CH}_3\text{COCH}_2\text{OH(aq)}$	k_exf(01, ind_ACETOL)	see general notes*
H43032b_a01	TrAa01ScC	$\text{CH}_3\text{COCH}_2\text{OH(aq)} \rightarrow \text{CH}_3\text{COCH}_2\text{OH}$	k_exb(01, ind_ACETOL)	see general notes*
H44000f_a01	TrAa01ScC	$\text{MACR} \rightarrow \text{MACR(aq)}$	k_exf(01, ind_MACR)	see general notes*
H44000b_a01	TrAa01ScC	$\text{MACR(aq)} \rightarrow \text{MACR}$	k_exb(01, ind_MACR)	see general notes*
H44001f_a01	TrAa01ScC	$\text{MVK} \rightarrow \text{MVK(aq)}$	k_exf(01, ind_MVK)	see general notes*
H44001b_a01	TrAa01ScC	$\text{MVK(aq)} \rightarrow \text{MVK}$	k_exb(01, ind_MVK)	see general notes*
H44002f_a01	TrAa01ScC	$\text{CH}_3\text{COCOCH}_2\text{O}_2 \rightarrow \text{CH}_3\text{COCOCH}_2\text{O}_2(\text{aq})$	k_exf(01, ind_BIACETO2)	see general notes*
H44002b_a01	TrAa01ScC	$\text{CH}_3\text{COCOCH}_2\text{O}_2(\text{aq}) \rightarrow \text{CH}_3\text{COCOCH}_2\text{O}_2$	k_exb(01, ind_BIACETO2)	see general notes*
H44003f_a01	TrAa01ScC	$\text{BIACETOH} \rightarrow \text{BIACETOH(aq)}$	k_exf(01, ind_BIACETOH)	see general notes*
H44003b_a01	TrAa01ScC	$\text{BIACETOH(aq)} \rightarrow \text{BIACETOH}$	k_exb(01, ind_BIACETOH)	see general notes*
H44004f_a01	TrAa01ScC	$\text{CH}_3\text{COCOCH}_2\text{OOH} \rightarrow \text{CH}_3\text{COCOCH}_2\text{OOH(aq)}$	k_exf(01, ind_BIACETOOTH)	see general notes*
H44004b_a01	TrAa01ScC	$\text{CH}_3\text{COCOCH}_2\text{OOH(aq)} \rightarrow \text{CH}_3\text{COCOCH}_2\text{OOH}$	k_exb(01, ind_BIACETOOTH)	see general notes*
H44005f_a01	TrAa01ScC	$\text{BUT2OLO} \rightarrow \text{BUT2OLO(aq)}$	k_exf(01, ind_BUT2OLO)	see general notes*
H44005b_a01	TrAa01ScC	$\text{BUT2OLO(aq)} \rightarrow \text{BUT2OLO}$	k_exb(01, ind_BUT2OLO)	see general notes*
H44006f_a01	TrAa01ScC	$\text{BUT2OLOOH} \rightarrow \text{BUT2OLOOOH(aq)}$	k_exf(01, ind_BUT2OLOOH)	see general notes*
H44006b_a01	TrAa01ScC	$\text{BUT2OLOOOH(aq)} \rightarrow \text{BUT2OLOOOH}$	k_exb(01, ind_BUT2OLOOH)	see general notes*
H44007f_a01	TrAa01ScC	$\text{BZFUUCO} \rightarrow \text{BZFUUCO(aq)}$	k_exf(01, ind_BZFUUCO)	see general notes*
H44007b_a01	TrAa01ScC	$\text{BZFUUCO(aq)} \rightarrow \text{BZFUUCO}$	k_exb(01, ind_BZFUUCO)	see general notes*
H44008f_a01	TrAa01ScC	$\text{BZFUOOH} \rightarrow \text{BZFUOOH(aq)}$	k_exf(01, ind_BZFUOOH)	see general notes*
H44008b_a01	TrAa01ScC	$\text{BZFUOOH(aq)} \rightarrow \text{BZFUOOH}$	k_exb(01, ind_BZFUOOH)	see general notes*
H44009f_a01	TrAa01ScC	$\text{C312COCO3H} \rightarrow \text{C312COCO3H(aq)}$	k_exf(01, ind_C312COCO3H)	see general notes*
H44009b_a01	TrAa01ScC	$\text{C312COCO3H(aq)} \rightarrow \text{C312COCO3H}$	k_exb(01, ind_C312COCO3H)	see general notes*
H44010f_a01	TrAa01ScCN	$\text{C312COPAN} \rightarrow \text{C312COPAN(aq)}$	k_exf(01, ind_C312COPAN)	see general notes*
H44010b_a01	TrAa01ScCN	$\text{C312COPAN(aq)} \rightarrow \text{C312COPAN}$	k_exb(01, ind_C312COPAN)	see general notes*
H44011f_a01	TrAa01ScC	$\text{C413COOOH} \rightarrow \text{C413COOOH(aq)}$	k_exf(01, ind_C413COOOH)	see general notes*
H44011b_a01	TrAa01ScC	$\text{C413COOOH(aq)} \rightarrow \text{C413COOOH}$	k_exb(01, ind_C413COOOH)	see general notes*
H44012f_a01	TrAa01ScC	$\text{C44OOH} \rightarrow \text{C44OOH(aq)}$	k_exf(01, ind_C44OOH)	see general notes*
H44012b_a01	TrAa01ScC	$\text{C44OOH(aq)} \rightarrow \text{C44OOH}$	k_exb(01, ind_C44OOH)	see general notes*
H44013f_a01	TrAa01ScC	$\text{C4CODIAL} \rightarrow \text{C4CODIAL(aq)}$	k_exf(01, ind_C4CODIAL)	see general notes*

Table 3: Reversible (Henry's law) equilibria and irreversible ("heterogenous") uptake

#	labels	reaction	rate coefficient	reference
H44013b_a01	TrAa01ScC	C4CODIAL(aq) → C4CODIAL	k_exb(01, ind_C4CODIAL)	see general notes*
H44014f_a01	TrAa01ScCN	C4PAN5 → C4PAN5(aq)	k_exf(01, ind_C4PAN5)	see general notes*
H44014b_a01	TrAa01ScCN	C4PAN5(aq) → C4PAN5	k_exb(01, ind_C4PAN5)	see general notes*
H44015f_a01	TrAa01ScC	CH ₃ COCHCO → CH ₃ COCHCO(aq)	k_exf(01, ind_CH3COCHCO)	see general notes*
H44015b_a01	TrAa01ScC	CH ₃ COCHCO(aq) → CH ₃ COCHCO	k_exb(01, ind_CH3COCHCO)	see general notes*
H44016f_a01	TrAa01ScC	CH ₃ COCOCO ₂ H → CH ₃ COCOCO ₂ H(aq)	k_exf(01, ind_CH3COCOCO2H)	see general notes*
H44016b_a01	TrAa01ScC	CH ₃ COCOCO ₂ H(aq) → CH ₃ COCOCO ₂ H	k_exb(01, ind_CH3COCOCO2H)	see general notes*
H44017f_a01	TrAa01ScC	CH ₃ COOHCHCHO CH ₃ COOHCHCHO(aq)	→ k_exf(01, ind_CH3COOHCHCHO)	see general notes*
H44017b_a01	TrAa01ScC	CH ₃ COOHCHCHO(aq) CH ₃ COOHCHCHO	→ k_exb(01, ind_CH3COOHCHCHO)	see general notes*
H44018f_a01	TrAa01ScC	CHOC3COO ₂ → CHOC3COO ₂ (aq)	k_exf(01, ind_CHOC3COO2)	see general notes*
H44018b_a01	TrAa01ScC	CHOC3COO ₂ (aq) → CHOC3COO ₂	k_exb(01, ind_CHOC3COO2)	see general notes*
H44019f_a01	TrAa01ScC	CO14O3CHO → CO14O3CHO(aq)	k_exf(01, ind_CO14O3CHO)	see general notes*
H44019b_a01	TrAa01ScC	CO14O3CHO(aq) → CO14O3CHO	k_exb(01, ind_CO14O3CHO)	see general notes*
H44020f_a01	TrAa01ScC	CO14O3CO ₂ H → CO14O3CO ₂ H(aq)	k_exf(01, ind_CO14O3CO2H)	see general notes*
H44020b_a01	TrAa01ScC	CO14O3CO ₂ H(aq) → CO14O3CO ₂ H	k_exb(01, ind_CO14O3CO2H)	see general notes*
H44021f_a01	TrAa01ScC	CH ₃ COCOCHO → CH ₃ COCOCHO(aq)	k_exf(01, ind_CO23C3CHO)	see general notes*
H44021b_a01	TrAa01ScC	CH ₃ COCOCHO(aq) → CH ₃ COCOCHO	k_exb(01, ind_CO23C3CHO)	see general notes*
H44022f_a01	TrAa01ScC	CO2C3CHO → CO2C3CHO(aq)	k_exf(01, ind_CO2C3CHO)	see general notes*
H44022b_a01	TrAa01ScC	CO2C3CHO(aq) → CO2C3CHO	k_exb(01, ind_CO2C3CHO)	see general notes*
H44023f_a01	TrAa01ScC	CO2C4DIAL → CO2C4DIAL(aq)	k_exf(01, ind_CO2C4DIAL)	see general notes*
H44023b_a01	TrAa01ScC	CO2C4DIAL(aq) → CO2C4DIAL	k_exb(01, ind_CO2C4DIAL)	see general notes*
H44024f_a01	TrAa01ScC	CO2H3CHO → CO2H3CHO(aq)	k_exf(01, ind_CO2H3CHO)	see general notes*
H44024b_a01	TrAa01ScC	CO2H3CHO(aq) → CO2H3CHO	k_exb(01, ind_CO2H3CHO)	see general notes*
H44025f_a01	TrAa01ScC	CO2H3CO ₂ H → CO2H3CO ₂ H(aq)	k_exf(01, ind_CO2H3CO2H)	see general notes*
H44025b_a01	TrAa01ScC	CO2H3CO ₂ H(aq) → CO2H3CO ₂ H	k_exb(01, ind_CO2H3CO2H)	see general notes*
H44026f_a01	TrAa01ScC	CO2H3CO ₃ H → CO2H3CO ₃ H(aq)	k_exf(01, ind_CO2H3CO3H)	see general notes*
H44026b_a01	TrAa01ScC	CO2H3CO ₃ H(aq) → CO2H3CO ₃ H	k_exb(01, ind_CO2H3CO3H)	see general notes*
H44027f_a01	TrAa01ScC	EPXC4DIAL → EPXC4DIAL(aq)	k_exf(01, ind_EPXC4DIAL)	see general notes*
H44027b_a01	TrAa01ScC	EPXC4DIAL(aq) → EPXC4DIAL	k_exb(01, ind_EPXC4DIAL)	see general notes*
H44028f_a01	TrAa01ScC	EPXDLCO ₂ H → EPXDLCO ₂ H(aq)	k_exf(01, ind_EPXDLCO2H)	see general notes*
H44028b_a01	TrAa01ScC	EPXDLCO ₂ H(aq) → EPXDLCO ₂ H	k_exb(01, ind_EPXDLCO2H)	see general notes*
H44029f_a01	TrAa01ScC	EPXDLCO ₃ H → EPXDLCO ₃ H(aq)	k_exf(01, ind_EPXDLCO3H)	see general notes*
H44029b_a01	TrAa01ScC	EPXDLCO ₃ H(aq) → EPXDLCO ₃ H	k_exb(01, ind_EPXDLCO3H)	see general notes*

Table 3: Reversible (Henry's law) equilibria and irreversible ("heterogenous") uptake

#	labels	reaction	rate coefficient	reference
H44030f_a01	TrAa01ScC	$\text{HCOCCH}_3\text{CHOOH}$ $\text{HCOCCH}_3\text{CHOOH(aq)}$	\rightarrow $k_{\text{exf}}(01, \text{ind_HCOCCH}_3\text{CHOOH})$	see general notes*
H44030b_a01	TrAa01ScC	$\text{HCOCCH}_3\text{CHOOH(aq)}$ $\text{HCOCCH}_3\text{CHOOH}$	\rightarrow $k_{\text{exb}}(01, \text{ind_HCOCCH}_3\text{CHOOH})$	see general notes*
H44031f_a01	TrAa01ScC	$\text{HCOCCH}_3\text{CO} \rightarrow \text{HCOCCH}_3\text{CO(aq)}$	$k_{\text{exf}}(01, \text{ind_HCOCCH}_3\text{CO})$	see general notes*
H44031b_a01	TrAa01ScC	$\text{HCOCCH}_3\text{CO(aq)} \rightarrow \text{HCOCCH}_3\text{CO}$	$k_{\text{exb}}(01, \text{ind_HCOCCH}_3\text{CO})$	see general notes*
H44032f_a01	TrAa01ScC	$\text{HMAC} \rightarrow \text{HMAC(aq)}$	$k_{\text{exf}}(01, \text{ind_HMAC})$	see general notes*
H44032b_a01	TrAa01ScC	$\text{HMAC(aq)} \rightarrow \text{HMAC}$	$k_{\text{exb}}(01, \text{ind_HMAC})$	see general notes*
H44033f_a01	TrAa01ScC	$\text{HO12CO3C4} \rightarrow \text{HO12CO3C4(aq)}$	$k_{\text{exf}}(01, \text{ind_HO12CO3C4})$	see general notes*
H44033b_a01	TrAa01ScC	$\text{HO12CO3C4(aq)} \rightarrow \text{HO12CO3C4}$	$k_{\text{exb}}(01, \text{ind_HO12CO3C4})$	see general notes*
H44034f_a01	TrAa01ScC	$\text{HOCOC4DIAL} \rightarrow \text{HOCOC4DIAL(aq)}$	$k_{\text{exf}}(01, \text{ind_HOCOC4DIAL})$	see general notes*
H44034b_a01	TrAa01ScC	$\text{HOCOC4DIAL(aq)} \rightarrow \text{HOCOC4DIAL}$	$k_{\text{exb}}(01, \text{ind_HOCOC4DIAL})$	see general notes*
H44035f_a01	TrAa01ScC	$\text{HVMK} \rightarrow \text{HVMK(aq)}$	$k_{\text{exf}}(01, \text{ind_HVMK})$	see general notes*
H44035b_a01	TrAa01ScC	$\text{HVMK(aq)} \rightarrow \text{HVMK}$	$k_{\text{exb}}(01, \text{ind_HVMK})$	see general notes*
H44036f_a01	TrAa01ScC	$\text{IBUTALOH} \rightarrow \text{IBUTALOH(aq)}$	$k_{\text{exf}}(01, \text{ind_IBUTALOH})$	see general notes*
H44036b_a01	TrAa01ScC	$\text{IBUTALOH(aq)} \rightarrow \text{IBUTALOH}$	$k_{\text{exb}}(01, \text{ind_IBUTALOH})$	see general notes*
H44037f_a01	TrAa01ScC	$\text{IBUTDIAL} \rightarrow \text{IBUTDIAL(aq)}$	$k_{\text{exf}}(01, \text{ind_IBUTDIAL})$	see general notes*
H44037b_a01	TrAa01ScC	$\text{IBUTDIAL(aq)} \rightarrow \text{IBUTDIAL}$	$k_{\text{exb}}(01, \text{ind_IBUTDIAL})$	see general notes*
H44038f_a01	TrAa01ScC	$\text{IBUTOLBOOH} \rightarrow \text{IBUTOLBOOH(aq)}$	$k_{\text{exf}}(01, \text{ind_IBUTOLBOOH})$	see general notes*
H44038b_a01	TrAa01ScC	$\text{IBUTOLBOOH(aq)} \rightarrow \text{IBUTOLBOOH}$	$k_{\text{exb}}(01, \text{ind_IBUTOLBOOH})$	see general notes*
H44039f_a01	TrAa01ScC	$\text{IPRHOCO2H} \rightarrow \text{IPRHOCO2H(aq)}$	$k_{\text{exf}}(01, \text{ind_IPRHOCO2H})$	see general notes*
H44039b_a01	TrAa01ScC	$\text{IPRHOCO2H(aq)} \rightarrow \text{IPRHOCO2H}$	$k_{\text{exb}}(01, \text{ind_IPRHOCO2H})$	see general notes*
H44040f_a01	TrAa01ScC	$\text{IPRHOCO3H} \rightarrow \text{IPRHOCO3H(aq)}$	$k_{\text{exf}}(01, \text{ind_IPRHOCO3H})$	see general notes*
H44040b_a01	TrAa01ScC	$\text{IPRHOCO3H(aq)} \rightarrow \text{IPRHOCO3H}$	$k_{\text{exb}}(01, \text{ind_IPRHOCO3H})$	see general notes*
H44041f_a01	TrAa01ScC	$\text{LBUT1ENOOSH} \rightarrow \text{LBUT1ENOOSH(aq)}$	$k_{\text{exf}}(01, \text{ind_LBUT1ENOOSH})$	see general notes*
H44041b_a01	TrAa01ScC	$\text{LBUT1ENOOSH(aq)} \rightarrow \text{LBUT1ENOOSH}$	$k_{\text{exb}}(01, \text{ind_LBUT1ENOOSH})$	see general notes*
H44042f_a01	TrAa01ScC	$\text{LHMVKABOOH} \rightarrow \text{LHMVKABOOH(aq)}$	$k_{\text{exf}}(01, \text{ind_LHMVKABOOH})$	see general notes*
H44042b_a01	TrAa01ScC	$\text{LHMVKABOOH(aq)} \rightarrow \text{LHMVKABOOH}$	$k_{\text{exb}}(01, \text{ind_LHMVKABOOH})$	see general notes*
H44043f_a01	TrAa01ScC	$\text{LMEKO OH} \rightarrow \text{LMEKO OH(aq)}$	$k_{\text{exf}}(01, \text{ind_LMEKO OH})$	see general notes*
H44043b_a01	TrAa01ScC	$\text{LMEKO OH(aq)} \rightarrow \text{LMEKO OH}$	$k_{\text{exb}}(01, \text{ind_LMEKO OH})$	see general notes*
H44044f_a01	TrAa01ScC	$\text{MACO2H} \rightarrow \text{MACO2H(aq)}$	$k_{\text{exf}}(01, \text{ind_MACO2H})$	see general notes*
H44044b_a01	TrAa01ScC	$\text{MACO2H(aq)} \rightarrow \text{MACO2H}$	$k_{\text{exb}}(01, \text{ind_MACO2H})$	see general notes*
H44045f_a01	TrAa01ScC	$\text{MACO3H} \rightarrow \text{MACO3H(aq)}$	$k_{\text{exf}}(01, \text{ind_MACO3H})$	see general notes*
H44045b_a01	TrAa01ScC	$\text{MACO3H(aq)} \rightarrow \text{MACO3H}$	$k_{\text{exb}}(01, \text{ind_MACO3H})$	see general notes*
H44046f_a01	TrAa01ScC	$\text{MACROH} \rightarrow \text{MACROH(aq)}$	$k_{\text{exf}}(01, \text{ind_MACROH})$	see general notes*

Table 3: Reversible (Henry's law) equilibria and irreversible ("heterogenous") uptake

#	labels	reaction	rate coefficient	reference
H44046b_a01	TrAa01ScC	MACROH(aq) → MACROH	k_exb(01, ind_MACROH)	see general notes*
H44047f_a01	TrAa01ScC	MACROOH → MACROOH(aq)	k_exf(01, ind_MACROOH)	see general notes*
H44047b_a01	TrAa01ScC	MACROOH(aq) → MACROOH	k_exb(01, ind_MACROOH)	see general notes*
H44048f_a01	TrAa01ScC	MALANHYOOH → MALANHYOOH(aq)	k_exf(01, ind_MALANHYOOH)	see general notes*
H44048b_a01	TrAa01ScC	MALANHYOOH(aq) → MALANHYOOH	k_exb(01, ind_MALANHYOOH)	see general notes*
H44049f_a01	TrAa01ScC	MALDALCO2H → MALDALCO2H(aq)	k_exf(01, ind_MALDALCO2H)	see general notes*
H44049b_a01	TrAa01ScC	MALDALCO2H(aq) → MALDALCO2H	k_exb(01, ind_MALDALCO2H)	see general notes*
H44050f_a01	TrAa01ScC	MALDALCO3H → MALDALCO3H(aq)	k_exf(01, ind_MALDALCO3H)	see general notes*
H44050b_a01	TrAa01ScC	MALDALCO3H(aq) → MALDALCO3H	k_exb(01, ind_MALDALCO3H)	see general notes*
H44051f_a01	TrAa01ScC	MALDIAL → MALDIAL(aq)	k_exf(01, ind_MALDIAL)	see general notes*
H44051b_a01	TrAa01ScC	MALDIAL(aq) → MALDIAL	k_exb(01, ind_MALDIAL)	see general notes*
H44052f_a01	TrAa01ScC	MALDIALOOH → MALDIALOOH(aq)	k_exf(01, ind_MALDIALOOH)	see general notes*
H44052b_a01	TrAa01ScC	MALDIALOOH(aq) → MALDIALOOH	k_exb(01, ind_MALDIALOOH)	see general notes*
H44053f_a01	TrAa01ScC	MALNHYOHCO → MALNHYOHCO(aq)	k_exf(01, ind_MALNHYOHCO)	see general notes*
H44053b_a01	TrAa01ScC	MALNHYOHCO(aq) → MALNHYOHCO	k_exb(01, ind_MALNHYOHCO)	see general notes*
H44054f_a01	TrAa01ScC	MECOACEOOH → MECOACEOOH(aq)	k_exf(01, ind_MECOACEOOH)	see general notes*
H44054b_a01	TrAa01ScC	MECOACEOOH(aq) → MECOACEOOH	k_exb(01, ind_MECOACEOOH)	see general notes*
H44055f_a01	TrAa01ScCN	MVKNO3 → MVKNO3(aq)	k_exf(01, ind_MVKNO3)	see general notes*
H44055b_a01	TrAa01ScCN	MVKNO3(aq) → MVKNO3	k_exb(01, ind_MVKNO3)	see general notes*
H44056f_a01	TrAa01ScCN	NBZFUOOH → NBZFUOOH(aq)	k_exf(01, ind_NBZFUOOH)	see general notes*
H44056b_a01	TrAa01ScCN	NBZFUOOH(aq) → NBZFUOOH	k_exb(01, ind_NBZFUOOH)	see general notes*
H44057f_a01	TrAa01ScCN	NC4DCO2H → NC4DCO2H(aq)	k_exf(01, ind_NC4DCO2H)	see general notes*
H44057b_a01	TrAa01ScCN	NC4DCO2H(aq) → NC4DCO2H	k_exb(01, ind_NC4DCO2H)	see general notes*
H45000f_a01	TrAa01ScC	ACCOMECHO → ACCOMECHO(aq)	k_exf(01, ind_ACCOMECHO)	see general notes*
H45000b_a01	TrAa01ScC	ACCOMECHO(aq) → ACCOMECHO	k_exb(01, ind_ACCOMECHO)	see general notes*
H45001f_a01	TrAa01ScC	ACCOMEKO3H → ACCOMEKO3H(aq)	k_exf(01, ind_ACCOMEKO3H)	see general notes*
H45001b_a01	TrAa01ScC	ACCOMEKO3H(aq) → ACCOMEKO3H	k_exb(01, ind_ACCOMEKO3H)	see general notes*
H45002f_a01	TrAa01ScC	C1ODC2O2C4OOH C1ODC2O2C4OOH(aq)	→ k_exf(01, ind_C1ODC2O2C4OOH)	see general notes*
H45002b_a01	TrAa01ScC	C1ODC2O2C4OOH(aq) C1ODC2O2C4OOH	→ k_exb(01, ind_C1ODC2O2C4OOH)	see general notes*
H45003f_a01	TrAa01ScC	C1ODC2OOHC4OD C1ODC2OOHC4OD(aq)	→ k_exf(01, ind_C1ODC2OOHC4OD)	see general notes*
H45003b_a01	TrAa01ScC	C1ODC2OOHC4OD(aq) C1ODC2OOHC4OD	→ k_exb(01, ind_C1ODC2OOHC4OD)	see general notes*

Table 3: Reversible (Henry's law) equilibria and irreversible ("heterogenous") uptake

#	labels	reaction	rate coefficient	reference
H45004f_a01	TrAa01ScC	C1ODC3O2C4OOH C1ODC3O2C4OOH(aq)	→ k_exf(01, ind_C10DC302C400H)	see general notes*
H45004b_a01	TrAa01ScC	C1ODC3O2C4OOH(aq) C1ODC3O2C4OOH	→ k_exb(01, ind_C10DC302C400H)	see general notes*
H45005f_a01	TrAa01ScC	C1OOHC2OOHC4OD C1OOHC2OOHC4OD(aq)	→ k_exf(01, ind_C100HC200HC40D)	see general notes*
H45005b_a01	TrAa01ScC	C1OOHC2OOHC4OD(aq) C1OOHC2OOHC4OD	→ k_exb(01, ind_C100HC200HC40D)	see general notes*
H45006f_a01	TrAa01ScC	C24O3CCO2H → C24O3CCO2H(aq)	k_exf(01, ind_C2403CC02H)	see general notes*
H45006b_a01	TrAa01ScC	C24O3CCO2H(aq) → C24O3CCO2H	k_exb(01, ind_C2403CC02H)	see general notes*
H45007f_a01	TrAa01ScC	C4CO2DBCO3 → C4CO2DBCO3(aq)	k_exf(01, ind_C4CO2DBCO3)	see general notes*
H45007b_a01	TrAa01ScC	C4CO2DBCO3(aq) → C4CO2DBCO3	k_exb(01, ind_C4CO2DBCO3)	see general notes*
H45008f_a01	TrAa01ScCN	C4CO2DBPAN → C4CO2DBPAN(aq)	k_exf(01, ind_C4CO2DBPAN)	see general notes*
H45008b_a01	TrAa01ScCN	C4CO2DBPAN(aq) → C4CO2DBPAN	k_exb(01, ind_C4CO2DBPAN)	see general notes*
H45009f_a01	TrAa01ScC	C4CO2DCO3H → C4CO2DCO3H(aq)	k_exf(01, ind_C4CO2DC03H)	see general notes*
H45009b_a01	TrAa01ScC	C4CO2DCO3H(aq) → C4CO2DCO3H	k_exb(01, ind_C4CO2DC03H)	see general notes*
H45010f_a01	TrAa01ScCN	C4MCONO3OH → C4MCONO3OH(aq)	k_exf(01, ind_C4MCONO3OH)	see general notes*
H45010b_a01	TrAa01ScCN	C4MCONO3OH(aq) → C4MCONO3OH	k_exb(01, ind_C4MCONO3OH)	see general notes*
H45011f_a01	TrAa01ScC	C511OOH → C511OOH(aq)	k_exf(01, ind_C511OOH)	see general notes*
H45011b_a01	TrAa01ScC	C511OOH(aq) → C511OOH	k_exb(01, ind_C511OOH)	see general notes*
H45012f_a01	TrAa01ScC	C512OOH → C512OOH(aq)	k_exf(01, ind_C512OOH)	see general notes*
H45012b_a01	TrAa01ScC	C512OOH(aq) → C512OOH	k_exb(01, ind_C512OOH)	see general notes*
H45013f_a01	TrAa01ScC	C5134CO2OH → C5134CO2OH(aq)	k_exf(01, ind_C5134CO20H)	see general notes*
H45013b_a01	TrAa01ScC	C5134CO2OH(aq) → C5134CO2OH	k_exb(01, ind_C5134CO20H)	see general notes*
H45014f_a01	TrAa01ScC	C513CO → C513CO(aq)	k_exf(01, ind_C513CO)	see general notes*
H45014b_a01	TrAa01ScC	C513CO(aq) → C513CO	k_exb(01, ind_C513CO)	see general notes*
H45015f_a01	TrAa01ScC	C513OOH → C513OOH(aq)	k_exf(01, ind_C51300H)	see general notes*
H45015b_a01	TrAa01ScC	C513OOH(aq) → C513OOH	k_exb(01, ind_C51300H)	see general notes*
H45016f_a01	TrAa01ScCN	C514NO3 → C514NO3(aq)	k_exf(01, ind_C514N03)	see general notes*
H45016b_a01	TrAa01ScCN	C514NO3(aq) → C514NO3	k_exb(01, ind_C514N03)	see general notes*
H45017f_a01	TrAa01ScC	C514OOH → C514OOH(aq)	k_exf(01, ind_C51400H)	see general notes*
H45017b_a01	TrAa01ScC	C514OOH(aq) → C514OOH	k_exb(01, ind_C51400H)	see general notes*
H45018f_a01	TrAa01ScC	C54CO → C54CO(aq)	k_exf(01, ind_C54CO)	see general notes*
H45018b_a01	TrAa01ScC	C54CO(aq) → C54CO	k_exb(01, ind_C54CO)	see general notes*
H45019f_a01	TrAa01ScC	C59OOH → C59OOH(aq)	k_exf(01, ind_C5900H)	see general notes*

Table 3: Reversible (Henry's law) equilibria and irreversible ("heterogenous") uptake

#	labels	reaction	rate coefficient	reference
H45019b_a01	TrAa01ScC	C59OOH(aq) → C59OOH	k_exb(01, ind_C59OOH)	see general notes*
H45020f_a01	TrAa01ScC	C5CO14OH → C5CO14OH(aq)	k_exf(01, ind_C5CO14OH)	see general notes*
H45020b_a01	TrAa01ScC	C5CO14OH(aq) → C5CO14OH	k_exb(01, ind_C5CO14OH)	see general notes*
H45021f_a01	TrAa01ScC	C5CO14OOH → C5CO14OOH(aq)	k_exf(01, ind_C5CO14OOH)	see general notes*
H45021b_a01	TrAa01ScC	C5CO14OOH(aq) → C5CO14OOH	k_exb(01, ind_C5CO14OOH)	see general notes*
H45022f_a01	TrAa01ScC	C5DIALCO → C5DIALCO(aq)	k_exf(01, ind_C5DIALCO)	see general notes*
H45022b_a01	TrAa01ScC	C5DIALCO(aq) → C5DIALCO	k_exb(01, ind_C5DIALCO)	see general notes*
H45023f_a01	TrAa01ScC	C5DIALOOH → C5DIALOOH(aq)	k_exf(01, ind_C5DIALOOH)	see general notes*
H45023b_a01	TrAa01ScC	C5DIALOOH(aq) → C5DIALOOH	k_exb(01, ind_C5DIALOOH)	see general notes*
H45024f_a01	TrAa01ScC	C5DICARB → C5DICARB(aq)	k_exf(01, ind_C5DICARB)	see general notes*
H45024b_a01	TrAa01ScC	C5DICARB(aq) → C5DICARB	k_exb(01, ind_C5DICARB)	see general notes*
H45025f_a01	TrAa01ScC	C5DICAROOH → C5DICAROOH(aq)	k_exf(01, ind_C5DICAROOH)	see general notes*
H45025b_a01	TrAa01ScC	C5DICAROOH(aq) → C5DICAROOH	k_exb(01, ind_C5DICAROOH)	see general notes*
H45026f_a01	TrAa01ScCN	C5PAN9 → C5PAN9(aq)	k_exf(01, ind_C5PAN9)	see general notes*
H45026b_a01	TrAa01ScCN	C5PAN9(aq) → C5PAN9	k_exb(01, ind_C5PAN9)	see general notes*
H45027f_a01	TrAa01ScC	CHOC3COOHOH → CHOC3COOHOH(aq)	k_exf(01, ind_CHOC3COOHOH)	see general notes*
H45027b_a01	TrAa01ScC	CHOC3COOHOH(aq) → CHOC3COOHOH	k_exb(01, ind_CHOC3COOHOH)	see general notes*
H45028f_a01	TrAa01ScCN	CHOC3COPAN → CHOC3COPAN(aq)	k_exf(01, ind_CHOC3COPAN)	see general notes*
H45028b_a01	TrAa01ScCN	CHOC3COPAN(aq) → CHOC3COPAN	k_exb(01, ind_CHOC3COPAN)	see general notes*
H45029f_a01	TrAa01ScC	CO13C4CHO → CO13C4CHO(aq)	k_exf(01, ind_CO13C4CHO)	see general notes*
H45029b_a01	TrAa01ScC	CO13C4CHO(aq) → CO13C4CHO	k_exb(01, ind_CO13C4CHO)	see general notes*
H45030f_a01	TrAa01ScC	CO23C4CHO → CO23C4CHO(aq)	k_exf(01, ind_CO23C4CHO)	see general notes*
H45030b_a01	TrAa01ScC	CO23C4CHO(aq) → CO23C4CHO	k_exb(01, ind_CO23C4CHO)	see general notes*
H45031f_a01	TrAa01ScC	CO23C4CO3H → CO23C4CO3H(aq)	k_exf(01, ind_CO23C4CO3H)	see general notes*
H45031b_a01	TrAa01ScC	CO23C4CO3H(aq) → CO23C4CO3H	k_exb(01, ind_CO23C4CO3H)	see general notes*
H45032f_a01	TrAa01ScCN	DB1NO3 → DB1NO3(aq)	k_exf(01, ind_DB1NO3)	see general notes*
H45032b_a01	TrAa01ScCN	DB1NO3(aq) → DB1NO3	k_exb(01, ind_DB1NO3)	see general notes*
H45033f_a01	TrAa01ScC	DB1OOH → DB1OOH(aq)	k_exf(01, ind_DB1OOH)	see general notes*
H45033b_a01	TrAa01ScC	DB1OOH(aq) → DB1OOH	k_exb(01, ind_DB1OOH)	see general notes*
H45034f_a01	TrAa01ScC	DB2OOH → DB2OOH(aq)	k_exf(01, ind_DB2OOH)	see general notes*
H45034b_a01	TrAa01ScC	DB2OOH(aq) → DB2OOH	k_exb(01, ind_DB2OOH)	see general notes*
H45035f_a01	TrAa01ScC	ISOPAOH → ISOPAOH(aq)	k_exf(01, ind_ISOPAOH)	see general notes*
H45035b_a01	TrAa01ScC	ISOPAOH(aq) → ISOPAOH	k_exb(01, ind_ISOPAOH)	see general notes*
H45036f_a01	TrAa01ScCN	ISOPBNO3 → ISOPBNO3(aq)	k_exf(01, ind_ISOPBNO3)	see general notes*
H45036b_a01	TrAa01ScCN	ISOPBNO3(aq) → ISOPBNO3	k_exb(01, ind_ISOPBNO3)	see general notes*

Table 3: Reversible (Henry's law) equilibria and irreversible ("heterogenous") uptake

#	labels	reaction	rate coefficient	reference
H45037f_a01	TrAa01ScC	ISOPBOH → ISOPBOH(aq)	k_exf(01, ind_ISOPBOH)	see general notes*
H45037b_a01	TrAa01ScC	ISOPBOH(aq) → ISOPBOH	k_exb(01, ind_ISOPBOH)	see general notes*
H45038f_a01	TrAa01ScC	ISOPBOOH → ISOPBOOH(aq)	k_exf(01, ind_ISOPBOOH)	see general notes*
H45038b_a01	TrAa01ScC	ISOPBOOH(aq) → ISOPBOOH	k_exb(01, ind_ISOPBOOH)	see general notes*
H45039f_a01	TrAa01ScCN	ISOPDNO3 → ISOPDNO3(aq)	k_exf(01, ind_ISOPDNO3)	see general notes*
H45039b_a01	TrAa01ScCN	ISOPDNO3(aq) → ISOPDNO3	k_exb(01, ind_ISOPDNO3)	see general notes*
H45040f_a01	TrAa01ScC	ISOPDOH → ISOPDOH(aq)	k_exf(01, ind_ISOPDOH)	see general notes*
H45040b_a01	TrAa01ScC	ISOPDOH(aq) → ISOPDOH	k_exb(01, ind_ISOPDOH)	see general notes*
H45041f_a01	TrAa01ScC	ISOPDOOH → ISOPDOOH(aq)	k_exf(01, ind_ISOPDOOH)	see general notes*
H45041b_a01	TrAa01ScC	ISOPDOOH(aq) → ISOPDOOH	k_exb(01, ind_ISOPDOOH)	see general notes*
H45042f_a01	TrAa01ScC	LC578OOH → LC578OOH(aq)	k_exf(01, ind_LC578OOH)	see general notes*
H45042b_a01	TrAa01ScC	LC578OOH(aq) → LC578OOH	k_exb(01, ind_LC578OOH)	see general notes*
H45043f_a01	TrAa01ScCN	LC5PAN1719 → LC5PAN1719(aq)	k_exf(01, ind_LC5PAN1719)	see general notes*
H45043b_a01	TrAa01ScCN	LC5PAN1719(aq) → LC5PAN1719	k_exb(01, ind_LC5PAN1719)	see general notes*
H45044f_a01	TrAa01ScC	LHC4ACCHO → LHC4ACCHO(aq)	k_exf(01, ind_LHC4ACCHO)	see general notes*
H45044b_a01	TrAa01ScC	LHC4ACCHO(aq) → LHC4ACCHO	k_exb(01, ind_LHC4ACCHO)	see general notes*
H45045f_a01	TrAa01ScC	LHC4ACCO2H → LHC4ACCO2H(aq)	k_exf(01, ind_LHC4ACCO2H)	see general notes*
H45045b_a01	TrAa01ScC	LHC4ACCO2H(aq) → LHC4ACCO2H	k_exb(01, ind_LHC4ACCO2H)	see general notes*
H45046f_a01	TrAa01ScC	LHC4ACCO3H → LHC4ACCO3H(aq)	k_exf(01, ind_LHC4ACCO3H)	see general notes*
H45046b_a01	TrAa01ScC	LHC4ACCO3H(aq) → LHC4ACCO3H	k_exb(01, ind_LHC4ACCO3H)	see general notes*
H45047f_a01	TrAa01ScC	LIEPOX → LIEPOX(aq)	k_exf(01, ind_LIEPOX)	see general notes*
H45047b_a01	TrAa01ScC	LIEPOX(aq) → LIEPOX	k_exb(01, ind_LIEPOX)	see general notes*
H45048f_a01	TrAa01ScCN	LISOPACNO3 → LISOPACNO3(aq)	k_exf(01, ind_LISOPACNO3)	see general notes*
H45048b_a01	TrAa01ScCN	LISOPACNO3(aq) → LISOPACNO3	k_exb(01, ind_LISOPACNO3)	see general notes*
H45049f_a01	TrAa01ScC	LISOPACCOOH → LISOPACCOOH(aq)	k_exf(01, ind_LISOPACCOOH)	see general notes*
H45049b_a01	TrAa01ScC	LISOPACCOOH(aq) → LISOPACCOOH	k_exb(01, ind_LISOPACCOOH)	see general notes*
H45050f_a01	TrAa01ScCN	LMBOABNO3 → LMBOABNO3(aq)	k_exf(01, ind_LMBOABNO3)	see general notes*
H45050b_a01	TrAa01ScCN	LMBOABNO3(aq) → LMBOABNO3	k_exb(01, ind_LMBOABNO3)	see general notes*
H45051f_a01	TrAa01ScC	LMBOABOOH → LMBOABOOH(aq)	k_exf(01, ind_LMBOABOOH)	see general notes*
H45051b_a01	TrAa01ScC	LMBOABOOH(aq) → LMBOABOOH	k_exb(01, ind_LMBOABOOH)	see general notes*
H45052f_a01	TrAa01ScCN	LNMBOABOOH → LNMBOABOOH(aq)	k_exf(01, ind_LNMBOABOOH)	see general notes*
H45052b_a01	TrAa01ScCN	LNMBOABOOH(aq) → LNMBOABOOH	k_exb(01, ind_LNMBOABOOH)	see general notes*
H45053f_a01	TrAa01ScC	MBO → MBO(aq)	k_exf(01, ind_MBO)	see general notes*
H45053b_a01	TrAa01ScC	MBO(aq) → MBO	k_exb(01, ind_MBO)	see general notes*
H45054f_a01	TrAa01ScC	MBOACO → MBOACO(aq)	k_exf(01, ind_MBOACO)	see general notes*

Table 3: Reversible (Henry's law) equilibria and irreversible ("heterogenous") uptake

#	labels	reaction	rate coefficient	reference
H45054b_a01	TrAa01ScC	MBOACO(aq) → MBOACO	k_exb(01, ind_MBOACO)	see general notes*
H45055f_a01	TrAa01ScC	MBOCOCO → MBOCOCO(aq)	k_exf(01, ind_MB0COCO)	see general notes*
H45055b_a01	TrAa01ScC	MBOCOCO(aq) → MBOCOCO	k_exb(01, ind_MB0COCO)	see general notes*
H45056f_a01	TrAa01ScC	MC3ODBCO2H → MC3ODBCO2H(aq)	k_exf(01, ind_MC3ODBCO2H)	see general notes*
H45056b_a01	TrAa01ScC	MC3ODBCO2H(aq) → MC3ODBCO2H	k_exb(01, ind_MC3ODBCO2H)	see general notes*
H45057f_a01	TrAa01ScC	3METHYLFURAN 3METHYLFURAN(aq)	→ k_exf(01, ind_ME3FURAN)	see general notes*
H45057b_a01	TrAa01ScC	3METHYLFURAN(aq) 3METHYLFURAN	→ k_exb(01, ind_ME3FURAN)	see general notes*
H45058f_a01	TrAa01ScC	MMALNHYOOH → MMALNHYOOH(aq)	k_exf(01, ind_MMALNHYOOH)	see general notes*
H45058b_a01	TrAa01ScC	MMALNHYOOH(aq) → MMALNHYOOH	k_exb(01, ind_MMALNHYOOH)	see general notes*
H45059f_a01	TrAa01ScCN	NC4MDCO2HN → NC4MDCO2HN(aq)	k_exf(01, ind_NC4MDCO2H)	see general notes*
H45059b_a01	TrAa01ScCN	NC4MDCO2HN(aq) → NC4MDCO2HN	k_exb(01, ind_NC4MDCO2H)	see general notes*
H45060f_a01	TrAa01ScCN	NC4OHCO3H → NC4OHCO3H(aq)	k_exf(01, ind_NC4OHCO3H)	see general notes*
H45060b_a01	TrAa01ScCN	NC4OHCO3H(aq) → NC4OHCO3H	k_exb(01, ind_NC4OHCO3H)	see general notes*
H45061f_a01	TrAa01ScCN	NC4OHC PAN → NC4OHC PAN(aq)	k_exf(01, ind_NC4OHC PAN)	see general notes*
H45061b_a01	TrAa01ScCN	NC4OHC PAN(aq) → NC4OHC PAN	k_exb(01, ind_NC4OHC PAN)	see general notes*
H45062f_a01	TrAa01ScCN	NISOPOOH → NISOPOOH(aq)	k_exf(01, ind_NISOPOOH)	see general notes*
H45062b_a01	TrAa01ScCN	NISOPOOH(aq) → NISOPOOH	k_exb(01, ind_NISOPOOH)	see general notes*
H45063f_a01	TrAa01ScCN	NMBOBCO → NMBOBCO(aq)	k_exf(01, ind_NMBOBCO)	see general notes*
H45063b_a01	TrAa01ScCN	NMBOBCO(aq) → NMBOBCO	k_exb(01, ind_NMBOBCO)	see general notes*
H45064f_a01	TrAa01ScCN	NTLFUOOH → NTLFUOOH(aq)	k_exf(01, ind_NTLFUOOH)	see general notes*
H45064b_a01	TrAa01ScCN	NTLFUOOH(aq) → NTLFUOOH	k_exb(01, ind_NTLFUOOH)	see general notes*
H45065f_a01	TrAa01ScC	TLFUOOH → TLFUOOH(aq)	k_exf(01, ind_TLFUOOH)	see general notes*
H45065b_a01	TrAa01ScC	TLFUOOH(aq) → TLFUOOH	k_exb(01, ind_TLFUOOH)	see general notes*
H45066f_a01	TrAa01ScC	LZCO3HC23DBCOD LZCO3HC23DBCOD(aq)	→ k_exf(01, ind_LZCO3HC23DBCOD)	see general notes*
H45066b_a01	TrAa01ScC	LZCO3HC23DBCOD(aq) LZCO3HC23DBCOD	→ k_exb(01, ind_LZCO3HC23DBCOD)	see general notes*
H45067f_a01	TrAa01ScC	C4MDIAL → C4MDIAL(aq)	k_exf(01, ind_C4MDIAL)	see general notes*
H45067b_a01	TrAa01ScC	C4MDIAL(aq) → C4MDIAL	k_exb(01, ind_C4MDIAL)	see general notes*
H46000f_a01	TrAa01ScCN	BZBIPERNO3 → BZBIPERNO3(aq)	k_exf(01, ind_BZBIPERNO3)	see general notes*
H46000b_a01	TrAa01ScCN	BZBIPERNO3(aq) → BZBIPERNO3	k_exb(01, ind_BZBIPERNO3)	see general notes*
H46001f_a01	TrAa01ScC	BZBIPEROOH → BZBIPEROOH(aq)	k_exf(01, ind_BZBIPEROOH)	see general notes*
H46001b_a01	TrAa01ScC	BZBIPEROOH(aq) → BZBIPEROOH	k_exb(01, ind_BZBIPEROOH)	see general notes*

Table 3: Reversible (Henry's law) equilibria and irreversible ("heterogenous") uptake

#	labels	reaction	rate coefficient	reference
H46002f_a01	TrAa01ScC	BZEMUCCO → BZEMUCCO(aq)	k_exf(01, ind_BZEMUCCO)	see general notes*
H46002b_a01	TrAa01ScC	BZEMUCCO(aq) → BZEMUCCO	k_exb(01, ind_BZEMUCCO)	see general notes*
H46003f_a01	TrAa01ScC	BZEMUCCO2H → BZEMUCCO2H(aq)	k_exf(01, ind_BZEMUCCO2H)	see general notes*
H46003b_a01	TrAa01ScC	BZEMUCCO2H(aq) → BZEMUCCO2H	k_exb(01, ind_BZEMUCCO2H)	see general notes*
H46004f_a01	TrAa01ScC	BZEMUCCO3H → BZEMUCCO3H(aq)	k_exf(01, ind_BZEMUCCO3H)	see general notes*
H46004b_a01	TrAa01ScC	BZEMUCCO3H(aq) → BZEMUCCO3H	k_exb(01, ind_BZEMUCCO3H)	see general notes*
H46005f_a01	TrAa01ScCN	BZEMUCNO3 → BZEMUCNO3(aq)	k_exf(01, ind_BZEMUCNO3)	see general notes*
H46005b_a01	TrAa01ScCN	BZEMUCNO3(aq) → BZEMUCNO3	k_exb(01, ind_BZEMUCNO3)	see general notes*
H46006f_a01	TrAa01ScC	BZEMUCOOH → BZEMUCOOH(aq)	k_exf(01, ind_BZEMUCOOH)	see general notes*
H46006b_a01	TrAa01ScC	BZEMUCOOH(aq) → BZEMUCOOH	k_exb(01, ind_BZEMUCOOH)	see general notes*
H46007f_a01	TrAa01ScC	BZEPOXMUC → BZEPOXMUC(aq)	k_exf(01, ind_BZEPOXMUC)	see general notes*
H46007b_a01	TrAa01ScC	BZEPOXMUC(aq) → BZEPOXMUC	k_exb(01, ind_BZEPOXMUC)	see general notes*
H46008f_a01	TrAa01ScC	BZOBIPEROH → BZOBIPEROH(aq)	k_exf(01, ind_BZOBIPEROH)	see general notes*
H46008b_a01	TrAa01ScC	BZOBIPEROH(aq) → BZOBIPEROH	k_exb(01, ind_BZOBIPEROH)	see general notes*
H46009f_a01	TrAa01ScCN	C5CO2DBPAN → C5CO2DBPAN(aq)	k_exf(01, ind_C5CO2DBPAN)	see general notes*
H46009b_a01	TrAa01ScCN	C5CO2DBPAN(aq) → C5CO2DBPAN	k_exb(01, ind_C5CO2DBPAN)	see general notes*
H46010f_a01	TrAa01ScC	C5CO2DCO3H → C5CO2DCO3H(aq)	k_exf(01, ind_C5CO2DCO3H)	see general notes*
H46010b_a01	TrAa01ScC	C5CO2DCO3H(aq) → C5CO2DCO3H	k_exb(01, ind_C5CO2DCO3H)	see general notes*
H46011f_a01	TrAa01ScCN	C5CO2OHPAN → C5CO2OHPAN(aq)	k_exf(01, ind_C5CO2OHPAN)	see general notes*
H46011b_a01	TrAa01ScCN	C5CO2OHPAN(aq) → C5CO2OHPAN	k_exb(01, ind_C5CO2OHPAN)	see general notes*
H46012f_a01	TrAa01ScC	C5COOHCO3H → C5COOHCO3H(aq)	k_exf(01, ind_C5COOHCO3H)	see general notes*
H46012b_a01	TrAa01ScC	C5COOHCO3H(aq) → C5COOHCO3H	k_exb(01, ind_C5COOHCO3H)	see general notes*
H46013f_a01	TrAa01ScC	C6125CO → C6125CO(aq)	k_exf(01, ind_C6125CO)	see general notes*
H46013b_a01	TrAa01ScC	C6125CO(aq) → C6125CO	k_exb(01, ind_C6125CO)	see general notes*
H46014f_a01	TrAa01ScC	C614CO → C614CO(aq)	k_exf(01, ind_C614CO)	see general notes*
H46014b_a01	TrAa01ScC	C614CO(aq) → C614CO	k_exb(01, ind_C614CO)	see general notes*
H46015f_a01	TrAa01ScCN	C614NO3 → C614NO3(aq)	k_exf(01, ind_C614NO3)	see general notes*
H46015b_a01	TrAa01ScCN	C614NO3(aq) → C614NO3	k_exb(01, ind_C614NO3)	see general notes*
H46016f_a01	TrAa01ScC	C614OOH → C614OOH(aq)	k_exf(01, ind_C614OOH)	see general notes*
H46016b_a01	TrAa01ScC	C614OOH(aq) → C614OOH	k_exb(01, ind_C614OOH)	see general notes*
H46017f_a01	TrAa01ScC	C615CO2OOH → C615CO2OOH(aq)	k_exf(01, ind_C615CO2OOH)	see general notes*
H46017b_a01	TrAa01ScC	C615CO2OOH(aq) → C615CO2OOH	k_exb(01, ind_C615CO2OOH)	see general notes*
H46018f_a01	TrAa01ScC	C6CO4DB → C6CO4DB(aq)	k_exf(01, ind_C6CO4DB)	see general notes*
H46018b_a01	TrAa01ScC	C6CO4DB(aq) → C6CO4DB	k_exb(01, ind_C6CO4DB)	see general notes*
H46019f_a01	TrAa01ScC	C6H5O → C6H5O(aq)	k_exf(01, ind_C6H5O)	see general notes*

Table 3: Reversible (Henry's law) equilibria and irreversible ("heterogenous") uptake

#	labels	reaction	rate coefficient	reference
H46019b_a01	TrAa01ScC	C6H5O(aq) → C6H5O	k_exb(01, ind_C6H5O)	see general notes*
H46020f_a01	TrAa01ScC	C6H5OOH → C6H5OOH(aq)	k_exf(01, ind_C6H5OOH)	see general notes*
H46020b_a01	TrAa01ScC	C6H5OOH(aq) → C6H5OOH	k_exb(01, ind_C6H5OOH)	see general notes*
H46021f_a01	TrAa01ScC	CATEC1O → CATEC1O(aq)	k_exf(01, ind_CATEC1O)	see general notes*
H46021b_a01	TrAa01ScC	CATEC1O(aq) → CATEC1O	k_exb(01, ind_CATEC1O)	see general notes*
H46022f_a01	TrAa01ScC	CATEC1OOH → CATEC1OOH(aq)	k_exf(01, ind_CATEC1OOH)	see general notes*
H46022b_a01	TrAa01ScC	CATEC1OOH(aq) → CATEC1OOH	k_exb(01, ind_CATEC1OOH)	see general notes*
H46023f_a01	TrAa01ScC	CATECHOL → CATECHOL(aq)	k_exf(01, ind_CATECHOL)	see general notes*
H46023b_a01	TrAa01ScC	CATECHOL(aq) → CATECHOL	k_exb(01, ind_CATECHOL)	see general notes*
H46024f_a01	TrAa01ScC	CO235C5CHO → CO235C5CHO(aq)	k_exf(01, ind_CO235C5CHO)	see general notes*
H46024b_a01	TrAa01ScC	CO235C5CHO(aq) → CO235C5CHO	k_exb(01, ind_CO235C5CHO)	see general notes*
H46025f_a01	TrAa01ScC	CO235C6OOH → CO235C6OOH(aq)	k_exf(01, ind_CO235C6OOH)	see general notes*
H46025b_a01	TrAa01ScC	CO235C6OOH(aq) → CO235C6OOH	k_exb(01, ind_CO235C6OOH)	see general notes*
H46026f_a01	TrAa01ScCN	DNPHEN → DNPHEN(aq)	k_exf(01, ind_DNPHEN)	see general notes*
H46026b_a01	TrAa01ScCN	DNPHEN(aq) → DNPHEN	k_exb(01, ind_DNPHEN)	see general notes*
H46027f_a01	TrAa01ScCN	DNPHENO OH → DNPHENO OH(aq)	k_exf(01, ind_DNPHENO OH)	see general notes*
H46027b_a01	TrAa01ScCN	DNPHENO OH(aq) → DNPHENO OH	k_exb(01, ind_DNPHENO OH)	see general notes*
H46028f_a01	TrAa01ScCN	NBZQOOH → NBZQOOH(aq)	k_exf(01, ind_NBZQOOH)	see general notes*
H46028b_a01	TrAa01ScCN	NBZQOOH(aq) → NBZQOOH	k_exb(01, ind_NBZQOOH)	see general notes*
H46029f_a01	TrAa01ScCN	NCATECHOL → NCATECHOL(aq)	k_exf(01, ind_NCATECHOL)	see general notes*
H46029b_a01	TrAa01ScCN	NCATECHOL(aq) → NCATECHOL	k_exb(01, ind_NCATECHOL)	see general notes*
H46030f_a01	TrAa01ScCN	NCATECOOH → NCATECOOH(aq)	k_exf(01, ind_NCATECOOH)	see general notes*
H46030b_a01	TrAa01ScCN	NCATECOOH(aq) → NCATECOOH	k_exb(01, ind_NCATECOOH)	see general notes*
H46031f_a01	TrAa01ScCN	NDNPHENO OH → NDNPHENO OH(aq)	k_exf(01, ind_NDNPHENO OH)	see general notes*
H46031b_a01	TrAa01ScCN	NDNPHENO OH(aq) → NDNPHENO OH	k_exb(01, ind_NDNPHENO OH)	see general notes*
H46032f_a01	TrAa01ScCN	NNCATECOOH → NNCATECOOH(aq)	k_exf(01, ind_NNCATECOOH)	see general notes*
H46032b_a01	TrAa01ScCN	NNCATECOOH(aq) → NNCATECOOH	k_exb(01, ind_NNCATECOOH)	see general notes*
H46033f_a01	TrAa01ScCN	NPHENO OH → NPHENO OH(aq)	k_exf(01, ind_NPHENO OH)	see general notes*
H46033b_a01	TrAa01ScCN	NPHENO OH(aq) → NPHENO OH	k_exb(01, ind_NPHENO OH)	see general notes*
H46034f_a01	TrAa01ScC	PBZQCO → PBZQCO(aq)	k_exf(01, ind_PBZQCO)	see general notes*
H46034b_a01	TrAa01ScC	PBZQCO(aq) → PBZQCO	k_exb(01, ind_PBZQCO)	see general notes*
H46035f_a01	TrAa01ScC	PBZQOOH → PBZQOOH(aq)	k_exf(01, ind_PBZQOOH)	see general notes*
H46035b_a01	TrAa01ScC	PBZQOOH(aq) → PBZQOOH	k_exb(01, ind_PBZQOOH)	see general notes*
H46036f_a01	TrAa01ScC	PHENOL → PHENOL(aq)	k_exf(01, ind_PHENOL)	see general notes*
H46036b_a01	TrAa01ScC	PHENOL(aq) → PHENOL	k_exb(01, ind_PHENOL)	see general notes*

Table 3: Reversible (Henry's law) equilibria and irreversible ("heterogenous") uptake

#	labels	reaction	rate coefficient	reference
H46037f_a01	TrAa01ScC	PHENOOH → PHENOOH(aq)	k_exf(01, ind_PHENOOH)	see general notes*
H46037b_a01	TrAa01ScC	PHENOOH(aq) → PHENOOH	k_exb(01, ind_PHENOOH)	see general notes*
H460MSf_a01	TrAa01ScCN	HOC6H4NO2 → HOC6H4NO2(aq)	k_exf(01, ind_HOC6H4NO2)	see general notes*
H461MSb_a01	TrAa01ScCN	HOC6H4NO2(aq) → HOC6H4NO2	k_exb(01, ind_HOC6H4NO2)	see general notes*
H47000f_a01	TrAa01ScC	C235C6CO3H → C235C6CO3H(aq)	k_exf(01, ind_C235C6CO3H)	see general notes*
H47000b_a01	TrAa01ScC	C235C6CO3H(aq) → C235C6CO3H	k_exb(01, ind_C235C6CO3H)	see general notes*
H47001f_a01	TrAa01ScCN	C6CO2OHPAN → C6CO2OHPAN(aq)	k_exf(01, ind_C6CO2OHPAN)	see general notes*
H47001b_a01	TrAa01ScCN	C6CO2OHPAN(aq) → C6CO2OHPAN	k_exb(01, ind_C6CO2OHPAN)	see general notes*
H47002f_a01	TrAa01ScC	C6COOHCO3H → C6COOHCO3H(aq)	k_exf(01, ind_C6COOHCO3H)	see general notes*
H47002b_a01	TrAa01ScC	C6COOHCO3H(aq) → C6COOHCO3H	k_exb(01, ind_C6COOHCO3H)	see general notes*
H47003f_a01	TrAa01ScC	C6H5CH2OOH → C6H5CH2OOH(aq)	k_exf(01, ind_C6H5CH2OOH)	see general notes*
H47003b_a01	TrAa01ScC	C6H5CH2OOH(aq) → C6H5CH2OOH	k_exb(01, ind_C6H5CH2OOH)	see general notes*
H47004f_a01	TrAa01ScC	C6H5CO3H → C6H5CO3H(aq)	k_exf(01, ind_C6H5CO3H)	see general notes*
H47004b_a01	TrAa01ScC	C6H5CO3H(aq) → C6H5CO3H	k_exb(01, ind_C6H5CO3H)	see general notes*
H47005f_a01	TrAa01ScC	C716OOH → C716OOH(aq)	k_exf(01, ind_C716OOH)	see general notes*
H47005b_a01	TrAa01ScC	C716OOH(aq) → C716OOH	k_exb(01, ind_C716OOH)	see general notes*
H47006f_a01	TrAa01ScC	C721OOH → C721OOH(aq)	k_exf(01, ind_C721OOH)	see general notes*
H47006b_a01	TrAa01ScC	C721OOH(aq) → C721OOH	k_exb(01, ind_C721OOH)	see general notes*
H47007f_a01	TrAa01ScC	C722OOH → C722OOH(aq)	k_exf(01, ind_C722OOH)	see general notes*
H47007b_a01	TrAa01ScC	C722OOH(aq) → C722OOH	k_exb(01, ind_C722OOH)	see general notes*
H47008f_a01	TrAa01ScC	C7CO4DB → C7CO4DB(aq)	k_exf(01, ind_C7CO4DB)	see general notes*
H47008b_a01	TrAa01ScC	C7CO4DB(aq) → C7CO4DB	k_exb(01, ind_C7CO4DB)	see general notes*
H47009f_a01	TrAa01ScCN	C7PAN3 → C7PAN3(aq)	k_exf(01, ind_C7PAN3)	see general notes*
H47009b_a01	TrAa01ScCN	C7PAN3(aq) → C7PAN3	k_exb(01, ind_C7PAN3)	see general notes*
H47010f_a01	TrAa01ScC	CO235C6CHO → CO235C6CHO(aq)	k_exf(01, ind_CO235C6CHO)	see general notes*
H47010b_a01	TrAa01ScC	CO235C6CHO(aq) → CO235C6CHO	k_exb(01, ind_CO235C6CHO)	see general notes*
H47011f_a01	TrAa01ScC	CRESOL → CRESOL(aq)	k_exf(01, ind_CRESOL)	see general notes*
H47011b_a01	TrAa01ScC	CRESOL(aq) → CRESOL	k_exb(01, ind_CRESOL)	see general notes*
H47012f_a01	TrAa01ScC	CRESOOH → CRESOOH(aq)	k_exf(01, ind_CRESOOH)	see general notes*
H47012b_a01	TrAa01ScC	CRESOOH(aq) → CRESOOH	k_exb(01, ind_CRESOOH)	see general notes*
H47013f_a01	TrAa01ScCN	DNCRES → DNCRES(aq)	k_exf(01, ind_DNCRES)	see general notes*
H47013b_a01	TrAa01ScCN	DNCRES(aq) → DNCRES	k_exb(01, ind_DNCRES)	see general notes*
H47014f_a01	TrAa01ScCN	DNCRESOOH → DNCRESOOH(aq)	k_exf(01, ind_DNCRESOOH)	see general notes*
H47014b_a01	TrAa01ScCN	DNCRESOOH(aq) → DNCRESOOH	k_exb(01, ind_DNCRESOOH)	see general notes*
H47015f_a01	TrAa01ScC	MCATEC1O → MCATEC1O(aq)	k_exf(01, ind_MCATEC1O)	see general notes*

Table 3: Reversible (Henry's law) equilibria and irreversible ("heterogenous") uptake

#	labels	reaction	rate coefficient	reference
H47015b_a01	TrAa01ScC	MCATEC1O(aq) → MCATEC1O	k_exb(01, ind_MCATEC1O)	see general notes*
H47016f_a01	TrAa01ScC	MCATEC1OOH → MCATEC1OOH(aq)	k_exf(01, ind_MCATEC1OOH)	see general notes*
H47016b_a01	TrAa01ScC	MCATEC1OOH(aq) → MCATEC1OOH	k_exb(01, ind_MCATEC1OOH)	see general notes*
H47017f_a01	TrAa01ScC	MCATECHOL → MCATECHOL(aq)	k_exf(01, ind_MCATECHOL)	see general notes*
H47017b_a01	TrAa01ScC	MCATECHOL(aq) → MCATECHOL	k_exb(01, ind_MCATECHOL)	see general notes*
H47018f_a01	TrAa01ScCN	MNCATECH → MNCATECH(aq)	k_exf(01, ind_MNCATECH)	see general notes*
H47018b_a01	TrAa01ScCN	MNCATECH(aq) → MNCATECH	k_exb(01, ind_MNCATECH)	see general notes*
H47019f_a01	TrAa01ScCN	MNCATECOOH → MNCATECOOH(aq)	k_exf(01, ind_MNCATECOOH)	see general notes*
H47019b_a01	TrAa01ScCN	MNCATECOOH(aq) → MNCATECOOH	k_exb(01, ind_MNCATECOOH)	see general notes*
H47020f_a01	TrAa01ScCN	MNNCATCOOH → MNNCATCOOH(aq)	k_exf(01, ind_MNNCATCOOH)	see general notes*
H47020b_a01	TrAa01ScCN	MNNCATCOOH(aq) → MNNCATCOOH	k_exb(01, ind_MNNCATCOOH)	see general notes*
H47021f_a01	TrAa01ScCN	NCRESOOH → NCRESOOH(aq)	k_exf(01, ind_NCRESOOH)	see general notes*
H47021b_a01	TrAa01ScCN	NCRESOOH(aq) → NCRESOOH	k_exb(01, ind_NCRESOOH)	see general notes*
H47022f_a01	TrAa01ScCN	NDNCRESOOH → NDNCRESOOH(aq)	k_exf(01, ind_NDNCRESOOH)	see general notes*
H47022b_a01	TrAa01ScCN	NDNCRESOOH(aq) → NDNCRESOOH	k_exb(01, ind_NDNCRESOOH)	see general notes*
H47023f_a01	TrAa01ScC	OXYL1OOH → OXYL1OOH(aq)	k_exf(01, ind_OXYL1OOH)	see general notes*
H47023b_a01	TrAa01ScC	OXYL1OOH(aq) → OXYL1OOH	k_exb(01, ind_OXYL1OOH)	see general notes*
H47024f_a01	TrAa01ScC	PHCOOH → PHCOOH(aq)	k_exf(01, ind_PHCOOH)	see general notes*
H47024b_a01	TrAa01ScC	PHCOOH(aq) → PHCOOH	k_exb(01, ind_PHCOOH)	see general notes*
H47025f_a01	TrAa01ScC	TLBIPEROOH → TLBIPEROOH(aq)	k_exf(01, ind_TLBIPEROOH)	see general notes*
H47025b_a01	TrAa01ScC	TLBIPEROOH(aq) → TLBIPEROOH	k_exb(01, ind_TLBIPEROOH)	see general notes*
H47026f_a01	TrAa01ScC	TLEMUCCO → TLEMUCCO(aq)	k_exf(01, ind_TLEMUCCO)	see general notes*
H47026b_a01	TrAa01ScC	TLEMUCCO(aq) → TLEMUCCO	k_exb(01, ind_TLEMUCCO)	see general notes*
H47027f_a01	TrAa01ScC	TLEMUCCO2H → TLEMUCCO2H(aq)	k_exf(01, ind_TLEMUCCO2H)	see general notes*
H47027b_a01	TrAa01ScC	TLEMUCCO2H(aq) → TLEMUCCO2H	k_exb(01, ind_TLEMUCCO2H)	see general notes*
H47028f_a01	TrAa01ScC	TLEMUCCO3H → TLEMUCCO3H(aq)	k_exf(01, ind_TLEMUCCO3H)	see general notes*
H47028b_a01	TrAa01ScC	TLEMUCCO3H(aq) → TLEMUCCO3H	k_exb(01, ind_TLEMUCCO3H)	see general notes*
H47029f_a01	TrAa01ScCN	TLEMUCNO3 → TLEMUCNO3(aq)	k_exf(01, ind_TLEMUCNO3)	see general notes*
H47029b_a01	TrAa01ScCN	TLEMUCNO3(aq) → TLEMUCNO3	k_exb(01, ind_TLEMUCNO3)	see general notes*
H47030f_a01	TrAa01ScC	TLEMUCOOH → TLEMUCOOH(aq)	k_exf(01, ind_TLEMUCOOH)	see general notes*
H47030b_a01	TrAa01ScC	TLEMUCOOH(aq) → TLEMUCOOH	k_exb(01, ind_TLEMUCOOH)	see general notes*
H47031f_a01	TrAa01ScC	TLOBIPEROH → TLOBIPEROH(aq)	k_exf(01, ind_TLOBIPEROH)	see general notes*
H47031b_a01	TrAa01ScC	TLOBIPEROH(aq) → TLOBIPEROH	k_exb(01, ind_TLOBIPEROH)	see general notes*
H47032f_a01	TrAa01ScC	TOL1O → TOL1O(aq)	k_exf(01, ind_TOL1O)	see general notes*
H47032b_a01	TrAa01ScC	TOL1O(aq) → TOL1O	k_exb(01, ind_TOL1O)	see general notes*

Table 3: Reversible (Henry's law) equilibria and irreversible ("heterogenous") uptake

#	labels	reaction	rate coefficient	reference
H470MSf_a01	TrAa01ScCN	TOL1OHNO2 → TOL1OHNO2(aq)	k_exf(01, ind_TOL1OHNO2)	see general notes*
H470MSb_a01	TrAa01ScCN	TOL1OHNO2(aq) → TOL1OHNO2	k_exb(01, ind_TOL1OHNO2)	see general notes*
H48000f_a01	TrAa01ScC	C721CHO → C721CHO(aq)	k_exf(01, ind_C721CHO)	see general notes*
H48000b_a01	TrAa01ScC	C721CHO(aq) → C721CHO	k_exb(01, ind_C721CHO)	see general notes*
H48001f_a01	TrAa01ScC	C721CO3H → C721CO3H(aq)	k_exf(01, ind_C721CO3H)	see general notes*
H48001b_a01	TrAa01ScC	C721CO3H(aq) → C721CO3H	k_exb(01, ind_C721CO3H)	see general notes*
H48002f_a01	TrAa01ScCN	C721PAN → C721PAN(aq)	k_exf(01, ind_C721PAN)	see general notes*
H48002b_a01	TrAa01ScCN	C721PAN(aq) → C721PAN	k_exb(01, ind_C721PAN)	see general notes*
H48003f_a01	TrAa01ScCN	C810NO3 → C810NO3(aq)	k_exf(01, ind_C810NO3)	see general notes*
H48003b_a01	TrAa01ScCN	C810NO3(aq) → C810NO3	k_exb(01, ind_C810NO3)	see general notes*
H48004f_a01	TrAa01ScC	C810OOH → C810OOH(aq)	k_exf(01, ind_C810OOH)	see general notes*
H48004b_a01	TrAa01ScC	C810OOH(aq) → C810OOH	k_exb(01, ind_C810OOH)	see general notes*
H48005f_a01	TrAa01ScC	C812OOH → C812OOH(aq)	k_exf(01, ind_C812OOH)	see general notes*
H48005b_a01	TrAa01ScC	C812OOH(aq) → C812OOH	k_exb(01, ind_C812OOH)	see general notes*
H48006f_a01	TrAa01ScC	C813OOH → C813OOH(aq)	k_exf(01, ind_C813OOH)	see general notes*
H48006b_a01	TrAa01ScC	C813OOH(aq) → C813OOH	k_exb(01, ind_C813OOH)	see general notes*
H48007f_a01	TrAa01ScC	C85OOH → C85OOH(aq)	k_exf(01, ind_C85OOH)	see general notes*
H48007b_a01	TrAa01ScC	C85OOH(aq) → C85OOH	k_exb(01, ind_C85OOH)	see general notes*
H48008f_a01	TrAa01ScC	C86OOH → C86OOH(aq)	k_exf(01, ind_C86OOH)	see general notes*
H48008b_a01	TrAa01ScC	C86OOH(aq) → C86OOH	k_exb(01, ind_C86OOH)	see general notes*
H48009f_a01	TrAa01ScCN	C89NO3 → C89NO3(aq)	k_exf(01, ind_C89NO3)	see general notes*
H48009b_a01	TrAa01ScCN	C89NO3(aq) → C89NO3	k_exb(01, ind_C89NO3)	see general notes*
H48010f_a01	TrAa01ScC	C89OOH → C89OOH(aq)	k_exf(01, ind_C89OOH)	see general notes*
H48010b_a01	TrAa01ScC	C89OOH(aq) → C89OOH	k_exb(01, ind_C89OOH)	see general notes*
H48011f_a01	TrAa01ScC	C8BC → C8BC(aq)	k_exf(01, ind_C8BC)	see general notes*
H48011b_a01	TrAa01ScC	C8BC(aq) → C8BC	k_exb(01, ind_C8BC)	see general notes*
H48012f_a01	TrAa01ScC	C8BCCO → C8BCCO(aq)	k_exf(01, ind_C8BCCO)	see general notes*
H48012b_a01	TrAa01ScC	C8BCCO(aq) → C8BCCO	k_exb(01, ind_C8BCCO)	see general notes*
H48013f_a01	TrAa01ScCN	C8BCNO3 → C8BCNO3(aq)	k_exf(01, ind_C8BCNO3)	see general notes*
H48013b_a01	TrAa01ScCN	C8BCNO3(aq) → C8BCNO3	k_exb(01, ind_C8BCNO3)	see general notes*
H48014f_a01	TrAa01ScC	C8BCOOH → C8BCOOH(aq)	k_exf(01, ind_C8BCOOH)	see general notes*
H48014b_a01	TrAa01ScC	C8BCOOH(aq) → C8BCOOH	k_exb(01, ind_C8BCOOH)	see general notes*
H48015f_a01	TrAa01ScC	NORPINIC → NORPINIC(aq)	k_exf(01, ind_NORPINIC)	see general notes*
H48015b_a01	TrAa01ScC	NORPINIC(aq) → NORPINIC	k_exb(01, ind_NORPINIC)	see general notes*
H48016f_a01	TrAa01ScC	STYRENOOH → STYRENOOH(aq)	k_exf(01, ind_STYRENOOH)	see general notes*

Table 3: Reversible (Henry's law) equilibria and irreversible ("heterogenous") uptake

#	labels	reaction	rate coefficient	reference
H48016b_a01	TrAa01ScC	STYRENOOH(aq) → STYRENOOH	k_exb(01, ind_STYRENOOH)	see general notes*
H49000f_a01	TrAa01ScC	C811CO3H → C811CO3H(aq)	k_exf(01, ind_C811CO3H)	see general notes*
H49000b_a01	TrAa01ScC	C811CO3H(aq) → C811CO3H	k_exb(01, ind_C811CO3H)	see general notes*
H49001f_a01	TrAa01ScCN	C811PAN → C811PAN(aq)	k_exf(01, ind_C811PAN)	see general notes*
H49001b_a01	TrAa01ScCN	C811PAN(aq) → C811PAN	k_exb(01, ind_C811PAN)	see general notes*
H49002f_a01	TrAa01ScC	C85CO3H → C85CO3H(aq)	k_exf(01, ind_C85CO3H)	see general notes*
H49002b_a01	TrAa01ScC	C85CO3H(aq) → C85CO3H	k_exb(01, ind_C85CO3H)	see general notes*
H49003f_a01	TrAa01ScC	C89CO2H → C89CO2H(aq)	k_exf(01, ind_C89CO2H)	see general notes*
H49003b_a01	TrAa01ScC	C89CO2H(aq) → C89CO2H	k_exb(01, ind_C89CO2H)	see general notes*
H49004f_a01	TrAa01ScC	C89CO3H → C89CO3H(aq)	k_exf(01, ind_C89CO3H)	see general notes*
H49004b_a01	TrAa01ScC	C89CO3H(aq) → C89CO3H	k_exb(01, ind_C89CO3H)	see general notes*
H49005f_a01	TrAa01ScCN	C89PAN → C89PAN(aq)	k_exf(01, ind_C89PAN)	see general notes*
H49005b_a01	TrAa01ScCN	C89PAN(aq) → C89PAN	k_exb(01, ind_C89PAN)	see general notes*
H49006f_a01	TrAa01ScCN	C96NO3 → C96NO3(aq)	k_exf(01, ind_C96NO3)	see general notes*
H49006b_a01	TrAa01ScCN	C96NO3(aq) → C96NO3	k_exb(01, ind_C96NO3)	see general notes*
H49007f_a01	TrAa01ScC	C96OOH → C96OOH(aq)	k_exf(01, ind_C96OOH)	see general notes*
H49007b_a01	TrAa01ScC	C96OOH(aq) → C96OOH	k_exb(01, ind_C96OOH)	see general notes*
H49008f_a01	TrAa01ScC	C97OOH → C97OOH(aq)	k_exf(01, ind_C97OOH)	see general notes*
H49008b_a01	TrAa01ScC	C97OOH(aq) → C97OOH	k_exb(01, ind_C97OOH)	see general notes*
H49009f_a01	TrAa01ScC	C98OOH → C98OOH(aq)	k_exf(01, ind_C98OOH)	see general notes*
H49009b_a01	TrAa01ScC	C98OOH(aq) → C98OOH	k_exb(01, ind_C98OOH)	see general notes*
H49010f_a01	TrAa01ScCN	C9PAN2 → C9PAN2(aq)	k_exf(01, ind_C9PAN2)	see general notes*
H49010b_a01	TrAa01ScCN	C9PAN2(aq) → C9PAN2	k_exb(01, ind_C9PAN2)	see general notes*
H49011f_a01	TrAa01ScC	NOPINDCO → NOPINDCO(aq)	k_exf(01, ind_NOPINDCO)	see general notes*
H49011b_a01	TrAa01ScC	NOPINDCO(aq) → NOPINDCO	k_exb(01, ind_NOPINDCO)	see general notes*
H49012f_a01	TrAa01ScC	NOPINDOOH → NOPINDOOH(aq)	k_exf(01, ind_NOPINDOOH)	see general notes*
H49012b_a01	TrAa01ScC	NOPINDOOH(aq) → NOPINDOOH	k_exb(01, ind_NOPINDOOH)	see general notes*
H49013f_a01	TrAa01ScC	NOPINONE → NOPINONE(aq)	k_exf(01, ind_NOPINONE)	see general notes*
H49013b_a01	TrAa01ScC	NOPINONE(aq) → NOPINONE	k_exb(01, ind_NOPINONE)	see general notes*
H49014f_a01	TrAa01ScC	NOPINOO → NOPINOO(aq)	k_exf(01, ind_NOPINOO)	see general notes*
H49014b_a01	TrAa01ScC	NOPINOO(aq) → NOPINOO	k_exb(01, ind_NOPINOO)	see general notes*
H49015f_a01	TrAa01ScC	NORPINAL → NORPINAL(aq)	k_exf(01, ind_NORPINAL)	see general notes*
H49015b_a01	TrAa01ScC	NORPINAL(aq) → NORPINAL	k_exb(01, ind_NORPINAL)	see general notes*
H49016f_a01	TrAa01ScC	NORPINENOL → NORPINENOL(aq)	k_exf(01, ind_NORPINENOL)	see general notes*
H49016b_a01	TrAa01ScC	NORPINENOL(aq) → NORPINENOL	k_exb(01, ind_NORPINENOL)	see general notes*

Table 3: Reversible (Henry's law) equilibria and irreversible ("heterogenous") uptake

#	labels	reaction	rate coefficient	reference
H49017f_a01	TrAa01ScC	PINIC → PINIC(aq)	k_exf(01, ind_PINIC)	see general notes*
H49017b_a01	TrAa01ScC	PINIC(aq) → PINIC	k_exb(01, ind_PINIC)	see general notes*
H410000f_a01	TrAa01ScCN	BPINANO3 → BPINANO3(aq)	k_exf(01, ind_BPINANO3)	see general notes*
H410000b_a01	TrAa01ScCN	BPINANO3(aq) → BPINANO3	k_exb(01, ind_BPINANO3)	see general notes*
H410001f_a01	TrAa01ScC	BPINAOOH → BPINAOOH(aq)	k_exf(01, ind_BPINAOOH)	see general notes*
H410001b_a01	TrAa01ScC	BPINAOOH(aq) → BPINAOOH	k_exb(01, ind_BPINAOOH)	see general notes*
H410002f_a01	TrAa01ScCN	C106NO3 → C106NO3(aq)	k_exf(01, ind_C106NO3)	see general notes*
H410002b_a01	TrAa01ScCN	C106NO3(aq) → C106NO3	k_exb(01, ind_C106NO3)	see general notes*
H410003f_a01	TrAa01ScC	C106OOH → C106OOH(aq)	k_exf(01, ind_C106OOH)	see general notes*
H410003b_a01	TrAa01ScC	C106OOH(aq) → C106OOH	k_exb(01, ind_C106OOH)	see general notes*
H410004f_a01	TrAa01ScC	C109CO → C109CO(aq)	k_exf(01, ind_C109CO)	see general notes*
H410004b_a01	TrAa01ScC	C109CO(aq) → C109CO	k_exb(01, ind_C109CO)	see general notes*
H410005f_a01	TrAa01ScC	C109OOH → C109OOH(aq)	k_exf(01, ind_C109OOH)	see general notes*
H410005b_a01	TrAa01ScC	C109OOH(aq) → C109OOH	k_exb(01, ind_C109OOH)	see general notes*
H410006f_a01	TrAa01ScCN	C10PAN2 → C10PAN2(aq)	k_exf(01, ind_C10PAN2)	see general notes*
H410006b_a01	TrAa01ScCN	C10PAN2(aq) → C10PAN2	k_exb(01, ind_C10PAN2)	see general notes*
H410007f_a01	TrAa01ScCN	LAPINABNO3 → LAPINABNO3(aq)	k_exf(01, ind_LAPINABNO3)	see general notes*
H410007b_a01	TrAa01ScCN	LAPINABNO3(aq) → LAPINABNO3	k_exb(01, ind_LAPINABNO3)	see general notes*
H410008f_a01	TrAa01ScC	LAPINABOOH → LAPINABOOH(aq)	k_exf(01, ind_LAPINABOOH)	see general notes*
H410008b_a01	TrAa01ScC	LAPINABOOH(aq) → LAPINABOOH	k_exb(01, ind_LAPINABOOH)	see general notes*
H410009f_a01	TrAa01ScCN	LNAPINABOOH → LNAPINABOOH(aq)	k_exf(01, ind_LNAPINABOOH)	see general notes*
H410009b_a01	TrAa01ScCN	LNAPINABOOH(aq) → LNAPINABOOH	k_exb(01, ind_LNAPINABOOH)	see general notes*
H410010f_a01	TrAa01ScCN	LNBPINABOOH → LNBPINABOOH(aq)	k_exf(01, ind_LNBPINABOOH)	see general notes*
H410010b_a01	TrAa01ScCN	LNBPINABOOH(aq) → LNBPINABOOH	k_exb(01, ind_LNBPINABOOH)	see general notes*
H410011f_a01	TrAa01ScC	MENTHEN6ONE → MENTHEN6ONE(aq)	k_exf(01, ind_MENTHEN6ONE)	see general notes*
H410011b_a01	TrAa01ScC	MENTHEN6ONE(aq) → MENTHEN6ONE	k_exb(01, ind_MENTHEN6ONE)	see general notes*
H410012f_a01	TrAa01ScC	2OHMENTHEN6ONE 2OHMENTHEN6ONE(aq)	→ k_exf(01, ind_OH2MENTHEN6ONE)	see general notes*
H410012b_a01	TrAa01ScC	2OHMENTHEN6ONE(aq) 2OHMENTHEN6ONE	→ k_exb(01, ind_OH2MENTHEN6ONE)	see general notes*
H410013f_a01	TrAa01ScC	PERPINONIC → PERPINONIC(aq)	k_exf(01, ind_PERPINONIC)	see general notes*
H410013b_a01	TrAa01ScC	PERPINONIC(aq) → PERPINONIC	k_exb(01, ind_PERPINONIC)	see general notes*
H410014f_a01	TrAa01ScC	PINAL → PINAL(aq)	k_exf(01, ind_PINAL)	see general notes*
H410014b_a01	TrAa01ScC	PINAL(aq) → PINAL	k_exb(01, ind_PINAL)	see general notes*
H410015f_a01	TrAa01ScCN	PINALNO3 → PINALNO3(aq)	k_exf(01, ind_PINALNO3)	see general notes*

Table 3: Reversible (Henry's law) equilibria and irreversible ("heterogenous") uptake

#	labels	reaction	rate coefficient	reference
H410015b_a01	TrAa01ScCN	PINALNO3(aq) → PINALNO3	k_exb(01, ind_PINALN03)	see general notes*
H410016f_a01	TrAa01ScC	PINALOOH → PINALOOH(aq)	k_exf(01, ind_PINALOOH)	see general notes*
H410016b_a01	TrAa01ScC	PINALOOH(aq) → PINALOOH	k_exb(01, ind_PINALOOH)	see general notes*
H410017f_a01	TrAa01ScC	PINEOL → PINEOL(aq)	k_exf(01, ind_PINENOL)	see general notes*
H410017b_a01	TrAa01ScC	PINEOL(aq) → PINEOL	k_exb(01, ind_PINENOL)	see general notes*
H410018f_a01	TrAa01ScC	PINONIC → PINONIC(aq)	k_exf(01, ind_PINONIC)	see general notes*
H410018b_a01	TrAa01ScC	PINONIC(aq) → PINONIC	k_exb(01, ind_PINONIC)	see general notes*
H410019f_a01	TrAa01ScCN	RO6R1NO3 → RO6R1NO3(aq)	k_exf(01, ind_RO6R1N03)	see general notes*
H410019b_a01	TrAa01ScCN	RO6R1NO3(aq) → RO6R1NO3	k_exb(01, ind_RO6R1N03)	see general notes*
H410020f_a01	TrAa01ScCN	ROO6R1NO3 → ROO6R1NO3(aq)	k_exf(01, ind_ROO6R1N03)	see general notes*
H410020b_a01	TrAa01ScCN	ROO6R1NO3(aq) → ROO6R1NO3	k_exb(01, ind_ROO6R1N03)	see general notes*
H60000f_a01	TrAa01MblScCl	Cl ₂ → Cl ₂ (aq)	k_exf(01, ind_Cl2)	see general notes*
H60000b_a01	TrAa01MblScCl	Cl ₂ (aq) → Cl ₂	k_exb(01, ind_Cl2)	see general notes*
H62000f_a01	TrAa01MblScScmCl	HCl → HCl(aq)	k_exf(01, ind_HCl)	see general notes*
H62000b_a01	TrAa01MblScScmCl	HCl(aq) → HCl	k_exb(01, ind_HCl)	see general notes*
H62001f_a01	TrAa01MblScCl	HOCl → HOCl(aq)	k_exf(01, ind_HOCl)	see general notes*
H62001b_a01	TrAa01MblScCl	HOCl(aq) → HOCl	k_exb(01, ind_HOCl)	see general notes*
H63001_a01	TrAa01MblClN	ClNO ₃ → HOCl(aq) + HNO ₃ (aq)	k_exf_ClNO3(01) * C(ind_H20_a01)	see general notes*
H63002_a01	TrAa01MblClN	ClNO ₃ + Cl ⁻ (aq) → Cl ₂ (aq) + NO ₃ ⁻ (aq)	k_exf_ClNO3(01) * 5.E2	see general notes*
H631MS_a01	TrAa01MblClN	ClNO ₂ → ClNO ₂ (aq)	k_exf(01, ind_ClNO2)	see general notes*
H632MS_a01	TrAa01MblClN	ClNO ₂ (aq) → ClNO ₂	k_exb(01, ind_ClNO2)	see general notes*
H633MS_a01	TrAa01MblClN	ClNO → ClNO(aq)	k_exf(01, ind_ClNO)	see general notes*
H634MS_a01	TrAa01MblClN	ClNO(aq) → ClNO	k_exb(01, ind_ClNO)	see general notes*
H70000f_a01	TrAa01MblScBr	Br ₂ → Br ₂ (aq)	k_exf(01, ind_Br2)	see general notes*
H70000b_a01	TrAa01MblScBr	Br ₂ (aq) → Br ₂	k_exb(01, ind_Br2)	see general notes*
H72000f_a01	TrAa01MblScScmBr	HBr → HBr(aq)	k_exf(01, ind_HBr)	see general notes*
H72000b_a01	TrAa01MblScScmBr	HBr(aq) → HBr	k_exb(01, ind_HBr)	see general notes*
H72001f_a01	TrAa01MblScBr	HOBr → HOBr(aq)	k_exf(01, ind_HOBr)	see general notes*
H72001b_a01	TrAa01MblScBr	HOBr(aq) → HOBr	k_exb(01, ind_HOBr)	see general notes*
H73000_a01	TrAa01MblBrN	N ₂ O ₅ + Br ⁻ (aq) → BrNO ₂ + NO ₃ ⁻ (aq)	k_exf_N205(01) * 3.E5	Behnke et al. (1994), Behnke et al. (1997)
H73001_a01	TrAa01MblBrN	BrNO ₃ → HOBr(aq) + HNO ₃ (aq)	k_exf_BrNO3(01) * C(ind_H20_a01)	see general notes*
H73002_a01	TrAa01MblBrN	BrNO ₃ + Br ⁻ (aq) → Br ₂ (aq) + NO ₃ ⁻ (aq)	k_exf_BrNO3(01) * 3.E5	see general notes*
H76000f_a01	TrAa01MblScBrCl	BrCl → BrCl(aq)	k_exf(01, ind_BrCl)	see general notes*
H76000b_a01	TrAa01MblScBrCl	BrCl(aq) → BrCl	k_exb(01, ind_BrCl)	see general notes*

Table 3: Reversible (Henry's law) equilibria and irreversible ("heterogenous") uptake

#	labels	reaction	rate coefficient	reference
H76001_a01	TrAa01MblBrClN	$\text{ClNO}_3 + \text{Br}^-(\text{aq}) \rightarrow \text{BrCl}(\text{aq}) + \text{NO}_3^-(\text{aq})$	$k_{\text{exf}}_{\text{ClNO}_3(01)} * 3.E5$	see general notes*
H76002_a01	TrAa01MblBrClN	$\text{BrNO}_3 + \text{Cl}^-(\text{aq}) \rightarrow \text{BrCl}(\text{aq}) + \text{NO}_3^-(\text{aq})$	$k_{\text{exf}}_{\text{BrNO}_3(01)} * 5.E2$	see general notes*
H91000f_a01	TrAa01MblScScmS	$\text{SO}_2 \rightarrow \text{SO}_2(\text{aq})$	$k_{\text{exf}}(01, \text{ind_SO}_2)$	see general notes*
H91000b_a01	TrAa01MblScScmS	$\text{SO}_2(\text{aq}) \rightarrow \text{SO}_2$	$k_{\text{exb}}(01, \text{ind_SO}_2)$	see general notes*
H92000_a01	TrAa01MblScScmS	$\text{H}_2\text{SO}_4 \rightarrow \text{H}_2\text{SO}_4(\text{aq})$	$xnom7sulf*k_{\text{exf}}(01, \text{ind_H}_2\text{SO}_4)$	see general notes*
H94000f_a01	TrAa01CS	$\text{DMSO} \rightarrow \text{DMSO}(\text{aq})$	$k_{\text{exf}}(01, \text{ind_DMSO})$	see general notes*
H94000b_a01	TrAa01CS	$\text{DMSO}(\text{aq}) \rightarrow \text{DMSO}$	$k_{\text{exb}}(01, \text{ind_DMSO})$	see general notes*
H94001_a01	TrAa01MblS	$\text{CH}_3\text{SO}_3\text{H} \rightarrow \text{CH}_3\text{SO}_3^-(\text{aq}) + \text{H}^+(\text{aq})$	$k_{\text{exf}}(01, \text{ind_CH}_3\text{SO}_3\text{H})$	see general notes*
H94002f_a01	TrAa01CS	$\text{DMS} \rightarrow \text{DMS}(\text{aq})$	$k_{\text{exf}}(01, \text{ind_DMS})$	see general notes*
H94002b_a01	TrAa01CS	$\text{DMS}(\text{aq}) \rightarrow \text{DMS}$	$k_{\text{exb}}(01, \text{ind_DMS})$	see general notes*

General notes

The forward (k_{exf}) and backward (k_{exb}) rate coefficients are calculated in subroutine `mecca_aero_calc_k_ex` in the file `messy_mecca_aero.f90` using accommodation coefficients and Henry's law constants from chemprop (see `chemprop.pdf`).

For uptake of X (X = N_2O_5 , ClNO_3 , or BrNO_3) and

subsequent reaction with H_2O , Cl^- , and Br^- in H3201, H6300, H6301, H6302, H7300, H7301, H7302, H7601, and H7602, we define:

$$k_{\text{exf}}(X) = \frac{k_{\text{mt}}(X) \times \text{LWC}}{[\text{H}_2\text{O}] + 5 \times 10^2 [\text{Cl}^-] + 3 \times 10^5 [\text{Br}^-]}$$

Here, k_{mt} = mass transfer coefficient, and LWC = liquid water content of the aerosol. The total uptake rate of X is only determined by k_{mt} . The factors only affect

the branching between hydrolysis and the halide reactions. The factor 5×10^2 was chosen such that the chloride reaction dominates over hydrolysis at about $[\text{Cl}^-] > 0.1 \text{ M}$ (see Fig. 3 in Behnke et al. (1997)), i.e. when the ratio $[\text{H}_2\text{O}]/[\text{Cl}^-]$ is less than 5×10^2 . The ratio $5 \times 10^2/3 \times 10^5$ was chosen such that the reactions with chloride and bromide are roughly equal for sea water composition (Behnke et al., 1994). These ratios were measured for uptake of N_2O_5 . Here, they are also used for ClNO_3 and BrNO_3 .

Table 4: Heterogeneous reactions

#	labels	reaction	rate coefficient	reference
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General notes

Heterogeneous reaction rates are calculated with an external module (e.g., MECCA_KHET) and then supplied to the MECCA chemistry (see www.messy-interface.org for details)

Table 5: Acid-base and other equilibria

#	labels	reaction	$K_0[M^{m-n}]$	$-\Delta H/R[K]$	reference
EQ2100_a01	TrAa01Sc	$\text{HO}_2 \rightleftharpoons \text{O}_2^- + \text{H}^+$	1.6E-5		Weinstein-Lloyd and Schwartz (1991)
EQ2101_a01	TrAa01MblScScm	$\text{H}_2\text{O} \rightleftharpoons \text{H}^+ + \text{OH}^-$	1.0E-16	-6716	Chameides (1984)
EQ2102_a01	TrAa01Sc	$\text{HO}_3 \rightleftharpoons \text{O}_3^- + \text{H}^+$	4.4E-9		Staehelin et al. (1984)
EQ3200_a01	TrAa01MblScScmN	$\text{NH}_4^+ \rightleftharpoons \text{H}^+ + \text{NH}_3$	5.88E-10	-2391	Chameides (1984)
EQ3201_a01	TrAa01ScN	$\text{HONO} \rightleftharpoons \text{H}^+ + \text{NO}_2^-$	5.1E-4	-1260	Schwartz and White (1981)
EQ3202_a01	TrAa01MblScScmN	$\text{HNO}_3 \rightleftharpoons \text{H}^+ + \text{NO}_3^-$	15	8700	Davis and de Bruin (1964)
EQ3203_a01	TrAa01ScN	$\text{HNO}_4 \rightleftharpoons \text{NO}_4^- + \text{H}^+$	1.E-5		Warneck (1999)
EQ4100_a01	TrAa01MblScScm	$\text{CO}_2 \rightleftharpoons \text{H}^+ + \text{HCO}_3^-$	4.3E-7	-913	Chameides (1984)*
EQ4101_a01	TrAa01ScScm	$\text{HCOOH} \rightleftharpoons \text{H}^+ + \text{HCOO}^-$	1.8E-4		Weast (1980)
EQ4150_a01	TrAa01Sc	$\text{HCHO} \rightleftharpoons \text{HOCH}_2\text{OH}$	4.11E-3	-3769	see note*
EQ4151_a01	TrAa01Sc	$\text{HCO}_3 \rightleftharpoons \text{HCOHOHO}_2$	1.08E1	-2936	see note*
EQ4201_a01	TrAa01ScC	$\text{CH}_3\text{C(O)OOH} \rightleftharpoons \text{CH}_3\text{COOO}^- + \text{H}^+$	6.3E-9		Schuchmann and von Sonntag (1988)
EQ4202_a01	TrAa01ScC	$\text{HOCH}_2\text{CO}_3\text{H} \rightleftharpoons \text{CH}_2\text{OHCO}_2\text{O}^- + \text{H}^+$	6.3E-9		Schuchmann and von Sonntag (1988)*
EQ4203_a01	TrAa01ScC	$\text{HOOCCOOH} \rightleftharpoons \text{H}^+ + \text{HOOCOO}^-$	5.6E-2		Martell (1977)
EQ4204_a01	TrAa01ScC	$\text{HOOCOO}^- \rightleftharpoons \text{H}^+ + \text{C}_2\text{O}_4^{2-}$	5.4E-5		Martell (1977)
EQ4205_a01	TrAa01ScC	$\text{HOOCH}_2\text{CO}_2\text{H} \rightleftharpoons \text{H}^+ + \text{CH}_2\text{OOHCO}_2^-$	1.754E-5		Fisher and Barnes (1972)*
EQ4206_a01	TrAa01ScC	$\text{CH}_2\text{OOCOOH} \rightleftharpoons \text{H}^+ + \text{CH}_2\text{OOCO}_2^-$	1.754E-5		Fisher and Barnes (1972)*
EQ4207_a01	TrAa01ScC	$\text{CHOOHOOCOOH} \rightleftharpoons \text{H}^+ + \text{CHOOHOOCO}_2^-$	1.754E-5		Fisher and Barnes (1972)*
EQ4208_a01	TrAa01ScC	$\text{HOCH}_2\text{CO}_2\text{H} \rightleftharpoons \text{H}^+ + \text{CH}_2\text{OHCO}_2^-$	1.5E-4		Rumble (2020)
EQ4209_a01	TrAa01ScC	$\text{CHOHOOCOOH} \rightleftharpoons \text{H}^+ + \text{CHOHOOCO}_2^-$	1.5E-4		Rumble (2020)*
EQ4210_a01	TrAa01ScC	$\text{CHOCOOH} \rightleftharpoons \text{H}^+ + \text{CHOCOO}^-$	1.754E-5		Fisher and Barnes (1972)
EQ4211_a01	TrAa01ScC	$\text{COOHCO}_3 \rightleftharpoons \text{H}^+ + \text{CO}_2^- \text{CO}_3$	1.754E-5		Fisher and Barnes (1972)
EQ4250_a01	TrAa01ScC	$\text{CH}_3\text{CHO} \rightleftharpoons \text{CH}_3\text{CHOHOH}$	1.22		Tur'yan (2000)
EQ4251_a01	TrAa01ScC	$\text{CHOHOOCOOH} \rightleftharpoons \text{H}^+ + \text{CHOHOOCO}_2^-$	1.57E1		see note*
EQ4252_a01	TrAa01ScC	$\text{CH}_2\text{OHCHO} \rightleftharpoons \text{CH}_2\text{OHCHOHOH}$	1.56E1		Doussin and Monod (2013)
EQ4253_a01	TrAa01ScC	$\text{GLYOX} \rightleftharpoons \text{CHOCHOHOH}$	3.5E2		Ervens and Volkamer (2010)
EQ4254_a01	TrAa01ScC	$\text{CHOCHOHOH} \rightleftharpoons \text{CHOHOCHHOHOH}$	2.0E2		Ervens and Volkamer (2010)
EQ4255_a01	TrAa01ScC	$\text{CHOCOOH} \rightleftharpoons \text{CHOOHOHCOOH}$	1.1E3		Doussin and Monod (2013)
EQ4256_a01	TrAa01ScC	$\text{CHOCOO}^- \rightleftharpoons \text{CHOHOHCO}_2^-$	6.6E1		Doussin and Monod (2013)
EQ4257_a01	TrAa01ScC	$\text{CO}_2^- \text{CO}_3 \rightleftharpoons \text{CO}_2^- \text{COHOHO}_2$	6.6E1		see note*
EQ4258_a01	TrAa01ScC	$\text{CH}_2\text{OOHCHO} \rightleftharpoons \text{HOOCH}_2\text{CHOHOH}$	1.56E1		see note*
EQ4300_a01	TrAa01ScScmC	$\text{CH}_3\text{COCOOH} \rightleftharpoons \text{H}^+ + \text{CH}_3\text{COCO}_2^-$	4.1E-3		Rumble (2020)
EQ4350_a01	TrAa01ScC	$\text{CH}_3\text{C(O)CHO} \rightleftharpoons \text{CH}_3\text{COCHOHOH}$	1.98E3		Doussin and Monod (2013)

Table 5: Acid-base and other equilibria

#	labels	reaction	$K_0[M^{m-n}]$	$-\Delta H/R[K]$	reference
EQ6000_a01	TrAa01Cl	$\text{Cl}_2^- \rightleftharpoons \text{Cl} + \text{Cl}^-$	7.3E-6		Yu (2004)
EQ6200_a01	TrAa01MblScScmCl	$\text{HCl} \rightleftharpoons \text{H}^+ + \text{Cl}^-$	1.7E6	6896	Marsh and McElroy (1985)
EQ6201_a01	TrAa01ScCl	$\text{HOCl} \rightleftharpoons \text{H}^+ + \text{ClO}^-$	3.2E-8		Lax (1969)
EQ7000_a01	TrAa01Br	$\text{Br}_2^- \rightleftharpoons \text{Br} + \text{Br}^-$	2.54E-6	-2256	Liu et al. (2002)
EQ7200_a01	TrAa01MblScScmBr	$\text{HBr} \rightleftharpoons \text{H}^+ + \text{Br}^-$	1.0E9		Lax (1969)
EQ7201_a01	TrAa01ScBr	$\text{HOBr} \rightleftharpoons \text{H}^+ + \text{BrO}^-$	2.3E-9	-3091	Kelley and Tartar (1956)*
EQ7600_a01	TrAa01MblBrCl	$\text{BrCl} + \text{Cl}^- \rightleftharpoons \text{BrCl}_2^-$	3.8	1191	Wang et al. (1994)
EQ7601_a01	TrAa01MblBrCl	$\text{BrCl} + \text{Br}^- \rightleftharpoons \text{Br}_2\text{Cl}^-$	1.8E4	7457	Wang et al. (1994)
EQ7602_a01	TrAa01MblBrCl	$\text{Br}_2 + \text{Cl}^- \rightleftharpoons \text{Br}_2\text{Cl}^-$	1.3	0	Wang et al. (1994)
EQ7603_a01	TrAa01MblBrCl	$\text{Br}^- + \text{Cl}_2 \rightleftharpoons \text{BrCl}_2^-$	4.2E6	14072	Wang et al. (1994)
EQ9200_a01	TrAa01MblScScmS	$\text{SO}_2 \rightleftharpoons \text{H}^+ + \text{HSO}_3^-$	1.7E-2	2090	Chameides (1984)
EQ9201_a01	TrAa01MblScScmS	$\text{HSO}_3^- \rightleftharpoons \text{H}^+ + \text{SO}_3^{2-}$	6.0E-8	1120	Chameides (1984)
EQ9202_a01	TrAa01MblScScmS	$\text{HSO}_4^- \rightleftharpoons \text{H}^+ + \text{SO}_4^{2-}$	1.2E-2	2720	Seinfeld and Pandis (1998)
EQ9203_a01	TrAa01MblScScmS	$\text{H}_2\text{SO}_4 \rightleftharpoons \text{H}^+ + \text{HSO}_4^-$	1.0E3		Seinfeld and Pandis (1998)
EQ11200_a01	TrAa01Fe	$\text{Fe}^{3+} \rightleftharpoons \text{FeOH}^{2+} + \text{H}^+$	2.34E-3		de Laat and Le (2006)*
EQ11201_a01	TrAa01Fe	$\text{FeOH}^{2+} \rightleftharpoons \text{Fe(OH)}_2^+ + \text{H}^+$	2E-4		de Laat and Le (2006)*
EQ11202_a01	TrAa01Fe	$\text{Fe}^{3+} + \text{H}_2\text{O}_2 \rightleftharpoons \text{FeHO}_2^{2+} + \text{H}^+$	3.1E-3		de Laat and Le (2006)
EQ11203_a01	TrAa01Fe	$\text{FeOH}^{2+} + \text{H}_2\text{O}_2 \rightleftharpoons \text{Fe(OH)(HO}_2)^+ + \text{H}^+$	2E-4		de Laat and Le (2006)
EQ11600_a01	TrAa01ClFe	$\text{Fe}^{3+} + \text{Cl}^- \rightleftharpoons \text{FeCl}^{2+}$	6.61		de Laat and Le (2006)*
EQ11601_a01	TrAa01ClFe	$\text{FeCl}^{2+} + \text{Cl}^- \rightleftharpoons \text{FeCl}_2^+$	1.6		de Laat and Le (2006)*
EQ11800_a01	TrAa01FeS	$\text{Fe}^{3+} + \text{SO}_4^{2-} \rightleftharpoons \text{FeSO}_4^+$	120		Brand and van Eldik (1995)*
EQ11801_a01	TrAa01FeS	$\text{FeOH}^{2+} + \text{HSO}_3^- \rightleftharpoons \text{FeSO}_3^+$	8.25E2		Warneck (2018)*
EQ11802_a01	TrAa01FeS	$\text{Fe}^{2+} + \text{SO}_3^- \rightleftharpoons \text{FeSO}_3^+$	1.6E7		Warneck (2018)

Specific notes

EQ4100_a01: For $pK_a(\text{CO}_2)$, see also Dickson and Millero (1987).

EQ4150_a01: Hydration from Winkelmann et al. (2000) and dehydration from Winkelmann et al. (2002). Bell and Evans (1966) found that acid catalysis is negligible.

EQ4151_a01: Assumed to be the same as for HCHO.

EQ4202_a01: Same as for CH3CO3H.

EQ4205_a01: Same as for CH3CO2H.

EQ4206_a01: Same as for CH3CO2H.

EQ4207_a01: Same as for CH3CO2H.

EQ4209_a01: Same as HOCH2CO2H.

EQ4251_a01: Calculated as $K_{\text{eq}} * k(\text{dehydration})$ where dehydration is assumed to be the same as for acetaldehyde.

EQ4257_a01: Assumed to be equal to CHOCO2m.

EQ4258_a01: Same as for HOCH₂CHO.

EQ7201_a01: For $pK_a(\text{HOBr})$, see also Keller-Rudek et al. (1992).

EQ11200_a01: See also K values listed in Tab. 2.5 of Brand and van Eldik (1995).

EQ11201_a01: Equilibrium calculated from K_1 and K_2 in Tab. 1 of de Laat and Le (2006). k for back reaction assumed. See also K values listed in Tab. 2.5 of Brand and van Eldik (1995).

EQ11600_a01: See also K values listed in Tab. 2.5 of Brand and van Eldik (1995).

EQ11601_a01: Equilibrium calculated from K_{29} and K_{30} in Tab. 2 of de Laat and Le (2006). k for forward reaction assumed. See also K values listed in Tab. 2.5 of Brand and van Eldik (1995).

EQ11800_a01: Equilibrium at $I = 1 \text{ M}$. k for back reaction assumed.

EQ11801_a01: Rate of equilibration assumed.

Table 6: Aqueous phase reactions

#	labels	reaction	$k_0 [M^{1-n}s^{-1}]$	$-E_a/R[K]$	reference
A10000_a01	TrAa01Sc	$O_3 + O_2^- \rightarrow O_3^- + O_2$	1.50E9		Staehelin et al. (1984)
A21000_a01	TrAa01Sc	$OH + O_2^- \rightarrow OH^-$	1.0E10		Sehested et al. (1968)
A21001_a01	TrAa01Sc	$OH + OH \rightarrow H_2O_2$	5.5E9		Buxton et al. (1988)
A21002_a01	TrAa01Sc	$HO_2 + O_2^- \rightarrow H_2O_2 + OH^-$	1.0E8	-900	Christensen and Sehested (1988)
A21003_a01	TrAa01Sc	$HO_2 + OH \rightarrow H_2O$	7.1E9		Sehested et al. (1968)
A21004_a01	TrAa01Sc	$HO_2 + HO_2 \rightarrow H_2O_2$	9.7E5	-2500	Christensen and Sehested (1988)
A21005_a01	TrAa01Sc	$H_2O_2 + OH \rightarrow HO_2$	2.7E7	-1684	Christensen et al. (1982)
A21006_a01	TrAa01Sc	$O_3 + OH \rightarrow HO_4$	1.10E8		Staehelin et al. (1984)
A21007_a01	TrAa01Sc	$O_3 + OH^- \rightarrow HO_2 + O_2^-$	7.00E1		Staehelin et al. (1984)
A21008_a01	TrAa01Sc	$HO_3 \rightarrow OH + O_2$	1.10E5		Staehelin et al. (1984)
A21009_a01	TrAa01Sc	$HO_4 \rightarrow HO_2 + O_2$	2.80E4		Staehelin et al. (1984)
A21010_a01	TrAa01Sc	$HO_4 + HO_4 \rightarrow H_2O_2 + 2 O_3$	5.00E9		Staehelin et al. (1984)
A21011_a01	TrAa01Sc	$HO_4 + HO_3 \rightarrow H_2O_2 + O_3 + O_2$	5.00E9		Staehelin et al. (1984)
A31000_a01	TrAa01ScN	$NO_2^- + O_3 \rightarrow NO_3^-$	5.0E5	-6950	Damschen and Martin (1983)
A31001_a01	TrAa01ScN	$NO_2 + NO_2 \rightarrow HNO_3 + HONO$	1.0E8		Lee and Schwartz (1981)
A31002_a01	TrAa01ScN	$NO_4^- \rightarrow NO_2^-$	8.0E1		Warneck (1999)
A310MS_a01	TrAa01ScN	$N_2O_5 \rightarrow NO_2^+ + NO_3^-$	1.5E5		Staudt et al. (2019)
A311MS_a01	TrAa01ScN	$NO_2^+ + NO_3^- \rightarrow N_2O_5$	2.7E8		Staudt et al. (2019), Bertram and Thornton (2009)*
A32000_a01	TrAa01ScN	$NO_2 + HO_2 \rightarrow HNO_4$	1.8E9		Warneck (1999)
A32001_a01	TrAa01ScN	$NO_2^- + OH \rightarrow NO_2 + OH^-$	1.0E10		Wingenter et al. (1999)
A32002_a01	TrAa01ScN	$NO_3 + OH^- \rightarrow NO_3^- + OH$	8.2E7	-2700	Exner et al. (1992)
A32003_a01	TrAa01ScN	$HONO + OH \rightarrow NO_2$	1.0E10		Barker et al. (1970)
A32004_a01	TrAa01ScN	$HONO + H_2O_2 + H^+ \rightarrow HNO_3 + H^+$	4.6E3	-6800	Damschen and Martin (1983)
A320MS_a01	TrAa01ScN	$NO_2^+ + H_2O \rightarrow HNO_3 + H^+$	1.6E7		Staudt et al. (2019)
A321MS_a01	TrAa01ScN	$HNO_3 + H^+ \rightarrow NO_2^+ + H_2O$	1.6E-9		Sampoli et al. (1985)*
A41000_a01	TrAa01Sc	$CO_3^- + O_2^- \rightarrow HCO_3^- + OH^-$	6.5E8		Ross et al. (1992)
A41001_a01	TrAa01Sc	$CO_3^- + H_2O_2 \rightarrow HCO_3^- + HO_2$	4.3E5		Ross et al. (1992)
A41002_a01	TrAa01Sc	$HCOO^- + CO_3^- \rightarrow 2 HCO_3^- + HO_2$	1.5E5		Ross et al. (1992)
A41003_a01	TrAa01Sc	$HCOO^- + OH \rightarrow O_2^- + H_2O + CO_2$	3.1E9	-1240	Chin and Wine (1994)
A41004_a01	TrAa01ScN	$HCOO^- + NO_3 \rightarrow NO_3^- + H^+ + O_2^- + CO_2$	5.119E+07	-2200	Exner et al. (1994)
A41005_a01	TrAa01Sc	$HCOO^- + O_3 \rightarrow OH + O_2^- + CO_2$	1.00E2		Hoigné and Bader (1983)
A41006_a01	TrAa01Sc	$HCO_3^- + OH \rightarrow CO_3^-$	8.5E6		Ross et al. (1992)
A41007_a01	TrAa01Sc	$HCHO + OH \rightarrow HCOOH + HO_2$	7.7E8	-1020	Chin and Wine (1994)

Table 6: Aqueous phase reactions (...continued)

#	labels	reaction	$k_0 [M^{1-n}s^{-1}]$	$-E_a/R[K]$	reference
A41008_a01	TrAa01Sc	$\text{HCOOH} + \text{OH} \rightarrow \text{HO}_2 + \text{CO}_2$	1.1E8	-991	Chin and Wine (1994)
A41009_a01	TrAa01ScN	$\text{HCOOH} + \text{NO}_3 \rightarrow \text{NO}_3^- + \text{H}^+ + \text{HO}_2 + \text{CO}_2$	3.8E5	-3400	Exner et al. (1994)
A41010_a01	TrAa01Sc	$\text{CH}_3\text{OO} + \text{HO}_2 \rightarrow \text{CH}_3\text{OOH}$	4.3E5		Jacob (1986)
A41011_a01	TrAa01Sc	$\text{CH}_3\text{OO} + \text{O}_2^- \rightarrow \text{CH}_3\text{OOH} + \text{OH}^-$	5.0E7		Jacob (1986)
A41012a_a01	TrAa01Sc	$\text{CH}_3\text{OO} + \text{CH}_3\text{OO} \rightarrow 2 \text{HCHO} + \text{H}_2\text{O}_2$	$0.20 \times 2.145\text{E+08}$	-2139	Herrmann et al. (1999b)
A41012b_a01	TrAa01Sc	$\text{CH}_3\text{OO} + \text{CH}_3\text{OO} \rightarrow 2 \text{HOCH}_2\text{O}_2$	$0.80 \times 2.145\text{E+08}$	-2139	Herrmann et al. (1999b)
A41013a_a01	TrAa01Sc	$\text{CH}_3\text{OH} + \text{OH} \rightarrow \text{HOCH}_2\text{O}_2 + \text{H}_2\text{O}$	$0.93 \times 9.70\text{E8}$	-600	Elliot and McCracken (1989)*
A41013b_a01	TrAa01Sc	$\text{CH}_3\text{OH} + \text{OH} \rightarrow \text{HCHO} + \text{HO}_2 + \text{H}_2\text{O}$	$0.07 \times 9.70\text{E8}$	-600	Elliot and McCracken (1989)
A41014_a01	TrAa01ScN	$\text{CH}_3\text{OH} + \text{NO}_3 \rightarrow \text{HOCH}_2\text{O}_2 + \text{NO}_3^- + \text{H}^+$	5.40E5	-4300	Herrmann and Zellner (1998)
A41015_a01	TrAa01Sc	$\text{CH}_3\text{OH} + \text{CO}_3^- \rightarrow \text{HOCH}_2\text{O}_2 + \text{HCO}_3^-$	5.431E+03	-3100	Clifton and Huie (1993)
A41016a_a01	TrAa01Sc	$\text{CH}_3\text{OOH} + \text{OH} \rightarrow \text{CH}_3\text{OO} + \text{H}_2\text{O}$	$0.25 \times 6.30\text{E8}$		Monod et al. (2007)*
A41016b_a01	TrAa01Sc	$\text{CH}_3\text{OOH} + \text{OH} \rightarrow \text{HCHO} + \text{OH} + \text{H}_2\text{O}$	$0.75 \times 6.30\text{E8}$		Monod et al. (2007)*
A41017a_a01	TrAa01ScN	$\text{CH}_3\text{OOH} + \text{NO}_3 \rightarrow \text{CH}_3\text{OO} + \text{NO}_3^- + \text{H}^+$	$0.25 \times 4.90\text{E6}$	-2000	see note*
A41017b_a01	TrAa01ScN	$\text{CH}_3\text{OOH} + \text{NO}_3 \rightarrow \text{HCHO} + \text{HO}_2 + \text{NO}_3^- + \text{H}^+$	$0.75 \times 4.90\text{E6}$	-2000	see note*
A41018a_a01	TrAa01Sc	$\text{CH}_3\text{OOH} + \text{CO}_3^- \rightarrow \text{CH}_3\text{OO} + \text{HCO}_3^-$	$0.25 \times 4.30\text{E5}$		see note*
A41018b_a01	TrAa01Sc	$\text{CH}_3\text{OOH} + \text{CO}_3^- \rightarrow \text{HCHO} + \text{HO}_2 + \text{HCO}_3^-$	$0.75 \times 4.30\text{E5}$		see note*
A41019a_a01	TrAa01Sc	$\text{HOCH}_2\text{OOH} + \text{OH} \rightarrow \text{HOCH}_2\text{O}_2 + \text{H}_2\text{O}$	$0.25 \times 6.30\text{E8}$		see note*
A41019b_a01	TrAa01Sc	$\text{HOCH}_2\text{OOH} + \text{OH} \rightarrow \text{CHOOOH} + \text{HO}_2 + \text{H}_2\text{O}$	$0.75 \times 6.30\text{E8}$		see note*
A41020a_a01	TrAa01ScN	$\text{HOCH}_2\text{OOH} + \text{NO}_3 \rightarrow \text{HOCH}_2\text{O}_2 + \text{NO}_3^- + \text{H}^+$	$0.25 \times 4.90\text{E6}$	-2000	see note*
A41020b_a01	TrAa01ScN	$\text{HOCH}_2\text{OOH} + \text{NO}_3 \rightarrow \text{CHOOOH} + \text{HO}_2 + \text{NO}_3^- + \text{H}^+$	$0.75 \times 4.90\text{E6}$	-2000	see note*
A41021_a01	TrAa01Sc	$\text{HOCH}_2\text{O}_2 \rightarrow \text{HCHO} + \text{HO}_2$	1.00E1		see note*
A41022_a01	TrAa01Sc	$\text{HOCH}_2\text{O}_2 + \text{HO}_2 \rightarrow \text{HOCH}_2\text{OOH} + \text{O}_2$	9.7E5	-2500	see note*
A41023_a01	TrAa01Sc	$\text{HOCH}_2\text{O}_2 + \text{O}_2^- \rightarrow \text{HOCH}_2\text{OOH} + \text{O}_2 + \text{OH}^-$	1.0E8	-900	see note*
A41024_a01	TrAa01Sc	$\text{HOCH}_2\text{O}_2 + \text{HOCH}_2\text{O}_2 \rightarrow 2 \text{HCOOH} + \text{H}_2\text{O}_2$	7.367E+08	-1395	Huie and Clifton (1993)
A41025_a01	TrAa01Sc	$\text{HCOOH} + \text{H}_2\text{O}_2 + \text{H}^+ \rightarrow \text{CHOOOH} + \text{H}_2\text{O}_2 + \text{H}^+$	3.080E-04	-5235	De Filippis et al. (2009)
A41026a_a01	TrAa01Sc	$\text{CHOOOH} + \text{H}^+ \rightarrow \text{HCOOH} + \text{H}_2\text{O}_2 + \text{H}^+$	3.790E-04	-5235	De Filippis et al. (2009)
A41026b_a01	TrAa01Sc	$\text{CHOOOH} + \text{H}^+ \rightarrow \text{CO}_2 + \text{H}_2\text{O} + \text{H}^+$	1.219E-03	-8735	De Filippis et al. (2009)
A41027_a01	TrAa01Sc	$\text{HOCH}_2\text{OH} + \text{OH} \rightarrow \text{HCOHOHO}_2 + \text{H}_2\text{O}$	7.70E8	-1000	Chin and Wine (1994)
A41028_a01	TrAa01Sc	$\text{HOCH}_2\text{OH} + \text{CO}_3^- \rightarrow \text{HCO}_3^- + \text{HCOHOHO}_2$	1.30E4		Zellner et al. (1996)
A41029_a01	TrAa01ScN	$\text{HOCH}_2\text{OH} + \text{NO}_3 \rightarrow \text{NO}_3^- + \text{H}^+ + \text{HCOHOHO}_2$	1.003E+06	-4500	Exner et al. (1993)
A41030_a01	TrAa01Sc	$\text{HCOHOHO}_2 \rightarrow \text{HCOOH} + \text{HO}_2$	1.00E6		see note*
A41030DT_a01	TrAa01ScN	$\text{HCOO}^- + \text{NO}_2^+ \rightarrow \text{HCOO}^- + \text{NO}_3^- + 2 \text{H}^+$	7.50E+09		Staudt et al. (2019)
A41030MS_a01	TrAa01ScN	$\text{CH}_3\text{OH} + \text{NO}_2^+ \rightarrow \text{CH}_3\text{ONO}_2 + \text{H}^+$	4.50E+08		Iraci et al. (2007)

Table 6: Aqueous phase reactions (...continued)

#	labels	reaction	$k_0 [M^{1-n}s^{-1}]$	$-E_a/R[K]$	reference
A42000a_a01	TrAa01ScC	$\text{CH}_3\text{CH}_2\text{OH} + \text{OH} \rightarrow \text{CH}_3\text{CHO} + \text{HO}_2 + \text{H}_2\text{O}$	$0.90 \times 2.002\text{E}+09$	-830	Monod et al. (2005)*
A42000b_a01	TrAa01ScC	$\text{CH}_3\text{CH}_2\text{OH} + \text{OH} \rightarrow \text{CH}_2\text{OHCH}_2\text{OO} + \text{H}_2\text{O}$	$0.10 \times 2.002\text{E}+09$	-830	Monod et al. (2005)
A42001a_a01	TrAa01ScCN	$\text{CH}_3\text{CH}_2\text{OH} + \text{NO}_3 \rightarrow \text{CH}_3\text{CHO} + \text{HO}_2 + \text{NO}_3^- + \text{H}^+$	$0.90 \times 2.184\text{E}+06$	-3300	Herrmann and Zellner (1998)*
A42001b_a01	TrAa01ScCN	$\text{CH}_3\text{CH}_2\text{OH} + \text{NO}_3 \rightarrow \text{CH}_2\text{OHCH}_2\text{OO} + \text{NO}_3^- + \text{H}^+$	$0.10 \times 2.184\text{E}+06$	-3300	Herrmann and Zellner (1998)
A42002a_a01	TrAa01ScC	$\text{CH}_2\text{OHCH}_2\text{OO} + \text{CH}_2\text{OHCH}_2\text{OO} \rightarrow \text{CH}_2\text{OHCHO} + \text{CH}_2\text{OHCHO} + \text{H}_2\text{O}_2$	$0.50 \times 1.00\text{E}8$		Piesiak et al. (1984)
A42002b_a01	TrAa01ScC	$\text{CH}_2\text{OHCH}_2\text{OO} + \text{CH}_2\text{OHCH}_2\text{OO} \rightarrow \text{CH}_2\text{OHCHO} + \text{ETHGLY}$	$0.33 \times 1.00\text{E}8$		Piesiak et al. (1984)
A42002c_a01	TrAa01ScC	$\text{CH}_2\text{OHCH}_2\text{OO} + \text{CH}_2\text{OHCH}_2\text{OO} \rightarrow 2 \text{ HOCH}_2\text{O}_2 + 2 \text{ HCHO}$	$0.17 \times 1.00\text{E}8$		Piesiak et al. (1984)
A42003_a01	TrAa01ScC	$\text{CH}_2\text{OHCH}_2\text{OO} + \text{O}_2^- \rightarrow \text{HYETHO2H} + \text{OH}^-$	1.0E8	-900	see note*
A42004_a01	TrAa01ScC	$\text{CH}_2\text{OHCH}_2\text{OO} + \text{HO}_2 \rightarrow \text{HYETHO2H}$	9.7E5	-2500	see note*
A42005_a01	TrAa01ScC	$\text{HYETHO2H} + \text{OH} \rightarrow \text{CH}_2\text{OHCHO}$	1.10E9		see note*
A42006_a01	TrAa01ScC	$\text{ETHGLY} + \text{OH} \rightarrow \text{CH}_2\text{OHCHO} + \text{HO}_2 + \text{H}_2\text{O}$	$1.657\text{E}+09$	-1191	Hoffmann et al. (2009)*
A42007_a01	TrAa01ScCN	$\text{ETHGLY} + \text{NO}_3 \rightarrow \text{CH}_2\text{OHCHO} + \text{HO}_2 + \text{NO}_3^- + \text{H}^+$	$5.856\text{E}+06$	-2117	Hoffmann et al. (2009)*
A42008_a01	TrAa01ScC	$\text{CH}_3\text{CHO} + \text{OH} \rightarrow \text{CH}_3\text{COOO} + \text{H}_2\text{O}$	3.60E9		Schuchmann and von Sonntag (1988)
A42009_a01	TrAa01ScCN	$\text{CH}_3\text{CHO} + \text{NO}_3 \rightarrow \text{CH}_3\text{COOO} + \text{NO}_3^- + \text{H}^+$	3.10E6		Rousse and George (2004)
A42010_a01	TrAa01ScC	$\text{CH}_3\text{COOO} + \text{CH}_3\text{COOO} \rightarrow \text{CH}_3\text{OO} + \text{CH}_3\text{OO} + \text{CO}_2 + \text{CO}_2$	$1.891\text{E}+08$	1563	see note*
A42011_a01	TrAa01ScC	$\text{CH}_3\text{COOO} + \text{O}_2^- \rightarrow \text{CH}_3\text{COOO}^- + \text{O}_2$	1.00E9		Schuchmann and von Sonntag (1988)
A42012_a01	TrAa01ScC	$\text{CH}_3\text{CHOHOH} + \text{OH} \rightarrow \text{CH}_3\text{COHOHO} + \text{H}_2\text{O}$	1.20E9		Schuchmann and von Sonntag (1988)
A42013_a01	TrAa01ScCN	$\text{CH}_3\text{CHOHOH} + \text{NO}_3 \rightarrow \text{CH}_3\text{COHOHO} + \text{NO}_3^- + \text{H}^+$	1.10E6		Rousse and George (2004)
A42014_a01	TrAa01ScC	$\text{CH}_3\text{COHOHO} \rightarrow \text{CH}_3\text{COOH} + \text{HO}_2$	1.00E6		see note*
A42015a_a01	TrAa01ScC	$\text{CH}_2\text{OHCHO} + \text{OH} \rightarrow \text{CH}_2\text{OHCO}_3 + \text{H}_2\text{O}$	$0.77 \times 1.40\text{E}9$		Doussin and Monod (2013)
A42015b_a01	TrAa01ScC	$\text{CH}_2\text{OHCHO} + \text{OH} \rightarrow \text{CHOHOOCCHO} + \text{H}_2\text{O}$	$0.23 \times 1.40\text{E}9$		Doussin and Monod (2013)
A42016a_a01	TrAa01ScCN	$\text{CH}_2\text{OHCHO} + \text{NO}_3 \rightarrow \text{CH}_2\text{OHCO}_3 + \text{NO}_3^- + \text{H}^+$	$0.77 \times 3.10\text{E}6$		see note*
A42016b_a01	TrAa01ScCN	$\text{CH}_2\text{OHCHO} + \text{NO}_3 \rightarrow \text{CHOHOOCCHO} + \text{NO}_3^- + \text{H}^+$	$0.23 \times 3.10\text{E}6$		see note*

Table 6: Aqueous phase reactions (...continued)

#	labels	reaction	$k_0 [M^{1-n}s^{-1}]$	$-E_a/R[K]$	reference
A42017_a01	TrAa01ScC	$\text{CH}_2\text{OHCO}_3 + \text{O}_2^- \rightarrow \text{CH}_2\text{OHCO}_2\text{O}^-$	1.00E9		see note*
A42018_a01	TrAa01ScC	$\text{CH}_2\text{OHCOHOHO}_2 \rightarrow \text{HOCH}_2\text{CO}_2\text{H} + \text{HO}_2$	1.00E6		see note*
A42019_a01	TrAa01ScC	$\text{CHOHOOCHO} \rightarrow \text{GLYOX} + \text{HO}_2$	1.90E2		see note*
A42020_a01	TrAa01ScC	$\text{CHOHOOCHOHOH} \rightarrow \text{CHOCHOHOH} + \text{HO}_2$	1.90E2		see note*
A42021a_a01	TrAa01ScC	$\text{CH}_2\text{OHCHOHOH} + \text{OH} \rightarrow \text{CH}_2\text{OHCOHOHO}_2 + \text{H}_2\text{O}$	$0.33 \times 1.10\text{E}9$		Doussin and Monod (2013)
A42021b_a01	TrAa01ScC	$\text{CH}_2\text{OHCHOHOH} + \text{OH} \rightarrow \text{CHOHOOCHOHOH} + \text{H}_2\text{O}$	$0.28 \times 1.10\text{E}9$		Doussin and Monod (2013)
A42021c_a01	TrAa01ScC	$\text{CH}_2\text{OHCHOHOH} + \text{OH} \rightarrow \text{HCOOH} + \text{HOCH}_2\text{O}_2 + \text{H}_2\text{O}$	$0.39 \times 1.10\text{E}9$		Doussin and Monod (2013)
A42022a_a01	TrAa01ScCN	$\text{CH}_2\text{OHCHOHOH} + \text{NO}_3 \rightarrow \text{CH}_2\text{OHCOHOHO}_2 + \text{NO}_3^- + \text{H}^+$	$0.33 \times 1.10\text{E}6$		see note*
A42022b_a01	TrAa01ScCN	$\text{CH}_2\text{OHCHOHOH} + \text{NO}_3 \rightarrow \text{CHOHOOCHOHOH} + \text{NO}_3^- + \text{H}^+$	$0.28 \times 1.10\text{E}6$		see note*
A42022c_a01	TrAa01ScCN	$\text{CH}_2\text{OHCHOHOH} + \text{NO}_3 \rightarrow \text{HCOOH} + \text{HOCH}_2\text{O}_2 + \text{NO}_3^- + \text{H}^+$	$0.39 \times 1.10\text{E}6$		see note*
A42023a_a01	TrAa01ScC	$\text{CHOHOHCHOHOH} + \text{OH} \rightarrow \text{CHOHOHCOHOHO}_2 + \text{H}_2\text{O}$	$0.27 \times 1.114\text{E}+09$	-1516	Buxton et al. (1997)
A42023b_a01	TrAa01ScC	$\text{CHOHOHCHOHOH} + \text{OH} \rightarrow \text{HCOOH} + \text{HCOHOHO}_2 + \text{HO}_2 + \text{H}_2\text{O}$	$0.73 \times 1.114\text{E}+09$	-1516	Buxton et al. (1997)*
A42024a_a01	TrAa01ScCN	$\text{CHOHOHCHOHOH} + \text{NO}_3 \rightarrow \text{CHOHOHCOHOHO}_2 + \text{NO}_3^- + \text{H}^+$	$0.27 \times 1.00\text{E}6$		see note*
A42024b_a01	TrAa01ScCN	$\text{CHOHOHCHOHOH} + \text{NO}_3 \rightarrow \text{HCOOH} + \text{HCOHOHO}_2 + \text{NO}_3^- + \text{H}^+$	$0.73 \times 1.00\text{E}6$		see note*
A42025_a01	TrAa01ScC	$\text{CHOHOHCOHOHO}_2 \rightarrow \text{CHOHOHCOOH} + \text{HO}_2$	$0.77 \times 1.00\text{E}6$		see note*
A42026_a01	TrAa01ScC	$\text{CH}_3\text{COOH} + \text{OH} \rightarrow \text{CH}_2\text{OOCOOH} + \text{H}_2\text{O}$	1.50E7	-1330	Chin and Wine (1994)
A42027_a01	TrAa01ScCN	$\text{CH}_3\text{COOH} + \text{NO}_3 \rightarrow \text{CH}_2\text{OOCOOH} + \text{NO}_3^- + \text{H}^+$	1.429E+04	-3800	Exner et al. (1994)
A42028_a01	TrAa01ScC	$\text{CH}_3\text{COO}^- + \text{OH} \rightarrow \text{CH}_2\text{OOCO}_2^- + \text{H}_2\text{O}$	1.00E8	-1800	Fisher and Hamill (1973)
A42029_a01	TrAa01ScCN	$\text{CH}_3\text{COO}^- + \text{NO}_3 \rightarrow \text{CH}_2\text{OOCO}_2^- + \text{NO}_3^- + \text{H}^+$	2.916E+06	-3800	Exner et al. (1994)
A42030DT_a01	TrAa01ScCN	$\text{CH}_3\text{COO}^- + \text{NO}_2^+ \rightarrow \text{CH}_3\text{COO}^- + \text{NO}_3^- + 2 \text{H}^+$	7.50E+09		Staudt et al. (2019)
A42030a_a01	TrAa01ScC	$\text{C}_2\text{H}_5\text{OOH} + \text{OH} \rightarrow \text{C}_2\text{H}_5\text{OO} + \text{H}_2\text{O}$	$0.80 \times 5.80\text{E}8$		Monod et al. (2007)
A42030b_a01	TrAa01ScC	$\text{C}_2\text{H}_5\text{OOH} + \text{OH} \rightarrow \text{CH}_3\text{COOH} + \text{HO}_2 + \text{H}_2\text{O}$	$0.20 \times 5.80\text{E}8$		Monod et al. (2007)*
A42031a_a01	TrAa01ScC	$\text{C}_2\text{H}_5\text{OO} + \text{C}_2\text{H}_5\text{OO} \rightarrow \text{CH}_3\text{CHO} + \text{CH}_3\text{CHO} + \text{H}_2\text{O}_2$	$0.20 \times 1.891\text{E}+08$	1563	Herrmann et al. (1999b)
A42031b_a01	TrAa01ScC	$\text{C}_2\text{H}_5\text{OO} + \text{C}_2\text{H}_5\text{OO} \rightarrow 2 \text{CH}_3\text{CHO} + 2 \text{HO}_2$	$0.80 \times 1.891\text{E}+08$	1563	Herrmann et al. (1999b)*

Table 6: Aqueous phase reactions (...continued)

#	labels	reaction	$k_0 [M^{1-n}s^{-1}]$	$-E_a/R[K]$	reference
A42032_a01	TrAa01ScC	$C_2H_5OO + O_2^- \rightarrow C_2H_5OOH + OH^-$	1.0E8	-900	see note*
A42033_a01	TrAa01ScC	$C_2H_5OO + HO_2 \rightarrow C_2H_5OOH$	9.7E5	-2500	see note*
A42034_a01	TrAa01ScC	$CH_2OOCOOH + HO_2 \rightarrow HOOCH_2CO_2H$	9.7E5	-2500	see note*
A42035_a01	TrAa01ScC	$CH_2OOCOOH + O_2^- + H^+ \rightarrow HOOCH_2CO_2H$	1.0E8	-900	see note*
A42036a_a01	TrAa01ScC	$CH_2OOCOOH + CH_2OOCOOH \rightarrow CHOCOOH + CHOCOOH + H_2O_2$	$0.30 \times 7.50E7$		Schuchmann et al. (1985)
A42036b_a01	TrAa01ScC	$CH_2OOCOOH + CH_2OOCOOH \rightarrow 2 HCHO + 2 CO_2 + H_2O_2$	$0.30 \times 7.50E7$		Schuchmann et al. (1985)
A42036c_a01	TrAa01ScC	$CH_2OOCOOH + CH_2OOCOOH \rightarrow CHOCOOH + HOCH_2CO_2H$	$0.30 \times 7.50E7$		Schuchmann et al. (1985)
A42036d_a01	TrAa01ScC	$CH_2OOCOOH + CH_2OOCOOH \rightarrow CHOHOOCOOH + CHOHOOCOOH$	$0.10 \times 7.50E7$		Schuchmann et al. (1985)
A42037_a01	TrAa01ScC	$CH_2OOCO_2^- + HO_2 \rightarrow CH_2OOHCO_2^- + O_2$	9.7E5	-2500	see note*
A42038_a01	TrAa01ScC	$CH_2OOCO_2^- + O_2^- + H^+ \rightarrow CH_2OOHCO_2^-$	1.0E8	-900	see note*
A42039a_a01	TrAa01ScC	$CH_2OOCO_2^- + CH_2OOCO_2^- \rightarrow CHOCOO^- + CHOCOO^- + H_2O_2$	$0.30 \times 7.50E7$		Schuchmann et al. (1985)
A42039b_a01	TrAa01ScC	$CH_2OOCO_2^- + CH_2OOCO_2^- \rightarrow 2 HCHO + 2 CO_2 + H_2O_2 + 2 OH^-$	$0.30 \times 7.50E7$		Schuchmann et al. (1985)
A42039c_a01	TrAa01ScC	$CH_2OOCO_2^- + CH_2OOCO_2^- \rightarrow CHOCOO^- + CH_2OHCC_2^-$	$0.30 \times 7.50E7$		Schuchmann et al. (1985)
A42039d_a01	TrAa01ScC	$CH_2OOCO_2^- + CH_2OOCO_2^- \rightarrow 2 CHOHOOCOO_2^-$	$0.10 \times 7.50E7$		Schuchmann et al. (1985)
A42040_a01	TrAa01ScC	$CH_2OOCO_2^- + O_3 \rightarrow O_3^- + HOCH_2OOH + CO_2$	2.00E9		Sehested et al. (1984)
A42141_a01	TrAa01ScC	$HOOCCOO^- + OH \rightarrow C_2O_4^{2-} + H_2O$	2.08E+08	-2800	Ervens et al. (2003)
A42142_a01	TrAa01ScC	$C_2O_4^{2-} + OH \rightarrow C_2O_4^- + OH^-$	2.508E+08	-4300	Ervens et al. (2003)
A42143_a01	TrAa01ScC	$C_2O_4^- + O_2 \rightarrow 2 CO_2 + O_2^-$	2.40E9		Hislop and Bolton (1999)
A42144a_a01	TrAa01ScC	$HOOCH_2CO_2H + OH \rightarrow CH_2OOCOOH + H_2O$	$0.80 \times 5.80E8$		see note*
A42144b_a01	TrAa01ScC	$HOOCH_2CO_2H + OH \rightarrow CHOOHOOCOOH + H_2O$	$0.20 \times 5.80E8$		see note*
A42145a_a01	TrAa01ScCN	$HOOCH_2CO_2H + NO_3 \rightarrow CH_2OOCOOH + NO_3^- + H^+$	$0.80 \times 1.70E6$		Herrmann and Zellner (1998)
A42145b_a01	TrAa01ScCN	$HOOCH_2CO_2H + NO_3 \rightarrow CHOOHOOCOOH + NO_3^- + H^+$	$0.20 \times 1.70E6$		Herrmann and Zellner (1998)
A42146a_a01	TrAa01ScC	$CH_2OOHCO_2^- + OH \rightarrow CH_2OOCO_2^- + H_2O$	$0.80 \times 5.80E8$		see note*
A42146b_a01	TrAa01ScC	$CH_2OOHCO_2^- + OH \rightarrow CHOOHOOCO_2^- + H_2O$	$0.20 \times 5.80E8$		see note*

Table 6: Aqueous phase reactions (...continued)

#	labels	reaction	$k_0 [M^{1-n}s^{-1}]$	$-E_a/R[K]$	reference
A42147a_a01	TrAa01ScCN	$\text{CH}_2\text{OOHCO}_2^- + \text{NO}_3 \rightarrow \text{CH}_2\text{OOCO}_2^- + \text{NO}_3^- + \text{H}^+$	$0.80 \times 7.10\text{E}6$		Herrmann and Zellner (1998)
A42147b_a01	TrAa01ScCN	$\text{CH}_2\text{OOHCO}_2^- + \text{NO}_3 \rightarrow \text{CHOHOOCO}_2^- + \text{NO}_3^- + \text{H}^+$	$0.20 \times 7.10\text{E}6$		Herrmann and Zellner (1998)
A42148_a01	TrAa01ScC	$\text{CHOHOOCOOH} \rightarrow \text{HOOCOOH} + \text{HO}_2$	1.90E2		see note*
A42149_a01	TrAa01ScC	$\text{CHOHOOCO}_2^- \rightarrow \text{HOOCOO}^- + \text{HO}_2$	1.90E2		see note*
A42150a_a01	TrAa01ScC	$\text{HOCH}_2\text{CO}_2\text{H} + \text{OH} \rightarrow \text{CHOHOOCOOH} + \text{H}_2\text{O}$	$0.62 \times 6.00\text{E}8$		see note*
A42150b_a01	TrAa01ScC	$\text{HOCH}_2\text{CO}_2\text{H} + \text{OH} \rightarrow \text{HCHO} + \text{CO}_2 + \text{HO}_2 + \text{H}_2\text{O}$	$0.38 \times 6.00\text{E}8$		see note*
A42151a_a01	TrAa01ScCN	$\text{HOCH}_2\text{CO}_2\text{H} + \text{NO}_3 \rightarrow \text{CHOHOOCOOH} + \text{NO}_3^- + \text{H}^+$	$0.62 \times 7.445\text{E}+05$	-3969	see note*
A42151b_a01	TrAa01ScCN	$\text{HOCH}_2\text{CO}_2\text{H} + \text{NO}_3 \rightarrow \text{HCHO} + \text{CO}_2 + \text{HO}_2 + \text{NO}_3^- + \text{H}^+$	$0.38 \times 7.445\text{E}+05$	-3969	see note*
A42152_a01	TrAa01ScC	$\text{CHOHOOCOOH} \rightarrow \text{CHOCOOH} + \text{HO}_2$	1.90E2		von Sonntag (1987)
A42153a_a01	TrAa01ScC	$\text{CH}_2\text{OHCO}_2^- + \text{OH} \rightarrow \text{CHOHOOCO}_2^- + \text{H}_2\text{O}$	$0.60 \times 8.60\text{E}8$		Buxton et al. (1988)
A42153b_a01	TrAa01ScC	$\text{CH}_2\text{OHCO}_2^- + \text{OH} \rightarrow \text{HCHO} + \text{CO}_2 + \text{O}_2^- + \text{H}_2\text{O}$	$0.19 \times 8.60\text{E}8$		Buxton et al. (1988)
A42153c_a01	TrAa01ScC	$\text{CH}_2\text{OHCO}_2^- + \text{OH} \rightarrow \text{HOCH}_2\text{O}_2 + \text{CO}_2 + \text{OH}^-$	$0.21 \times 8.60\text{E}8$		Buxton et al. (1988)
A42154a_a01	TrAa01ScCN	$\text{CH}_2\text{OHCO}_2^- + \text{NO}_3 \rightarrow \text{CHOHOOCO}_2^- + \text{NO}_3^- + \text{H}^+$	$0.76 \times 7.502\text{E}+06$	-3007	Gaillard de Sémainville et al. (2007)
A42154b_a01	TrAa01ScCN	$\text{CH}_2\text{OHCO}_2^- + \text{NO}_3 \rightarrow \text{HCHO} + \text{CO}_2 + \text{O}_2^- + \text{NO}_3^- + \text{H}^+$	$0.24 \times 7.502\text{E}+06$	-3007	Gaillard de Sémainville et al. (2007)
A42155_a01	TrAa01ScC	$\text{CHOHOOCO}_2^- \rightarrow \text{CHOCOO}^- + \text{HO}_2$	1.90E2		von Sonntag (1987)
A42156a_a01	TrAa01ScC	$\text{CHOHOHC}(\text{OOH}) + \text{OH} \rightarrow \text{COOHCOHOHO}_2 + \text{H}_2\text{O}$	$0.15 \times 2.830\text{E}+08$	-1000	Ervens et al. (2003)
A42156b_a01	TrAa01ScC	$\text{CHOHOHC}(\text{OOH}) + \text{OH} \rightarrow \text{HCOOH} + \text{CO}_2 + \text{HO}_2 + \text{H}_2\text{O}$	$0.85 \times 2.830\text{E}+08$	-1000	Ervens et al. (2003)*
A42157a_a01	TrAa01ScCN	$\text{CHOHOHC}(\text{OOH}) + \text{NO}_3 \rightarrow \text{COOHCOHOHO}_2 + \text{NO}_3^- + \text{H}^+$	$0.15 \times 1.00\text{E}6$		see note*
A42157b_a01	TrAa01ScCN	$\text{CHOHOHC}(\text{OOH}) + \text{NO}_3 \rightarrow \text{HCOOH} + \text{CO}_2 + \text{HO}_2 + \text{NO}_3^- + \text{H}^+$	$0.85 \times 1.00\text{E}6$		see note*
A42158a_a01	TrAa01ScC	$\text{CHOHOHC}(\text{OOH}) + \text{OH} \rightarrow \text{CO}_2^- \text{COHOHO}_2 + \text{H}_2\text{O}$	$0.26 \times 3.271\text{E}+09$	-4300	Ervens et al. (2003)
A42158b_a01	TrAa01ScC	$\text{CHOHOHC}(\text{OOH}) + \text{OH} \rightarrow \text{HCOOH} + \text{CO}_2 + \text{O}_2^- + \text{H}_2\text{O}$	$0.74 \times 3.271\text{E}+09$	-4300	Ervens et al. (2003)
A42159a_a01	TrAa01ScCN	$\text{CHOHOHC}(\text{OOH}) + \text{NO}_3 \rightarrow \text{CO}_2^- \text{COHOHO}_2 + \text{NO}_3^- + \text{H}^+$	$0.26 \times 1.80\text{E}5$		Herrmann and Zellner (1998)

Table 6: Aqueous phase reactions (...continued)

#	labels	reaction	$k_0 [M^{1-n}s^{-1}]$	$-E_a/R[K]$	reference
A42159b_a01	TrAa01ScCN	$\text{CHOHOHCO}_2^- + \text{NO}_3 \rightarrow \text{HCOOH} + \text{CO}_2 + \text{O}_2^- + \text{NO}_3^- + \text{H}^+$	$0.74 \times 1.80\text{E}5$		Herrmann and Zellner (1998)
A42160_a01	TrAa01ScC	$\text{CHOHOHCO}_2^- + \text{H}_2\text{O}_2 \rightarrow \text{HCOO}^- + \text{CO}_2 + \text{H}_2\text{O} + \text{H}^+$	1.10E-1		Schöne and Herrmann (2014)
A42161_a01	TrAa01ScC	$\text{COOHCOHOHO}_2 \rightarrow \text{HOOCOOH} + \text{HO}_2$	1.00E6		see note*
A42162_a01	TrAa01ScC	$\text{CO}_2^- \text{COHOHO}_2 \rightarrow \text{HOOCOO}^- + \text{HO}_2$	1.00E6		see note*
A42163a_a01	TrAa01ScC	$\text{CH}_2\text{OOHCHO} + \text{OH} \rightarrow \text{HCHO} + \text{CO} + \text{OH} + \text{H}_2\text{O}$	$0.77 \times 1.40\text{E}9$		see note*
A42163b_a01	TrAa01ScC	$\text{CH}_2\text{OOHCHO} + \text{OH} \rightarrow \text{GLYOX} + \text{HO}_2 + \text{H}_2\text{O}$	$0.23 \times 1.40\text{E}9$		see note*
A42164a_a01	TrAa01ScCN	$\text{CH}_2\text{OOHCHO} + \text{NO}_3 \rightarrow \text{HCHO} + \text{CO} + \text{NO}_3^- + \text{H}_2\text{O}$	$0.77 \times 3.10\text{E}6$		see note*
A42164b_a01	TrAa01ScCN	$\text{CH}_2\text{OOHCHO} + \text{NO}_3 \rightarrow \text{GLYOX} + \text{NO}_3^- + \text{H}_2\text{O} + \text{H}^+$	$0.23 \times 3.10\text{E}6$		see note*
A42165a_a01	TrAa01ScC	$\text{HOOCH}_2\text{CHOHOH} + \text{OH} \rightarrow \text{HOOCH}_2\text{CO}_2\text{H} + \text{HO}_2 + \text{H}_2\text{O}$	$0.33 \times 1.10\text{E}9$		see note*
A42165b_a01	TrAa01ScC	$\text{HOOCH}_2\text{CHOHOH} + \text{OH} \rightarrow \text{CHOCHOHOH} + \text{OH} + \text{H}_2\text{O}$	$0.28 \times 1.10\text{E}9$		see note*
A42165c_a01	TrAa01ScC	$\text{HOOCH}_2\text{CHOHOH} + \text{OH} \rightarrow \text{HCOOH} + \text{HCHO} + \text{OH} + \text{H}_2\text{O}$	$0.39 \times 1.10\text{E}9$		see note*
A42166a_a01	TrAa01ScCN	$\text{HOOCH}_2\text{CHOHOH} + \text{NO}_3 \rightarrow \text{HOOCH}_2\text{CO}_2\text{H} + \text{NO}_3^- + \text{H}_2\text{O} + \text{H}^+$	$0.33 \times 1.10\text{E}6$		see note*
A42166b_a01	TrAa01ScCN	$\text{HOOCH}_2\text{CHOHOH} + \text{NO}_3 \rightarrow \text{CHOCHOHOH} + \text{NO}_3^- + \text{H}_2\text{O}$	$0.28 \times 1.10\text{E}6$		see note*
A42166c_a01	TrAa01ScCN	$\text{HOOCH}_2\text{CHOHOH} + \text{NO}_3 \rightarrow \text{HCOOH} + \text{HCHO} + \text{NO}_3^- + \text{H}_2\text{O}$	$0.39 \times 1.10\text{E}6$		see note*
A42167_a01	TrAa01ScC	$\text{HOCH}_2\text{OH} + \text{HOCH}_2\text{OH} \rightarrow \text{MG2} + \text{H}_2\text{O}$	see note	see note	Hahnenstein et al. (1995)*
A42168_a01	TrAa01ScC	$\text{MG2} + \text{H}_2\text{O} \rightarrow \text{HOCH}_2\text{OH} + \text{HOCH}_2\text{OH}$	see note	see note	Hahnenstein et al. (1995)
A42169_a01	TrAa01ScC	$\text{MG2} + \text{OH} \rightarrow \text{HMF} + \text{HO}_2$	1.54E9	-1020	see note*
A42470_a01	TrAa01ScC	$\text{CH}_3\text{COOO} + \text{H}_2\text{O} \rightarrow \text{CH}_3\text{COOH} + \text{HO}_2$	7.0E5		Villalta et al. (1996)
A42471_a01	TrAa01ScC	$\text{CH}_2\text{OHCO}_3 + \text{H}_2\text{O} \rightarrow \text{HOCH}_2\text{CO}_2\text{H} + \text{HO}_2$	7.0E5		see note*
A42472_a01	TrAa01ScC	$\text{CHOCO}_3 + \text{H}_2\text{O} \rightarrow \text{CHOCOOH} + \text{HO}_2$	7.0E5		see note*
A42473_a01	TrAa01ScC	$\text{COOHCO}_3 + \text{H}_2\text{O} \rightarrow \text{HOOCOOH} + \text{HO}_2$	7.0E5		see note*
A43000a_a01	TrAa01ScC	$\text{CH}_3\text{COCHOHOH} + \text{OH} \rightarrow \text{CH}_3\text{COCOOH} + \text{HO}_2$	$0.29 \times 9.215\text{E}+08$	-1235	Schaefer et al. (2015)*
A43000b_a01	TrAa01ScC	$\text{CH}_3\text{COCHOHOH} + \text{OH} \rightarrow \text{HCOOH} + \text{CH}_3\text{COOO}$	$0.71 \times 9.215\text{E}+08$	-1235	Schaefer et al. (2015)
A43001_a01	TrAa01ScCN	$\text{CH}_3\text{COCHOHOH} + \text{NO}_3 \rightarrow \text{CH}_3\text{COCOOH} + \text{NO}_3^- + \text{H}^+$	4.539E+06	-4213	Schaefer et al. (2015)*

Table 6: Aqueous phase reactions (...continued)

#	labels	reaction	$k_0 [M^{1-n}s^{-1}]$	$-E_a/R[K]$	reference
A43002_a01	TrAa01ScC	$\text{CH}_3\text{COCOOH} + \text{OH} \rightarrow \text{CH}_3\text{COOH} + \text{HO}_2 + \text{CO}_2$	2.592E+08	-1804	Schaefer et al. (2012)*
A43003_a01	TrAa01ScCN	$\text{CH}_3\text{COCOOH} + \text{NO}_3 \rightarrow \text{CH}_3\text{COOH} + \text{NO}_3^- + \text{CO}_2 + \text{H}^+$	2.828E+06	-1804	Gaillard de Sémainville et al. (2007)
A43004_a01	TrAa01ScC	$\text{CH}_3\text{COCO}_2^- + \text{OH} \rightarrow \text{CH}_3\text{COO}^- + \text{HO}_2 + \text{CO}_2$	6.252E+08	-3007	Schaefer et al. (2012)*
A43005_a01	TrAa01ScCN	$\text{CH}_3\text{COCO}_2^- + \text{NO}_3 \rightarrow \text{CH}_3\text{COO}^- + \text{NO}_3^- + \text{CO}_2 + \text{H}^+$	2.306E+07	-2887	Gaillard de Sémainville et al. (2007)
A43006_a01	TrAa01ScC	$\text{CH}_3\text{COCH}_3 + \text{OH} \rightarrow \text{CH}_3\text{COCH}_2\text{O}_2$	1.80E8		Gligorovski et al. (2009)
A43007_a01	TrAa01ScCN	$\text{CH}_3\text{COCH}_3 + \text{NO}_3 \rightarrow \text{CH}_3\text{COCH}_2\text{O}_2 + \text{NO}_3^- + \text{H}^+$	3.721E+03	-4332	Herrmann and Zellner (1998)
A43008a_a01	TrAa01ScC	$\text{CH}_3\text{COCH}_2\text{O}_2 + \text{CH}_3\text{COCH}_2\text{O}_2 \rightarrow \text{CH}_3\text{COCH}_2\text{OH} + \text{CH}_3\text{C(O)CHO}$	$0.20 \times 4.00\text{E}8$		Zegota et al. (1986)
A43008b_a01	TrAa01ScC	$\text{CH}_3\text{COCH}_2\text{O}_2 + \text{CH}_3\text{COCH}_2\text{O}_2 \rightarrow 2.0 \text{ CH}_3\text{C(O)CHO} + \text{H}_2\text{O}_2$	0.45 × 4.00E8		Zegota et al. (1986)
A43009c_a01	TrAa01ScC	$\text{CH}_3\text{COCH}_2\text{O}_2 + \text{CH}_3\text{COCH}_2\text{O}_2 \rightarrow 2.0 \text{ HCHO} + 2.0 \text{ CH}_3\text{COOO}$	$0.15 \times 4.00\text{E}8$		Zegota et al. (1986)
A43009d_a01	TrAa01ScC	$\text{CH}_3\text{COCH}_2\text{O}_2 + \text{CH}_3\text{COCH}_2\text{O}_2 \rightarrow 2.0 \text{ CH}_3\text{C(O)CHO} + 2.0 \text{ HO}_2$	$0.20 \times 4.00\text{E}8$		Zegota et al. (1986)
A43010a_a01	TrAa01ScC	$\text{CH}_3\text{COCH}_2\text{OH} + \text{OH} \rightarrow \text{CH}_3\text{COCHOHO}_2$	$0.85 \times 5.10\text{E}8$		Doussin and Monod (2013)*
A43010b_a01	TrAa01ScC	$\text{CH}_3\text{COCH}_2\text{OH} + \text{OH} \rightarrow \text{HCHO} + \text{CH}_3\text{COOO}$	$0.15 \times 5.10\text{E}8$		Doussin and Monod (2013)
A43011_a01	TrAa01ScCN	$\text{CH}_3\text{COCH}_2\text{OH} + \text{NO}_3 \rightarrow \text{CH}_3\text{COCHOHO}_2 + \text{NO}_3^- + \text{H}^+$	2.108E+07	-1564	Gaillard de Sémainville et al. (2007)
A43012_a01	TrAa01ScC	$\text{CH}_3\text{COCHOHO}_2 \rightarrow \text{CH}_3\text{C(O)CHO} + \text{HO}_2$	1.90E2		von Sonntag (1987)
A43013_a01	TrAa01ScC	$\text{IPROPOL} + \text{OH} \rightarrow \text{CH}_3\text{COCH}_3 + \text{HO}_2$	1.90E9		see note*
A43014_a01	TrAa01ScCN	$\text{IPROPOL} + \text{NO}_3 \rightarrow \text{CH}_3\text{COCH}_3 + \text{NO}_3^- + \text{H}^+ + \text{HO}_2$	3.70E6		see note*
A43015a_a01	TrAa01ScC	$\text{CH}_3\text{COCH}_2\text{O}_2\text{H} + \text{OH} \rightarrow \text{CH}_3\text{C(O)CHO} + \text{OH}$	1.80E8		see note*
A43015b_a01	TrAa01ScC	$\text{CH}_3\text{COCH}_2\text{O}_2\text{H} + \text{OH} \rightarrow \text{CH}_3\text{COCH}_2\text{O}_2$	4.70E8		see note*
A43016_a01	TrAa01ScCN	$\text{CH}_3\text{COCH}_2\text{O}_2\text{H} + \text{NO}_3 \rightarrow \text{CH}_3\text{COCH}_2\text{O}_2 + \text{NO}_3^- + \text{H}^+$	4.50E6		see note*
A43017_a01	TrAa01ScC	$\text{CH}_3\text{COCH}_2\text{O}_2 + \text{HO}_2 \rightarrow \text{CH}_3\text{COCH}_2\text{O}_2\text{H}$	4.30E5		see note*
A43018_a01	TrAa01ScC	$\text{CH}_3\text{COCH}_2\text{O}_2 + \text{O}_2^- \rightarrow \text{CH}_3\text{COCH}_2\text{O}_2\text{H} + \text{O}_2 + \text{OH}^-$	5.00E7		see note*
A43019a_a01	TrAa01ScC	$\text{iC}_3\text{H}_7\text{OOH} + \text{OH} \rightarrow \text{CH}_3\text{COCH}_3 + \text{OH}$	9.90E8		see note*
A43019b_a01	TrAa01ScC	$\text{iC}_3\text{H}_7\text{OOH} + \text{OH} \rightarrow \text{iC}_3\text{H}_7\text{O}_2$	1.80E8		see note*
A43020_a01	TrAa01ScCN	$\text{iC}_3\text{H}_7\text{OOH} + \text{NO}_3 \rightarrow \text{iC}_3\text{H}_7\text{O}_2 + \text{NO}_3^- + \text{H}^+$	4.50E6		see note*
A43021_a01	TrAa01ScC	$\text{iC}_3\text{H}_7\text{O}_2 + \text{HO}_2 \rightarrow \text{iC}_3\text{H}_7\text{OOH}$	4.30E5		see note*

Table 6: Aqueous phase reactions (...continued)

#	labels	reaction	$k_0 [M^{1-n}s^{-1}]$	$-E_a/R[K]$	reference
A43022_a01	TrAa01ScC	$iC_3H_7O_2 + O_2^- \rightarrow iC_3H_7OOH + O_2 + OH^-$	5.00E7		see note*
A43023_a01	TrAa01ScC	$HOCH_2OH + MG2 \rightarrow MG3 + H_2O$	see note	see note	Hahnenstein et al. (1995)*
A43024_a01	TrAa01ScC	$MG3 + H_2O \rightarrow HOCH_2OH + MG2$	see note	see note	Hahnenstein et al. (1995)
A43025_a01	TrAa01ScC	$MG3 + OH \rightarrow HM2F + HO_2$	1.54E9	-1020	see note*
A44000_a01	TrAa01ScC	$MACR + OH \rightarrow CH_2OHCO_2CH_3CHO$	9.905E+09	-1203	Schöne and Herrmann (2014)
A44001_a01	TrAa01ScC	$CH_2OHCO_2CH_3CHO + CH_2OHCO_2CH_3CHO \rightarrow CH_3C(O)CHO + CH_3COCH_2OH + HOCH_2O_2 + HCOHOHO_2$	4.00E8		Schöne and Herrmann (2014)
A44002_a01	TrAa01ScC	$MVK + OH \rightarrow CH_3COCHO_2CH_2OH$	7.117E+09	-1443	Schöne and Herrmann (2014)
A44003_a01	TrAa01ScC	$CH_3COCHO_2CH_2OH + CH_3COCHO_2CH_2OH \rightarrow 1.1 CH_2COCOCH_2OH + .2 CH_2OHCHOOHCOCH_3 + .35 CH_2OHCHO + .35 CH_3C(O)CHO + .35 HOCH_2O_2 + .35 CH_3COOO + .45 H_2O_2$	4.00E8		Schöne and Herrmann (2014)
A44004_a01	TrAa01ScC	$GLYOX + CHOCHOHOH \rightarrow GOLIG1 + H_2O$	1.00E2		Ervens and Volkamer (2010)
A44005_a01	TrAa01ScC	$GOLIG1 + H_2O \rightarrow GLYOX + CHOCHOHOH$	1.00E-1		Ervens and Volkamer (2010)
A44006_a01	TrAa01ScC	$CHOCHOHOH + CHOCHOHOH \rightarrow GOLIG2 + H_2O$	1.00E2		Ervens and Volkamer (2010)
A44007_a01	TrAa01ScC	$GOLIG2 + H_2O \rightarrow CHOCHOHOH + CHOCHOHOH$	1.00E-1		Ervens and Volkamer (2010)
A44008_a01	TrAa01ScC	$CHOHOHCHOHOH + CHOCHOHOH \rightarrow GOLIG3 + H_2O$	1.00E2		Ervens and Volkamer (2010)
A44009_a01	TrAa01ScC	$GOLIG3 + H_2O \rightarrow CHOHOHCHOHOH + CHOCHOHOH$	1.00E-1		Ervens and Volkamer (2010)
A44010_a01	TrAa01ScC	$GOLIG1 + OH \rightarrow GOLIGO1 + HO_2$	1.610E+09	-1516	see note*
A44011_a01	TrAa01ScC	$GOLIG2 + OH \rightarrow GOLIGO2 + HO_2$	1.610E+09	-1516	see note*
A44012_a01	TrAa01ScC	$GOLIG3 + OH \rightarrow GOLIGO3 + HO_2$	1.610E+09	-1516	see note*
A46000_a01	TrAa01ScC	$CH_3C(O)CHO + CH_3COCHOHOH \rightarrow CH_3COCHOHOCHOHOCH_3 + H_2O$	1.00E2		Ervens and Volkamer (2010)*
A46001_a01	TrAa01ScC	$CH_3COCHOHOCHOHOCH_3 + H_2O \rightarrow CH_3COCHOHOH + CH_3C(O)CHO$	1.00E-1		Ervens and Volkamer (2010)*
A46002_a01	TrAa01ScC	$CH_3COCHOHOCHOHOCH_3 + OH \rightarrow CH_3COCOOHOHOCH_3 + HO_2$	1.303E+09	-1235	see note*
A46003_a01	TrAa01ScC	$CH_3COCHOHOH + CH_3COCHOHOH \rightarrow CH_3COCHOHOCHOHC_3CHOHOH + H_2O$	1.00E2		Ervens and Volkamer (2010)*

Table 6: Aqueous phase reactions (...continued)

#	labels	reaction	$k_0 [M^{1-n}s^{-1}]$	$-E_a/R[K]$	reference
A46004_a01	TrAa01ScC	$\text{CH}_3\text{COCHOHOCOHC}_3\text{CHOHOH} + \text{H}_2\text{O} \rightarrow 2 \text{CH}_3\text{COCHOHOH}$	1.00E-1		Ervens and Volkamer (2010)*
A46005_a01	TrAa01ScC	$\text{CH}_3\text{COCHOHOCOHC}_3\text{CHOHOH} + \text{OH} \rightarrow \text{CH}_3\text{COCOHOOCOHC}_3\text{COHOH} + \text{HO}_2$	1.303E+09	-1235	see note*
A460MS_a01	TrAa01CN	$\text{PHENOL} + \text{NO}_2^+ \rightarrow \text{HOC}_6\text{H}_4\text{NO}_2 + \text{H}^+$	7.5E9		Ryder et al. (2015), Heal et al. (2007)*
A471MS_a01	TrAa01CN	$\text{CRESOL} + \text{NO}_2^+ \rightarrow \text{TOL1OHNO}_2 + \text{H}^+$	7.5E9		Coombes et al. (1979)
A60000_a01	TrAa01Cl	$\text{Cl} + \text{Cl} \rightarrow \text{Cl}_2$	8.8E7		Wu et al. (1980)
A60001_a01	TrAa01Cl	$\text{Cl}_2^- + \text{Cl}_2 \rightarrow \text{Cl}_2 + 2 \text{Cl}^-$	3.5E9		Yu (2004)
A61000_a01	TrAa01Cl	$\text{Cl}^- + \text{O}_3 \rightarrow \text{ClO}^-$	3.0E-3		Hoigné et al. (1985)
A61001_a01	TrAa01Cl	$\text{Cl}_2 + \text{O}_2^- \rightarrow \text{Cl}_2^-$	1.0E9		Bjergbakke et al. (1981)
A61002_a01	TrAa01Cl	$\text{Cl}_2^- + \text{O}_2^- \rightarrow 2 \text{Cl}^-$	1.0E9		Jacobi (1996)*
A62000_a01	TrAa01Cl	$\text{Cl} \rightarrow \text{H}^+ + \text{ClOH}^-$	1.8E5		Yu (2004)
A62001_a01	TrAa01Cl	$\text{Cl} + \text{H}_2\text{O}_2 \rightarrow \text{HO}_2 + \text{Cl}^- + \text{H}^+$	2.7E7	-1684	Christensen et al. (1982)
A62002_a01	TrAa01Cl	$\text{Cl}^- + \text{OH} \rightarrow \text{ClOH}^-$	4.2E9		Yu (2004)
A62003_a01	TrAa01Cl	$\text{Cl}_2 + \text{HO}_2 \rightarrow \text{Cl}_2^- + \text{H}^+$	1.0E9		Bjergbakke et al. (1981)
A62004_a01	TrAa01MblCl	$\text{Cl}_2 \rightarrow \text{Cl}^- + \text{HOCl} + \text{H}^+$	21.8	-8012	Wang and Margerum (1994)
A62005_a01	TrAa01Cl	$\text{Cl}_2^- + \text{HO}_2 \rightarrow 2 \text{Cl}^- + \text{H}^+$	1.3E10		Jacobi (1996)
A62006_a01	TrAa01Cl	$\text{HOCl} + \text{O}_2^- \rightarrow \text{Cl} + \text{OH}^-$	7.5E6		Long and Bielski (1980)
A62007_a01	TrAa01Cl	$\text{HOCl} + \text{HO}_2 \rightarrow \text{Cl}$	7.5E6		Long and Bielski (1980)
A62008_a01	TrAa01MblCl	$\text{HOCl} + \text{Cl}^- + \text{H}^+ \rightarrow \text{Cl}_2$	2.2E4	-3508	Wang and Margerum (1994)
A62009_a01	TrAa01Cl	$\text{ClOH}^- \rightarrow \text{Cl}^- + \text{OH}$	6.0E9		Yu (2004)
A62010_a01	TrAa01Cl	$\text{ClOH}^- + \text{H}^+ \rightarrow \text{Cl}$	2.4E10		Yu (2004)
A620MS_a01	TrAa01Cl	$\text{ClOH}^- + \text{ClOH}^- \rightarrow \text{Cl}_2 + 2 \text{OH}^-$	1.8E9		Knipping et al. (2000)
A621MS_a01	TrAa01Cl	$\text{ClOH}^- + \text{Cl}^- \rightarrow \text{Cl}_2^- + \text{OH}^-$	1.0E4		Grigorev et al. (1987)
A622MS_a01	TrAa01Cl	$\text{OH}^- + \text{Cl}_2^- \rightarrow \text{ClOH}^- + \text{Cl}^-$	4.5E7		Grigorev et al. (1987)
A63000_a01	TrAa01ClN	$\text{Cl} + \text{NO}_3^- \rightarrow \text{NO}_3 + \text{Cl}^-$	1.0E8		Buxton et al. (1999b)
A63001_a01	TrAa01ClN	$\text{Cl}^- + \text{NO}_3 \rightarrow \text{NO}_3^- + \text{Cl}$	3.4E8		Buxton et al. (1999b)*
A63002_a01	TrAa01ClN	$\text{Cl}_2^- + \text{NO}_2^- \rightarrow 2 \text{Cl}^- + \text{NO}_2$	6.0E7		Jacobi et al. (1996)
A630MS_a01	TrAa01MblClN	$\text{ClNO}_2 \rightarrow \text{Cl}^- + \text{NO}_2^+$	2.70E2		Behnke et al. (1997)
A631MS_a01	TrAa01MblClN	$\text{Cl}^- + \text{NO}_2^+ \rightarrow \text{ClNO}_2$	7.5E9		Staudt et al. (2019)
A632MS_a01	TrAa01MblClN	$\text{ClNO}_2 + \text{Cl}^- \rightarrow \text{Cl}_2 + \text{NO}_2^-$	1.0E7		Roberts et al. (2008)
A64000_a01	TrAa01ScCl	$\text{Cl}_2^- + \text{CH}_3\text{OOH} \rightarrow 2 \text{Cl}^- + \text{H}^+ + \text{CH}_3\text{OO}$	6.20E5		see note*
A70000_a01	TrAa01Br	$\text{Br}_2^- + \text{Br}_2^- \rightarrow 2 \text{Br}^- + \text{Br}_2$	1.9E9		Ross et al. (1992)

Table 6: Aqueous phase reactions (...continued)

#	labels	reaction	$k_0 [M^{1-n}s^{-1}]$	$-E_a/R[K]$	reference
A71000_a01	TrAa01Br	$\text{Br}^- + \text{O}_3 \rightarrow \text{BrO}^-$	2.1E2	-4450	Haag and Hoigné (1983)
A71001_a01	TrAa01Br	$\text{Br}_2 + \text{O}_2^- \rightarrow \text{Br}_2^-$	5.6E9		Sutton and Downes (1972)
A71002_a01	TrAa01Br	$\text{Br}_2^- + \text{O}_2^- \rightarrow 2 \text{Br}^-$	1.7E8		Wagner and Strehlow (1987)
A72000_a01	TrAa01Br	$\text{Br}^- + \text{OH} \rightarrow \text{BrOH}^-$	1.1E10		Zehavi and Rabani (1972)
A72001_a01	TrAa01Br	$\text{Br}_2 + \text{HO}_2 \rightarrow \text{Br}_2^- + \text{H}^+$	1.1E8		Sutton and Downes (1972)
A72002_a01	TrAa01MblBr	$\text{Br}_2 \rightarrow \text{Br}^- + \text{HOBr} + \text{H}^+$	9.7E1	-7457	Beckwith et al. (1996)
A72003_a01	TrAa01Br	$\text{Br}_2^- + \text{HO}_2 \rightarrow \text{Br}_2 + \text{H}_2\text{O}_2 + \text{OH}^-$	4.4E9		Matthew et al. (2003)
A72004_a01	TrAa01Br	$\text{Br}_2^- + \text{H}_2\text{O}_2 \rightarrow 2 \text{Br}^- + \text{H}^+ + \text{HO}_2$	1.0E5		Jacobi (1996)
A72005_a01	TrAa01Br	$\text{HOBr} + \text{O}_2^- \rightarrow \text{Br} + \text{OH}^-$	3.5E9		Schwarz and Bielski (1986)
A72006_a01	TrAa01Br	$\text{HOBr} + \text{HO}_2 \rightarrow \text{Br}$	1.0E9		Herrmann et al. (1999a)
A72007_a01	TrAa01Br	$\text{HOBr} + \text{H}_2\text{O}_2 \rightarrow \text{Br}^- + \text{H}^+$	1.2E6		Bichsel and von Gunten (1999)
A72008_a01	TrAa01MblBr	$\text{HOBr} + \text{Br}^- + \text{H}^+ \rightarrow \text{Br}_2$	1.6E10		Beckwith et al. (1996)
A72009a_a01	TrAa01Br	$\text{BrOH}^- \rightarrow \text{Br}^- + \text{OH}$	3.3E7		Zehavi and Rabani (1972)
A72009b_a01	TrAa01Br	$\text{BrOH}^- \rightarrow \text{Br} + \text{OH}^-$	4.2E6		Zehavi and Rabani (1972)
A72010_a01	TrAa01Br	$\text{BrOH}^- + \text{H}^+ \rightarrow \text{Br}$	4.4E10		Zehavi and Rabani (1972)
A73000_a01	TrAa01BrN	$\text{Br}^- + \text{NO}_3 \rightarrow \text{Br} + \text{NO}_3^-$	4.0E9		Neta and Huie (1986)
A73001_a01	TrAa01BrN	$\text{Br}_2^- + \text{NO}_2^- \rightarrow 2 \text{Br}^- + \text{NO}_2$	1.7E7	-1720	Shoute et al. (1991)
A74000_a01	TrAa01Br	$\text{Br}_2^- + \text{CH}_3\text{OOH} \rightarrow 2 \text{Br}^- + \text{H}^+ + \text{CH}_3\text{OO}$	1.0E5		Jacobi (1996)*
A76001_a01	TrAa01BrCl	$\text{Br}^- + \text{ClO}^- + \text{H}^+ \rightarrow \text{BrCl} + \text{OH}^-$	3.7E10		Kumar and Margerum (1987)
A76002_a01	TrAa01MblBrCl	$\text{Br}^- + \text{HOCl} + \text{H}^+ \rightarrow \text{BrCl}$	1.32E6		Kumar and Margerum (1987)
A76003_a01	TrAa01MblBrCl	$\text{HOBr} + \text{Cl}^- + \text{H}^+ \rightarrow \text{BrCl}$	2.3E10		Liu and Margerum (2001)*
A76004_a01	TrAa01MblBrCl	$\text{BrCl} \rightarrow \text{Cl}^- + \text{HOBr} + \text{H}^+$	3.0E6		Liu and Margerum (2001)
A91000_a01	TrAa01ScS	$\text{SO}_3^- + \text{O}_2 \rightarrow \text{SO}_5^-$	1.5E9		Huie and Neta (1987)
A91001_a01	TrAa01MblScScmS	$\text{SO}_3^{2-} + \text{O}_3 \rightarrow \text{SO}_4^{2-}$	1.5E9	-5300	Hoffmann (1986)
A91002_a01	TrAa01ScS	$\text{SO}_4^- + \text{O}_2^- \rightarrow \text{SO}_4^{2-}$	3.5E9		Jiang et al. (1992)
A91003_a01	TrAa01ScS	$\text{SO}_4^- + \text{SO}_3^{2-} \rightarrow \text{SO}_3^- + \text{SO}_4^{2-}$	4.6E8		Huie and Neta (1987)
A91004_a01	TrAa01ScS	$\text{SO}_5^- + \text{O}_2^- \rightarrow \text{HSO}_5^- + \text{OH}^-$	2.3E8		Buxton et al. (1996)
A91005_a01	TrAa01S	$\text{SO}_5^- + \text{SO}_3^{2-} \rightarrow .72 \text{SO}_3^- + .72 \text{SO}_4^{2-} + .28 \text{SO}_3^- + .28 \text{HSO}_5^- + .28 \text{OH}^-$	1.3E7		Huie and Neta (1987), Deister and Warneck (1990)*
A91006_a01	TrAa01S	$\text{SO}_5^- + \text{SO}_5^- \rightarrow \text{O}_2 + \text{SO}_4^{2-} + \text{LSULFUR}$	1.0E8		Ross et al. (1992)*
A92000_a01	TrAa01ScS	$\text{SO}_3^{2-} + \text{OH} \rightarrow \text{SO}_3^- + \text{OH}^-$	5.5E9		Buxton et al. (1988)
A92001_a01	TrAa01ScS	$\text{SO}_4^- + \text{OH} \rightarrow \text{HSO}_5^-$	1.0E9		Jiang et al. (1992)
A92002_a01	TrAa01ScS	$\text{SO}_4^- + \text{HO}_2 \rightarrow \text{SO}_4^{2-} + \text{H}^+$	3.5E9		Jiang et al. (1992)
A92003_a01	TrAa01ScS	$\text{SO}_4^- + \text{H}_2\text{O} \rightarrow \text{SO}_4^{2-} + \text{H}^+ + \text{OH}$	1.1E1	-1110	Herrmann et al. (1995)

Table 6: Aqueous phase reactions (...continued)

#	labels	reaction	$k_0 [M^{1-n}s^{-1}]$	$-E_a/R[K]$	reference
A92004_a01	TrAa01ScS	$\text{SO}_4^- + \text{H}_2\text{O}_2 \rightarrow \text{SO}_4^{2-} + \text{H}^+ + \text{HO}_2$	1.2E7		Wine et al. (1989)
A92005_a01	TrAa01ScS	$\text{HSO}_3^- + \text{O}_2^- \rightarrow \text{SO}_4^{2-} + \text{OH}$	3.0E3		see note*
A92006_a01	TrAa01MblScScmS	$\text{HSO}_3^- + \text{O}_3 \rightarrow \text{SO}_4^{2-} + \text{H}^+$	3.7E5	-5500	Hoffmann (1986)
A92007_a01	TrAa01ScS	$\text{HSO}_3^- + \text{OH} \rightarrow \text{SO}_3^-$	4.5E9		Buxton et al. (1988)
A92008_a01	TrAa01ScS	$\text{HSO}_3^- + \text{HO}_2 \rightarrow \text{SO}_4^{2-} + \text{OH} + \text{H}^+$	3.0E3		see note*
A92009_a01	TrAa01MblScScmS	$\text{HSO}_3^- + \text{H}_2\text{O}_2 \rightarrow \text{SO}_4^{2-} + \text{H}^+$	5.2E6	-3650	Martin and Damschen (1981)
A92010_a01	TrAa01ScS	$\text{HSO}_3^- + \text{SO}_4^- \rightarrow \text{SO}_3^- + \text{SO}_4^{2-} + \text{H}^+$	8.0E8		Huie and Neta (1987)
A92011_a01	TrAa01S	$\text{HSO}_3^- + \text{SO}_5^- \rightarrow .75 \text{SO}_4^- + .75 \text{SO}_4^{2-} + .75 \text{H}^+ + .25 \text{SO}_3^- + .25 \text{HSO}_5^-$	1.0E5		Huie and Neta (1987)
A92012_a01	TrAa01ScS	$\text{HSO}_3^- + \text{HSO}_5^- + \text{H}^+ \rightarrow 2 \text{HSO}_4^- + \text{H}^+$	7.1E6		Betterton and Hoffmann (1988)
A93001_a01	TrAa01ScNS	$\text{SO}_4^- + \text{NO}_3^- \rightarrow \text{SO}_4^{2-} + \text{NO}_3$	5.0E4		Exner et al. (1992)
A93002_a01	TrAa01ScNS	$\text{SO}_4^{2-} + \text{NO}_3 \rightarrow \text{NO}_3^- + \text{SO}_4^-$	1.0E5		Løgager et al. (1993)
A93004_a01	TrAa01ScNS	$\text{HSO}_3^- + \text{NO}_3 \rightarrow \text{SO}_3^- + \text{NO}_3^- + \text{H}^+$	1.4E9	-2000	Exner et al. (1992)
A93005_a01	TrAa01ScNS	$\text{HSO}_3^- + \text{HNO}_4 \rightarrow \text{HSO}_4^- + \text{NO}_3^- + \text{H}^+$	3.1E5		Warneck (1999)
A930MS_a01	TrAa01ScNS	$\text{SO}_4^{2-} + \text{NO}_2^+ \rightarrow \text{SO}_4^- + \text{NO}_3^- + 2 \text{H}^+$	7.5E9		Staudt et al. (2019)*
A94100_a01	TrAa01ScS	$\text{SO}_3^{2-} + \text{HCHO} \rightarrow \text{CH}_2\text{OHSO}_3^- + \text{OH}^-$	1.4E4		Boyce and Hoffmann (1984)*
A94101_a01	TrAa01ScS	$\text{SO}_3^{2-} + \text{CH}_3\text{OOH} + \text{H}^+ \rightarrow \text{SO}_4^{2-} + \text{H}^+ + \text{CH}_3\text{OH}$	1.6E7	-3800	Lind et al. (1987)
A94102_a01	TrAa01ScS	$\text{HSO}_3^- + \text{HCHO} \rightarrow \text{CH}_2\text{OHSO}_3^-$	4.3E-1		Boyce and Hoffmann (1984)*
A94103_a01	TrAa01ScS	$\text{HSO}_3^- + \text{CH}_3\text{OOH} + \text{H}^+ \rightarrow \text{HSO}_4^- + \text{H}^+ + \text{CH}_3\text{OH}$	1.6E7	-3800	Lind et al. (1987)
A94104_a01	TrAa01ScS	$\text{HSO}_3^- + \text{CH}_3\text{OO} \rightarrow \text{SO}_3^- + \text{CH}_3\text{OOH}$	5.00E5		Herrmann et al. (1999b)
A94105_a01	TrAa01ScS	$\text{SO}_4^- + \text{HCOO}^- \rightarrow \text{SO}_4^{2-} + \text{CO}_2 + \text{HO}_2$	1.7E8	-1500	Jacob (1986)
A94106_a01	TrAa01ScS	$\text{SO}_4^- + \text{HCOOH} \rightarrow \text{SO}_4^{2-} + \text{CO}_2 + \text{H}^+ + \text{HO}_2$	1.7E8	-1500	Jacob (1986)
A94107_a01	TrAa01ScS	$\text{SO}_4^- + \text{CH}_3\text{OH} \rightarrow \text{SO}_4^{2-} + \text{HOCH}_2\text{O}_2 + \text{H}^+$	9.039E+06	-2190	Clifton and Huie (1989)
A94108a_a01	TrAa01ScS	$\text{SO}_4^- + \text{CH}_3\text{OOH} \rightarrow \text{SO}_4^{2-} + \text{CH}_3\text{OO} + \text{H}^+$	$0.25 \times 1.20\text{E}7$		see note*
A94108b_a01	TrAa01ScS	$\text{SO}_4^- + \text{CH}_3\text{OOH} \rightarrow \text{SO}_4^{2-} + \text{HCHO} + \text{HO}_2 + \text{H}^+$	$0.75 \times 1.20\text{E}7$		see note*
A94109_a01	TrAa01ScS	$\text{SO}_4^- + \text{HOCH}_2\text{OH} \rightarrow \text{SO}_4^{2-} + \text{HCOHOHO}_2 + \text{H}^+$	1.40E7	-1300	Buxton et al. (1990)
A94110_a01	TrAa01ScS	$\text{SO}_5^- + \text{HCOO}^- \rightarrow \text{HSO}_5^- + \text{CO}_2 + \text{O}_2^-$	1.4E4	-4000	Jacob (1986)
A94111_a01	TrAa01ScS	$\text{CH}_2\text{OHSO}_3^- + \text{OH}^- \rightarrow \text{SO}_3^{2-} + \text{HCHO}$	3.6E3		Seinfeld and Pandis (1998)
A94200_a01	TrAa01ScCS	$\text{HSO}_3^- + \text{CH}_2\text{OOCOOH} \rightarrow \text{SO}_3^- + \text{HOOCH}_2\text{CO}_2\text{H}$	5.00E5		see note*
A94201_a01	TrAa01ScCS	$\text{HSO}_3^- + \text{CH}_2\text{OOCO}_2^- \rightarrow \text{SO}_3^- + \text{CH}_2\text{OOHC}_2^-$	5.00E5		see note*
A94202a_a01	TrAa01ScCS	$\text{SO}_4^- + \text{CH}_3\text{CH}_2\text{OH} \rightarrow \text{SO}_4^{2-} + \text{CH}_3\text{CHO} + \text{HO}_2 + \text{H}^+$	$0.90 \times 4.236\text{E}+07$	-1750	Clifton and Huie (1989)*
A94202b_a01	TrAa01ScCS	$\text{SO}_4^- + \text{CH}_3\text{CH}_2\text{OH} \rightarrow \text{SO}_4^{2-} + \text{CH}_2\text{OHCH}_2\text{OO} + \text{H}^+$	$0.10 \times 4.236\text{E}+07$	-1750	Clifton and Huie (1989)

Table 6: Aqueous phase reactions (...continued)

#	labels	reaction	$k_0 [M^{1-n}s^{-1}]$	$-E_a/R[K]$	reference
A94203a_a01	TrAa01ScCS	$\text{SO}_4^- + \text{CHOHOHCHOHOH} \rightarrow \text{SO}_4^{2-} + \text{CHOHOHCOHOHO}_2 + \text{H}^+$	$0.27 \times 2.40\text{E}7$		George et al. (2001)
A94203b_a01	TrAa01ScCS	$\text{SO}_4^- + \text{CHOHOHCHOHOH} \rightarrow \text{SO}_4^{2-} + \text{HCOHOHO}_2 + \text{HCOOH} + \text{HO}_2 + \text{H}^+$	$0.73 \times 2.40\text{E}7$		George et al. (2001)*
A94204_a01	TrAa01ScCS	$\text{SO}_4^- + \text{CH}_3\text{COO}^- \rightarrow \text{SO}_4^{2-} + \text{CH}_2\text{OOOCO}_2^- + \text{H}^+$	5.10E6		Huie and Clifton (1990)
A94205_a01	TrAa01ScCS	$\text{SO}_4^- + \text{HOOCCOO}^- \rightarrow \text{SO}_4^{2-} + \text{C}_2\text{O}_4^- + \text{H}^+$	1.70E6		Grgić et al. (2007)
A94206_a01	TrAa01ScCS	$\text{SO}_4^- + \text{C}_2\text{O}_4^{2-} \rightarrow \text{SO}_4^{2-} + \text{C}_2\text{O}_4^-$	1.30E7		Grgić et al. (2007)
A96000_a01	TrAa01ClS	$\text{SO}_3^{2-} + \text{Cl}_2^- \rightarrow \text{SO}_3^- + 2 \text{Cl}^-$	6.2E7		Jacobi et al. (1996)
A96001_a01	TrAa01MblClS	$\text{SO}_3^{2-} + \text{HOCl} \rightarrow \text{Cl}^- + \text{HSO}_4^-$	7.6E8		Fogelman et al. (1989)
A96002_a01	TrAa01ClS	$\text{SO}_4^- + \text{Cl}^- \rightarrow \text{SO}_4^{2-} + \text{Cl}$	2.5E8		Buxton et al. (1999a)
A96003_a01	TrAa01ClS	$\text{SO}_4^{2-} + \text{Cl} \rightarrow \text{SO}_4^- + \text{Cl}^-$	2.1E8		Buxton et al. (1999a)
A96004_a01	TrAa01ClS	$\text{HSO}_3^- + \text{Cl}_2^- \rightarrow \text{SO}_3^- + 2 \text{Cl}^- + \text{H}^+$	4.7E8	-1082	Shoute et al. (1991)
A96005_a01	TrAa01MblClS	$\text{HSO}_3^- + \text{HOCl} \rightarrow \text{Cl}^- + \text{HSO}_4^- + \text{H}^+$	7.6E8		see note*
A96006_a01	TrAa01ClS	$\text{HSO}_5^- + \text{Cl}^- \rightarrow \text{HOCl} + \text{SO}_4^{2-}$	1.8E-3	-7352	Fortnum et al. (1960)
A97000_a01	TrAa01BrS	$\text{SO}_3^{2-} + \text{Br}_2^- \rightarrow 2 \text{Br}^- + \text{SO}_3^-$	2.2E8	-649	Shoute et al. (1991)
A97001_a01	TrAa01BrS	$\text{SO}_3^{2-} + \text{BrO}^- \rightarrow \text{Br}^- + \text{SO}_4^{2-}$	1.0E8		Troy and Margerum (1991)
A97002_a01	TrAa01MblBrS	$\text{SO}_3^- + \text{HOBr} \rightarrow \text{Br}^- + \text{HSO}_4^-$	5.0E9		Troy and Margerum (1991)
A97003_a01	TrAa01BrS	$\text{SO}_4^- + \text{Br}^- \rightarrow \text{Br} + \text{SO}_4^{2-}$	2.1E9		Jacobi (1996)
A97004_a01	TrAa01BrS	$\text{HSO}_3^- + \text{Br}_2^- \rightarrow 2 \text{Br}^- + \text{H}^+ + \text{SO}_3^-$	6.3E7	-782	Shoute et al. (1991)
A97005_a01	TrAa01MblBrS	$\text{HSO}_3^- + \text{HOBr} \rightarrow \text{Br}^- + \text{HSO}_4^- + \text{H}^+$	5.0E9		see note*
A97006_a01	TrAa01BrS	$\text{HSO}_5^- + \text{Br}^- \rightarrow \text{HOBr} + \text{SO}_4^{2-}$	1.0E0	-5338	Fogelman et al. (1989)
A111001_a01	TrAa01Fe	$\text{Fe}^{2+} + \text{O}_2^- \rightarrow \text{Fe}^{3+} + \text{HO}_2^- + \text{OH}^-$	1E7		de Laat and Le (2006)
A111002_a01	TrAa01Fe	$\text{Fe}^{3+} + \text{O}_2^- \rightarrow \text{O}_2 + \text{Fe}^{2+}$	5E7		de Laat and Le (2006)
A111003_a01	TrAa01Fe	$\text{Fe}^{2+} + \text{O}_3 \rightarrow \text{FeO}^{2+} + \text{O}_2$	8.2E5		Løgager et al. (1992)
A112001a_a01	TrAa01Fe	$\text{Fe}^{2+} + \text{OH} \rightarrow \text{Fe}^{3+} + \text{OH}^-$	2.7E8		de Laat and Le (2006)
A112001b_a01	TrAa01Fe	$\text{FeOH}^+ + \text{OH} \rightarrow \text{Fe}^{3+} + 2 \text{OH}^-$	2.7E8		de Laat and Le (2006)
A112002a_a01	TrAa01Fe	$\text{Fe}^{2+} + \text{H}_2\text{O}_2 \rightarrow \text{Fe}^{3+} + \text{OH} + \text{OH}^-$	5.5E1		de Laat and Le (2006)
A112002b_a01	TrAa01Fe	$\text{FeOH}^+ + \text{H}_2\text{O}_2 \rightarrow \text{Fe}^{3+} + \text{OH} + 2 \text{OH}^-$	5.9E6		de Laat and Le (2006)
A112003_a01	TrAa01Fe	$\text{FeHO}_2^{2+} \rightarrow \text{Fe}^{2+} + \text{HO}_2$	2.3E-3		de Laat and Le (2006)
A112004_a01	TrAa01Fe	$\text{Fe(OH)(HO}_2)^+ \rightarrow \text{Fe}^{2+} + \text{HO}_2 + \text{OH}^-$	2.3E-3		de Laat and Le (2006)
A112006_a01	TrAa01Fe	$\text{Fe}^{2+} + \text{HO}_2 \rightarrow \text{Fe}^{3+} + \text{HO}_2^-$	1.2E6		de Laat and Le (2006)
A112008a_a01	TrAa01Fe	$\text{FeOH}^{2+} + \text{O}_2^- \rightarrow \text{Fe}^{2+} + \text{O}_2 + \text{OH}^-$	1.5E8		Rush and Bielski (1985)
A112008b_a01	TrAa01Fe	$\text{Fe(OH)}_2^+ + \text{O}_2^- \rightarrow \text{Fe}^{2+} + \text{O}_2 + 2 \text{OH}^-$	1.5E8		Rush and Bielski (1985)
A112009_a01	TrAa01Fe	$\text{Fe}^{2+} + \text{O}_2^- \rightarrow \text{Fe}^{3+} + \text{H}_2\text{O}_2 + 2 \text{OH}^-$	1.0E7		Rush and Bielski (1985)

Table 6: Aqueous phase reactions (...continued)

#	labels	reaction	$k_0 [M^{1-n}s^{-1}]$	$-E_a/R[K]$	reference
A112010_a01	TrAa01Fe	$Fe^{2+} + OH \rightarrow FeOH^{2+}$	4.3E8		Christensen and Sehested (1981)
A112011_a01	TrAa01Fe	$FeO^{2+} + H_2O_2 \rightarrow Fe^{3+} + HO_2 + OH^-$	9.5E3		L��gager et al. (1992)
A112012_a01	TrAa01Fe	$FeO^{2+} \rightarrow Fe^{3+} + OH + OH^-$	1.3E-2		L��gager et al. (1992)
A112013_a01	TrAa01Fe	$FeO^{2+} + HO_2 \rightarrow Fe^{3+} + O_2 + OH^-$	2.0E6		L��gager et al. (1992)
A112014_a01	TrAa01Fe	$FeO^{2+} + OH \rightarrow Fe^{3+} + HO^{2-}$	1.0E7		L��gager et al. (1992)
A112015_a01	TrAa01Fe	$FeO^{2+} + Fe^{2+} \rightarrow 2 Fe^{3+} + 2 OH^-$	1.4E5		L��gager et al. (1992)
A112016_a01	TrAa01Fe	$FeO^{2+} + Fe^{2+} \rightarrow Fe(OH)_2Fe^{4+}$	1.8E4		Jacobsen et al. (1997)
A112017_a01	TrAa01Fe	$Fe(OH)_2Fe^{4+} + H^+ \rightarrow 2 Fe^{3+} + OH^-$	2.0		Jacobsen et al. (1997)
A112018_a01	TrAa01Fe	$Fe(OH)_2Fe^{4+} \rightarrow 2 Fe^{3+} + 2 OH^-$	0.49		Jacobsen et al. (1997)
A113001_a01	TrAa01FeN	$FeO^{2+} + HONO \rightarrow Fe^{3+} + NO_2 + OH^-$	1.1E4		Jacobsen et al. (1998)
A113002_a01	TrAa01FeN	$Fe^{2+} + NO_3 \rightarrow Fe^{3+} + NO_3^-$	8.0E6		Herrmann et al. (2000)
A116001_a01	TrAa01ClFe	$Fe^{2+} + Cl \rightarrow Fe^{3+} + Cl^-$	5.9E9		Jayson et al. (1973)
A116002a_a01	TrAa01ClFe	$Fe^{2+} + Cl^- \rightarrow Fe^{3+} + 2 Cl^-$	1E7		Thornton and Laurence (1973)
A116002b_a01	TrAa01ClFe	$Fe^{2+} + Cl_2^- \rightarrow FeCl^{2+} + Cl^-$	4E6		Thornton and Laurence (1973)
A116003a_a01	TrAa01ClFe	$FeCl^+ + HO_2 \rightarrow Fe^{3+} + Cl^- + HO^{2-}$	1.2E6		de Laat and Le (2006)
A116003b_a01	TrAa01ClFe	$FeCl^+ + O_2^- \rightarrow Fe^{3+} + Cl^- + HO^{2-} + OH^-$	1E7		de Laat and Le (2006)
A116004a_a01	TrAa01ClFe	$FeCl^{2+} + HO_2 \rightarrow Fe^{2+} + Cl^- + O_2 + H^+$	2E4		de Laat and Le (2006)
A116004b_a01	TrAa01ClFe	$FeCl_2^+ + HO_2 \rightarrow Fe^{2+} + 2 Cl^- + O_2 + H^+$	2E4		de Laat and Le (2006)
A116004c_a01	TrAa01ClFe	$FeCl^{2+} + O_2^- \rightarrow Fe^{2+} + Cl^- + O_2$	5E7		de Laat and Le (2006)
A116004d_a01	TrAa01ClFe	$FeCl_2^+ + O_2^- \rightarrow Fe^{2+} + 2 Cl^- + O_2$	5E7		de Laat and Le (2006)
A116005_a01	TrAa01ClFe	$FeO^{2+} + Cl^- \rightarrow Fe^{3+} + Cl + 2 OH^-$	1E2		Jacobsen et al. (1998)*
A117001_a01	TrAa01BrFe	$Fe^{2+} + Br_2^- \rightarrow Fe^{3+} + 2 Br^-$	3.6E6		Thornton and Laurence (1973)
A119001_a01	TrAa01FeS	$FeO^{2+} + SO_2 \rightarrow Fe^{3+} + SO_3^-$	4.5E5		Jacobsen et al. (1998)*
A119002_a01	TrAa01FeS	$FeO^{2+} + HSO_3^- \rightarrow Fe^{3+} + SO_3^- + OH^-$	2.5E5		Jacobsen et al. (1998)*
A119003_a01	TrAa01FeS	$FeOH^{2+} + HSO_3^- \rightarrow Fe^{2+} + SO_3^- + H_2O$	30		Ziajka et al. (1994)
A119004_a01	TrAa01FeS	$Fe^{2+} + SO_5^- \rightarrow FeOH^{2+} + HSO_5^-$	8E5		Ziajka et al. (1994)*
A119005_a01	TrAa01FeS	$Fe^{2+} + HSO_5^- \rightarrow FeOH^{2+} + SO_4^-$	3.0E4		Gilbert and Stell (1990)
A119006_a01	TrAa01FeS	$Fe^{2+} + SO_4^- \rightarrow FeSO_4^+$	3.6E7		McElroy and Waygood (1990)*
A119007_a01	TrAa01FeS	$FeOH^{2+} + SO_3^- \rightarrow Fe^{2+} + HSO_4^-$	3E7		Warneck (2018)
A119008_a01	TrAa01FeS	$FeSO_3^+ + SO_3^- \rightarrow Fe^{2+} + SO_4^{2-} + SO_2$	2.16E6		Warneck (2018)*

Specific notes

Staudt et al. (2019).

(1985).

A311MS_a01: Rate constant derived from $k_3/k_2b = 6.E-2$ in Bertram and Thornton (2009) and $k_3 = 1.7E5$ inA321MS_a01: Rate constant is derived from $k(NO_2p + H_2O)/K(NO_2p)$, where $K(NO_2p)$ is 1E16 Sampoli et al.

A41013a_a01: Branching ratios taken from Asmus et al. (1973)

A41016a_a01: Branching ratio explaining the HCOOH yield by Monod et al. (2007) who originally assigned it to the channel for the methylic H-abstraction. However, Monod et al. (2007), differently from Herrmann et al. (1999b), assumed that the self-reaction of CH_3O_2 would only produce 2 CH_3O radicals and thus $\text{HCHO} + \text{HO}_2$. Instead, the latter reaction has a 0.8 yield of HOCH_2O_2 , which is a precursor of hydroxymethyl hydroperoxide and thus HCOOH.

A41016b_a01: The CH_2OOH radical has a lifetime of 10^{-9} s in the gas phase decomposing to HCHO and OH . O_2 -addition in the aqueous-phase seems unlikely. It is hard to imagine how the HOOCH_2O_2 radical would decompose into HCOOH and HO_2 .

A41017a_a01: $k(\text{H}_2\text{O}_2+\text{NO}_3)$, branching ratio as for $\text{CH}_3\text{OOH} + \text{OH}$

A41017b_a01: See branch a.

A41018a_a01: $k(\text{H}_2\text{O}_2+\text{CO}_3^-)$, branching ratio as for $\text{CH}_3\text{OOH} + \text{OH}$

A41018b_a01: See branch a.

A41019a_a01: Branching ratio as for $\text{CH}_3\text{OOH} + \text{OH}$

A41019b_a01: HOCHOHO_2 is assumed to directly decompose into CHOOOH and HO_2

A41020a_a01: $k(\text{H}_2\text{O}_2+\text{NO}_3)$, branching ratio as for $\text{CH}_3\text{OOH} + \text{OH}$

A41020b_a01: HOCHOHO_2 is assumed to directly decompose into CHOOOH and HO_2

A41021_a01: HO_2 elimination

A41022_a01: $k(\text{HO}_2+\text{HO}_2)$

A41023_a01: $k(\text{HO}_2+\text{O}_2^-)$

A41030_a01: HO_2 elimination

A42000a_a01: $\text{CH}_3\text{CHOHO}_2$ is assumed to directly decompose into $\text{CH}_3\text{CHO} + \text{HO}_2$

A42001a_a01: $\text{CH}_3\text{CHOHO}_2$ is assumed to directly decompose into $\text{CH}_3\text{CHO} + \text{HO}_2$

A42003_a01: $k(\text{HO}_2+\text{O}_2^-)$

A42004_a01: $k(\text{HO}_2+\text{HO}_2)$

A42005_a01: k approximated from $(k(\text{CH}_3\text{OOH}+\text{OH}) / k(\text{CH}_3\text{OH}+\text{OH}))$

A42006_a01: $\text{CH}_2\text{OHCHOHO}_2$ is assumed to directly decompose into $\text{HOCH}_2\text{CHO} + \text{HO}_2$

A42007_a01: $\text{CH}_2\text{OHCHOHO}_2$ is assumed to directly decompose into $\text{HOCH}_2\text{CHO} + \text{HO}_2$

A42010_a01: k based on Monod et al. (2005): $k=k(2 \text{CH}_3\text{CH}_2(\text{OO}))$

A42014_a01: HO_2 elimination

A42016a_a01: k assumed to be the same as for $\text{CH}_3\text{CHO} + \text{NO}_3$

A42016b_a01: See branch a.

A42017_a01: $k(\text{CH}_3\text{CHOHOH}+\text{O}_2^-)$

A42018_a01: HO_2 elimination

A42019_a01: k based on von Sonntag (1987)

A42020_a01: k based on von Sonntag (1987)

A42022a_a01: $k(\text{CH}_3\text{CHOHOH}+\text{NO}_3)$

A42022b_a01: See branch a.

A42022c_a01: See branch a.

A42023b_a01: CHOHOHO_2 directly decomposes into $\text{HCOOH} + \text{HO}_2$

A42024a_a01: k based on Neta and Huie (1986)

A42024b_a01: CHOHOHO_2 directly decomposes into $\text{HCOOH} + \text{HO}_2$

A42025_a01: HO_2 elimination

A42030b_a01: $\text{CH}_3\text{CHOHO}_2$ is assumed to directly decompose into $\text{CH}_3\text{CO}_2\text{H}$ and HO_2

A42031b_a01: $\text{CH}_3\text{CHOHO}_2$ is assumed to directly decompose into $\text{CH}_3\text{CHO} + \text{HO}_2$

A42032_a01: $k(\text{HO}_2+\text{O}_2^-)$

A42033_a01: $k(\text{HO}_2+\text{HO}_2)$

A42034_a01: $k(\text{HO}_2+\text{HO}_2)$

A42035_a01: $k(\text{HO}_2+\text{O}_2^-)$

A42037_a01: $k(\text{HO}_2+\text{HO}_2)$

A42038_a01: $k(\text{HO}_2+\text{O}_2^-)$

A42144a_a01: k assumed to be the same as $\text{C}_2\text{H}_5\text{OOH} + \text{OH}$ based on Monod et al. (2007)

A42144b_a01: See branch a.

A42146a_a01: k assumed to be the same as $\text{C}_2\text{H}_5\text{OOH} + \text{OH}$ based on Monod et al. (2007)

A42146b_a01: See branch a.

A42148_a01: HO_2 elimination

A42149_a01: HO_2 elimination

A42150a_a01: COOHOO is not formed but directly dissociates into $\text{CO}_2 + \text{HO}_2$. Rate coefficient based on Buxton et al. (1988)

A42150b_a01: See branch a.

A42151a_a01: COOHOO is not formed but directly dissociates into $\text{CO}_2 + \text{HO}_2$. Rate coefficient based on Gaillard de Sémainville et al. (2007)

A42151b_a01: See branch a.

A42156b_a01: COOHOO is not formed but directly dissociates into $\text{CO}_2 + \text{HO}_2$.

A42157a_a01: $k(\text{CHOHOHCHOHOH}+\text{NO}_3)$

A42157b_a01: COOHOO is not formed but directly dissociates into $\text{CO}_2 + \text{HO}_2$

A42161_a01: HO_2 elimination

A42162_a01: HO_2 elimination

A42163a_a01: $k(\text{HOCH}_2\text{CHO} + \text{OH})$

A42163b_a01: See branch a.

- A42164a_a01: $k(\text{HOCH}_2\text{CHO} + \text{NO}_3)$
- A42164b_a01: See branch a.
- A42165a_a01: $k(\text{HOCH}_2\text{CHOHOH} + \text{OH})$
- A42165b_a01: See branch a.
- A42165c_a01: See branch a.
- A42166a_a01: $k(\text{HOCH}_2\text{CHOHOH} + \text{NO}_3)$
- A42166b_a01: See branch a.
- A42166c_a01: See branch a.
- A42167_a01: pH-dependent
- A42169_a01: $k = 2 \times k(\text{HOCH}_2\text{OH} + \text{OH})$
- A42471_a01: Assumed to be the same as $\text{CH}_3\text{CO}_3 + \text{H}_2\text{O}$, following Villalta et al. (1996)
- A42472_a01: Assumed to be the same as $\text{CH}_3\text{CO}_3 + \text{H}_2\text{O}$, following Villalta et al. (1996)
- A42473_a01: Assumed to be the same as $\text{CH}_3\text{CO}_3 + \text{H}_2\text{O}$, following Villalta et al. (1996)
- A43000a_a01: Intermediate reaction with O_2^- and $\text{CH}(\text{OH})_2\text{COCH}_2\text{O}_2$ neglected
- A43001_a01: $\text{CH}(\text{OH})_2\text{COCH}_2\text{O}_2$ neglected
- A43002_a01: CO_2 added for mass balance intermediate reactions neglected
- A43004_a01: CO_2 added for mass balance intermediate reactions neglected
- A43010a_a01: $\text{CH}_2(\text{OH})\text{COCH}_2\text{O}_2$ was neglected with a branching ratio 0.16 added to $\text{CH}_3\text{COCHOHO}_2$
- A43013_a01: There is an intermediate reaction with branching ratio 0.87 and 0.13, the minor compound is neglected (Monod et al., 2005)
- A43014_a01: There is an intermediate reaction with branching ratio 0.87 and 0.13, the minor compound is neglected (Herrmann et al., 1994)
- A43015a_a01: k calculated comparing the rates $(\text{CH}_3\text{OH} + \text{OH}/\text{CH}_3\text{OOH} + \text{OH})$ and $(\text{ACETOL} + \text{OH}/\text{HYPERACET} + \text{OH})$
- A43015b_a01: k from $\text{CH}_3\text{OOH} + \text{OH} \rightarrow \text{HCHO}$.
- A43016_a01: k taken from the reaction of the hydrated form of MGLYOX and NO_3^-
- A43017_a01: k from $\text{CH}_3\text{O}_2 + \text{HO}_2$
- A43018_a01: k from $\text{CH}_3\text{O}_2 + \text{O}_2^-$
- A43019a_a01: k calculated comparing $\text{CH}_3\text{OH} + \text{OH} / \text{CH}_3\text{OOH} + \text{OH}$ with IPROPL + OH
- A43019b_a01: k calculated comparing $\text{CH}_3\text{OH} + \text{OH} / \text{CH}_3\text{OOH} + \text{OH}$ with $\text{ACETOL} + \text{OH} / \text{HYPERCET} + \text{OH}$
- A43020_a01: k taken from the reaction of the hydrated form of MGLYOX and NO_3^-
- A43021_a01: k from $\text{CH}_3\text{O}_2 + \text{HO}_2$
- A43022_a01: k from $\text{CH}_3\text{O}_2 + \text{O}_2^-$
- A43023_a01: pH-dependent
- A43025_a01: $k = 2 \times k(\text{HOCH}_2\text{OH} + \text{OH})$
- A44010_a01: $k = 2 \times k(\text{CHOHOHCHOHOH} + \text{OH})$
- A44011_a01: $k = 2 \times k(\text{CHOHOHCHOHOH} + \text{OH})$
- A44012_a01: $k = 2 \times k(\text{CHOHOHCHOHOH} + \text{OH})$
- A46000_a01: Assumed to be the same as for glyoxal
- A46001_a01: Assumed to be the same as for glyoxal
- A46002_a01: $k = 2 \times k(\text{CH}_3\text{COCHOHOH} + \text{OH})$
- A46003_a01: Assumed to be the same as for glyoxal
- A46004_a01: Assumed to be the same as for glyoxal
- A46005_a01: $k = 2 \times k(\text{CH}_3\text{COCHOHOH} + \text{OH})$
- A460MS_a01: Ryder et al. say that k is the same as for $\text{NO}_2 + \text{chloride}$; In reality 6% 4-nitrophenol, 80% 4-nitrosophenol and 14% 2-nitrophenol is formed but we

assume that only 2-nitrophenol is formed like in the gas phase.

A61002_a01: Jacobi (1996) found an upper limit of 6E9 and cite an upper limit from another study of 2E9. Here, we set the rate coefficient to 1E9.

A63001_a01: There is also an earlier study by Exner et al. (1992) which found a smaller rate coefficient but did not consider the back reaction.

A64000_a01: k taken from $\text{H}_2\text{O}_2 + \text{Cl}_2^-$ (Yu, 2004).

A74000_a01: Assumed to be the same as for $\text{Br}_2^- + \text{H}_2\text{O}_2$.

A76003_a01: The rate coefficient is defined as backward reaction divided by equilibrium constant.

A91005_a01: The rate coefficient for the sum of the paths (leading to either HSO_5^- or SO_4^{2-}) is from Huie and Neta (1987), the ratio 0.28/0.72 is from Deister and Warneck (1990).

A91006_a01: See also: (Huie and Neta, 1987; Warneck, 1991). If this reaction produces a lot of SO_4^- , it will have an effect. However, we currently assume only the stable $\text{S}_2\text{O}_8^{2-}$ as product. Since $\text{S}_2\text{O}_8^{2-}$ is not treated explicitly in the mechanism, SO_4^{2-} is used as a proxy and the second sulfur atom is put into the lumped LSULFUR.

A92005_a01: D. Sedlak, pers. comm. (1993).

A92008_a01: D. Sedlak, pers. comm. (1993).

A930MS_a01: As suggested in Staudt et al., 2019 intermediate product is NO_2SO_4^- , which readily reacts with H_2O and produce final products

A94100_a01: $2.48 \times 10^7 \times 5.5 \times 10^{-4}$, considering the hydrated form of HCHO .

A94102_a01: $790 \times 5.5 \times 10^{-4}$, considering the hydrated form of HCHO .

A94108a_a01: $k(\text{H}_2\text{O}_2 + \text{SO}_4^-)$, branching ratio as for $\text{CH}_3\text{OOH} + \text{OH}$

A94108b_a01:	See branch a.	A96005_a01:	Assumed to be the same as for $\text{SO}_3^{2-} + \text{HOCl}$.	A119004_a01:	Assumed. Note that CAPRAM 2.4 lists $k=4.3\text{E}7$ from Herrmann Air Pollution Research Report 57 and it also lists $k= 2.65\text{E}7$ from Williams PhD 1996 http://lib.leeds.ac.uk/record=b1835184~S5 . Brand and van Eldik (1995) also list $k=3.56\text{E}4$ from Waygood EUROTRAC 1992 report.
A94200_a01:	$k(\text{CH}_3\text{OO} + \text{HSO}_3^-)$	A97005_a01:	Assumed to be the same as for $\text{SO}_3^{2-} + \text{HOBr}$.	A119006_a01:	$3\text{E}8 * 6500 / (48000 + 6500)$
A94201_a01:	$k(\text{CH}_3\text{OO} + \text{HSO}_3^-)$	A116005_a01:	products assumed	A119008_a01:	Assuming that the intermediate $\text{S}_2\text{O}_6^{2-}$ dissociates quickly.
A94202a_a01:	$\text{CH}_3\text{CHOHO}_2$ is assumed to directly decompose into $\text{CH}_3\text{CHO} + \text{HO}_2$	A119001_a01:	products assumed		
A94203b_a01:	CHOHOHO_2 directly decomposes into $\text{HCOOH} + \text{HO}_2$	A119002_a01:	products assumed		

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