

Emissions in UKCA: What goes in must... do something?!

Introduction to UKCA, 9th-13th January 2017, Cambridge

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Garbage in garbage out!

- It's as simple as that really. Numerical models take inputs, operate on these and produce outputs.
- If you want high quality, meaningful outputs then you need to think about BOTH what your model is doing and what you are putting into your model.





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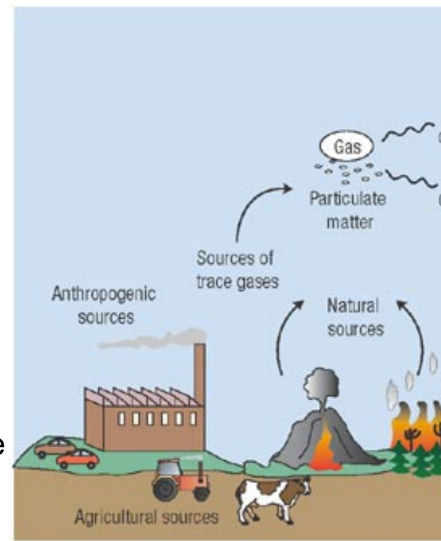
Sources of emissions:

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- There are mainly two sources of emissions:
 - Natural

• Anthropogenic

 We can also divide emissions by their phase (gas vs aerosol).

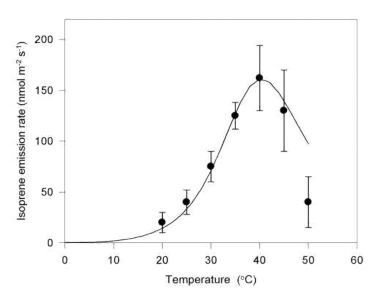


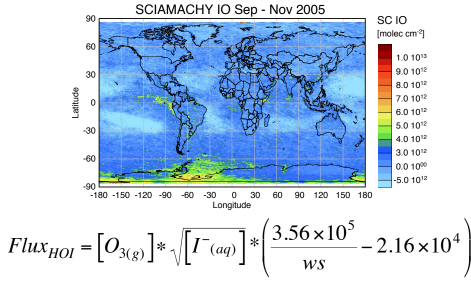


Natural emissions

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- Produced through natural processes e.g. photosynthesis, respiration, wild fires etc. So often have a dependence on environmental factors
 - Some of the most interesting "new" research topics deal with the feedbacks between climate and natural emissions.







Anthropogenic emissions

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- Produced by man-made processes e.g. pasture burning, agriculture, industry etc. Can have "environmental dependence" but generally not.
- For some compounds anthropogenic emissions are the dominate source.
- Predicted to change due to socio-economic factors.



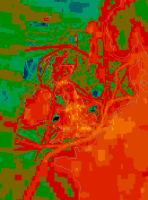




- 1. Introduction
- 2. Ancillary emission system
- 3. NetCDF emission system
- 4. Final notes on UKCA
- 5. Getting hold of emissions data



- Offline: Pre-computed fluxes
 - Read in from ancillary or NetCDF files
 - 12-monthly varying 2-D or 3-D fields



- Online: Computed in real-time using fields from the UM. Examples:
 - Lightning NO_x
 - Sea-salt (in GLOMAP-mode)
 - Wetland CH₄ (from JULES)
 - Interactive BVOC (from JULES; for isoprene, terpenes, methanol, acetone; since UM vn9.2).



Emission Species	Units	Std	Trop-	RAQ	Strat	Strat+
		Trop	Isop	Chem	Chem	Trop
NOx Surface Emissions	$kg(NO2)/m^2/s$	Y	Υ	Υ	Y	Υ
CH4 Surface Emissions	$kg(CH4)/m^2/s$	Y	Υ	Υ	Υ	Y
CO Surface Emissions	$ m kg/m^2/s$	Y	Υ	Υ	Υ	Υ
HCHO Surface Emissions	$ m kg/m^2/s$	Y	Υ	Υ	Y	Υ
C2H6 Surface Emissions	$ m kg/m^2/s$	Y	Υ	Υ		Υ
C3H8 Surface Emissions	$ m kg/m^2/s$	Y	Υ	Υ		Y
CH3COCH3 Surface Emissions	$ m kg/m^2/s$	Y	Υ	Υ		Υ
CH3CHO Surface Emissions	$ m kg/m^2/s$	Y	Υ	Υ		Y
C5H8 (Isop) Surface Emiss	$\rm kg(C)/m^2/s$		Y	Υ		Y
H2 Surface Emissions	$kg/m^2/s$			Υ		
C4H10 Surface Emissions	$ m kg/m^2/s$			Υ		
C2H4 Surface Emissions	$ m kg/m^2/s$			Υ		
C3H6 Surface Emissions	$ m kg/m^2/s$			Υ		
Toluene Surface Emissions	$ m kg/m^2/s$			Υ		
o-xylene Surface Emissions	$ m kg/m^2/s$			Υ		
CH3OH Surface Emissions	$kg(C)/m^2/s$			Υ		
NOx Aircraft Emissions (3D)	$kg(NO2)/m^2/s$	Y	Y	Υ	Υ	Υ



Aerosol & online emissions

Emission Species	Units	Std	Trop-	RAQ	Strat	Strat+
		Trop	Isop	Chem	Chem	Trop
If using Aerosol Chem						
Monoterpene Surface Emiss	$kg(C)/m^2/s$	Y	Y			Y
NVOC Surface Emiss	$kg(C)/m^2/s$		Υ			Y
SO2 Surface Emissions	$kg(S)/m^2/s$	Y	Υ		Υ	Y
DMS Surf Emiss (Land, ocean)	$\rm kg(S)/m^2/s$	Υ	Υ		Y	Y
NH3 Surface Emissions	$ m kg/m^2/s$	Υ	Υ			Y
SO2 High (Ind,forest,ship)	$kg(S)/m^2/s$	Υ	Υ		Υ	Y
SO2 Volcanic Emissions (3D)	$\rm kg(S)/m^2/s$	Y	Υ		Y	Y
for GLOMAP-mode (if using 5-mode setup)						
BC Fossil Fuel Emiss	$kg(C)/m^2/s$	Y	Y			Y
OC Fossil Fuel Emiss	$\rm kg(C)/m^2/s$	Y	Υ			Y
BC Biofuel Emissions	$\rm kg(C)/m^2/s$	Y	Υ			Y
OC Biofuel Emissions	$kg(C)/m^2/s$	Y	Υ			Y
BC Biomass burn Emiss (3D)	$kg(C)/m^2/s$	Y	Υ			Y
OC Biomass burn Emiss (3D)	$kg(C)/m^2/s$	Υ	Υ			Y
Online Emissions						
CH4 Wetland Emiss (in LSH - Optional)	$kg(C)/m^2/s$	Y	Y	Y	Y	Y
Lightning NOx (in UKCA)	kg(NO2)/kg(air)/cell/s	Υ	Υ	Y	Υ	Y
Sea-Salt Emiss (in GLOMAP-mode)	$number/m^2/s$	Υ	Y		Y	Υ



2. Ancillary emission system

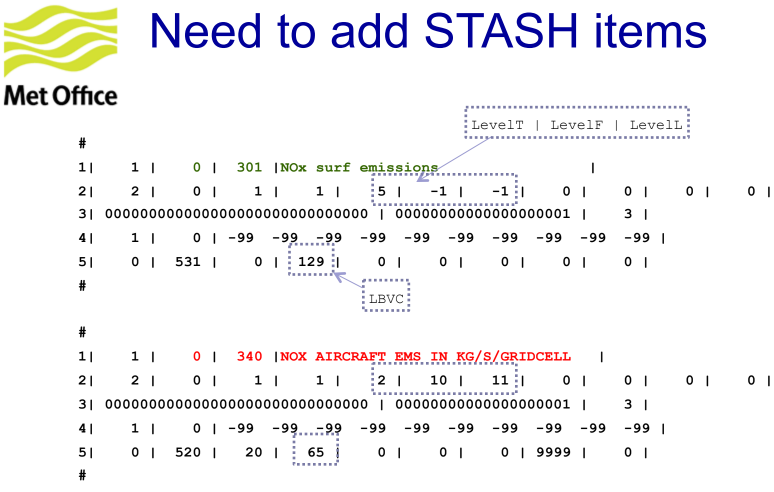


UKCA Makes use of user ancillary files:

- single-level: items 301-320 (STASH section 0)
- multi-level: items 321-340 (STASH section 0)

STASH	Emission Species	Units	Std	Trop-	RAQ	Strat	Strat+
code			Trop	Isop	Chem	Chem	Trop
301	NOx Surface Emissions	$kg(NO2)/m^2/s$	Y	Y	Y	Y	Y
302	CH4 Surface Emissions	$kg(CH4)/m^2/s$	Y	Y	Υ	Y	Y
303	CO Surface Emissions	$ m kg/m^2/s$	Y	Υ	Υ	Υ	Y
304	HCHO Surface Emissions	$ m kg/m^2/s$	Y	Y	Υ	Υ	Y
305	C2H6 Surface Emissions	$ m kg/m^2/s$	Y	Y	Υ		Y
306	C3H8 Surface Emissions	$ m kg/m^2/s$	Y	Υ	Υ		Y
307	CH3COCH3 Surface Emissions	$ m kg/m^2/s$	Y	Y	Υ		Y
308	CH3CHO Surface Emissions	$ m kg/m^2/s$	Y	Y	Υ		Y
309	C5H8 (Isop) Surface Emiss	$kg(C)/m^2/s$		Y	Υ		Y
	·		I	l	i I	1	, , , , , , , , , , , , , , , , , , ,
340	NOx Aircraft Emissions (3D)	$kg(NO2)/m^2/s$	Y	Y	Y	Y	Y

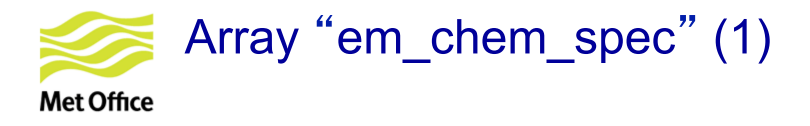
Rest of items in section 0 reserved for UM prognostics



Meaning:

- Item 301: Single level data (surface level)
- Item 340: Data on atmosphere theta levels (from first to last tracer level)

More info in UMDP C4 "Storage Handling And Diagnostic System (STASH)" Basically 2-D fields will be like 301 and 3-D like 340.



- Array with list of emitted species/fields
 Scheme specific. Declared in ukca_d1_defs.F90:
 CHARACTER(LEN=10), DIMENSION(:), ALLOCATABLE, SAVE :: em_chem_spec
- Allocated & filled in ukca_setd1defs.F90

(IF block for different chemistry schemes)

 NO_x emissions: Expressed as kg(NO_2) m⁻² s⁻¹ but assigned to the NO tracer!

. . .

Array "em chem spec" (2) Met Office

 NO_x emissions: Expressed as kg(NO_2) m⁻² s⁻¹ but assigned to the NO tracer!

If adding new emission field:

- Look for your chemistry scheme in the IF block of UKCA_SETD1DEFS. Update em_chem_spec.
- Need new STASH code?
- Need to update long block towards the end of UKCA_SETD1DEFS?

(See tutorial in the afternoon)





Both cases accept online emissions



Beginning of the routine → Some initialisations & IF blocks affecting different UKCA configurations/options. Examples:

```
! Regrid interative BVOC emissions from landpoints (1D) to 2D-grid
IF (L_bvoc_emis) THEN
  We get isoprene_2D, terpene_2D, methanol_2D, acetone_2D
END IF
```

Then very **long loop through all gas phase tracers** to (i) **add surface emissions**, including unit conversions if needed, and (ii) do **boundary layer mixing of tracers**. See next slide ...



ukca_emission_ctl.F90 (2)

! Check if tracer has **surface emissions** and set emission. Otherwise emission field is zero from initialisation.

DO k = 1, jpctr ! loop over tracers

DO I = 1, n_chem_emissions

! Rest of emissions
ELSE IF (advt(k) == em_chem_spec(I)) THEN
em_field(:,:,k) = all_emissions(:,:,I)
ENDIF ! end advt(k)
END DO ! I=1,n_use_emissions



!

ukca emission ctl.F90 (2)

- ! Check if tracer has surface emissions and set emission. Otherwise emission field is zero from initialisation.
- DO k = 1, jpctr ! loop over tracers DOI = 1, n chem emissions IF (advt(k) == em_chem_spec(l) .AND. em_chem_spec(l) == 'NO ') THEN Convert from kg NO2/m2/s to kg NO/m2/s em field (:,:,k) = all_emissions (:,:,l) * m_no / m_no2

```
!
      Rest of emissions
      ELSE IF (advt(k) == em_chem_spec(I) ) THEN
       em_field(:,:,k) = all_emissions(:,:,l)
                   ! end advt(k)
      ENDIF
  END DO
              ! I=1,n_use_emissions
```

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!

!

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. . . .

ukca_emission_ctl.F90 (2)

- Check if tracer has surface emissions and set emission. Otherwise emission field is zero from initialisation.
 DO k = 1, jpctr ! loop over tracers
 - DO I = 1, n_chem_emissions IF (advt(k) == em_chem_spec(I) .AND. em_chem_spec(I) == 'NO ') THEN Convert from kg NO2/m2/s to kg NO/m2/s em_field (:,:,k) = all_emissions (:,:,I) * m_no / m_no2

ELSE IF ... ! Many other checks (for SO2_low, DMS, Monoterp, C5H8, MeOH, Me2CO, ...). Note that BVOC emissions might be interactive

If you update the code check if there are specific unit conversions for your species and make sure that is consistent with your emission fields.

```
! Rest of emissions
ELSE IF (advt(k) == em_chem_spec(I) ) THEN
em_field(:,:,k) = all_emissions(:,:,I)
ENDIF ! end advt(k)
END DO ! I=1,n_use_emissions
```

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!

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. . . .

ukca_emission_ctl.F90 (2)

initialisation.
DO k = 1, jpctr ! loop over tracers
DO I = 1, n_chem_emissions
IF (advt(k) == em_chem_spec(I) .AND. em_chem_spec(I) == 'NO ') THEN
! Convert from kg NO2/m2/s to kg NO/m2/s
em_field (:,:,k) = all_emissions (:,:,l) * m_no / m_no2

! Check if tracer has surface emissions and set emission. Otherwise emission field is zero from

ELSE IF ... ! Many other checks (for SO2_low, DMS, Monoterp, C5H8, MeOH, Me2CO, ...). Note that BVOC emissions might be interactive

If you update the code check if there are specific unit conversions for your species and make sure that is consistent with your emission fields.

```
! Rest of emissions
ELSE IF (advt(k) == em_chem_spec(I) ) THEN
em_field(:,:,k) = all_emissions(:,:,I)
ENDIF ! end advt(k)
END DO ! I=1,n_use_emissions
```

IF (advt(k) == 'CH4 ' ...) THEN Add wetland CH4 emissions or prescribe the surface mixing ratio

IF (L_ukca_strat .OR. L_ukca_stratcfc .OR. L_ukca_strattrop) THEN

! Treatment of long-lived species with lower boundary condition

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!

1

. . . .

ukca_emission_ctl.F90 (2)

initialisation.
DO k = 1, jpctr ! loop over tracers
DO I = 1, n_chem_emissions
IF (advt(k) == em_chem_spec(I) .AND. em_chem_spec(I) == 'NO ') THEN
! Convert from kg NO2/m2/s to kg NO/m2/s
em_field (:,:,k) = all_emissions (:,:,I) * m_no / m_no2

! Check if tracer has surface emissions and set emission. Otherwise emission field is zero from

ELSE IF ... ! Many other checks (for SO2_low, DMS, Monoterp, C5H8, MeOH, Me2CO, ...). Note that BVOC emissions might be interactive

If you update the code check if there are specific unit conversions for your species and make sure that is consistent with your emission fields.

```
! Rest of emissions
ELSE IF (advt(k) == em_chem_spec(I) ) THEN
em_field(:,:,k) = all_emissions(:,:,I)
ENDIF ! end advt(k)
END DO ! I=1,n_use_emissions
```

IF (advt(k) == 'CH4 ' ...) THEN

! Add wetland CH4 emissions or prescribe the surface mixing ratio

IF (L_ukca_strat .OR. L_ukca_stratcfc .OR. L_ukca_strattrop) THEN

! Treatment of long-lived species with lower boundary condition

```
    Call boundary layer mixing and add surface emissions.
    CALL TR_MIX ( ..., em_field (:, :, k), ..., tracers (:, :, 1:bl_levels, k), ....)
```



ukca_emission_ctl.F90 (3a)

Similar stuff for GLOMAP-mode

! Beginning of ukca_emission_ctl. Call	routine for primary emissions for UKCA-MODE
IF (L_ukca_mode) THEN	
CALL ukca_mode_ems_um -	<pre>em_field_mode is filled with nr & mass</pre>
END IF	emission fluxes for each tracer

- 1. Initial emission arrays for aerosols are created from the **input data** (depending on the model set-up)
- 2. CALL ukca_mode_ems: returns mass & number emission arrays for sulphate, sea-salt, OC, BC and dust (as required)
- 3. The number and mass fluxes are then assembled for each tracer in the **array em_field_mode**
- 4. **Diagnostics** for emitted component mass are stored in STASH
 - 38 | 201 | PRIMARY SO4 TO AITKEN (SOL)
 38 | 202 | PRIMARY SO4 TO ACCUMULATION (SOL)
 38 | 203 | PRIMARY SO4 TO COARSE (SOL)
 38 | 204 | PRIMARY SEA-SALT TO ACCUMULTN (SOL)
 38 | 205 | PRIMARY SEA-SALT TO COARSE (SOL)
 38 | 206 | PRIMARY BLACK CARBON TO AITKEN (SOL)
 38 | 207 | PRIMARY BLACK CARBON TO AITKEN (INS)
 38 | 208 | PRIMARY ORG. CARBON TO AITKEN (SOL)
 38 | 209 | PRIMARY ORG. CARBON TO AITKEN (INS)
 38 | 210 | PRIMARY DUST TO ACCUMULATION (SOL)
 38 | 211 | PRIMARY DUST TO ACCUMULATION (INS)
 - 38 | 212 | PRIMARY DUST TO COARSE (SOLUBLE)



Similar stuff for GLOMAP-mode

! Beginning of ukca emission ctl. Call routine for primary emissions for UKCA-MODE

IF (L_ukca_mode) THEN CALL ukca_mode_ems_um → em_field_mode is filled with nr & mass END IF emission fluxes for each tracer

! After doing injection & mixing of gas tracers. Do same for MODE aerosols

IF (L ukca mode) THEN

! Call boundary layer mixing and add surface emissions ! for GLOMAP-mode aerosol tracers

DO kaer = 1, n_mode_tracers ! loop over tracers

CALL **TR_MIX** (..., em_field_mode (:, :, 1, kaer), ..., mode_tracers (:, :, 1:bl_levels, kaer), ...)

END DO ! end of loop over tracers (kaer)

END IF ! if L UKCA MODE



Similar stuff for GLOMAP-mode

! Beginning of ukca emission ctl. Call routine for primary emissions for UKCA-MODE

IF (L_ukca_mode) THEN CALL ukca_mode_ems_um → em_field_mode is filled with nr & mass END IF emission fluxes for each tracer

! After doing injection & mixing of gas tracers. Do same for MODE aerosols

IF (L ukca mode) THEN

END DO ! end of loop over tracers (kaer)

END IF ! if L UKCA MODE

ukca_emission_ctl.F90 (4)

! Diagnose NO2 lightning emissions

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```
lightningem_n_gridbox = 0.0
lightningem_no2_to_air = 0.0
CALL UKCA_LIGHT_CTL( ...
lightningem_n_gridbox (1:row_length,1:rows,1:model_levels),
lightningem_no2_to_air (1:row_length,1:rows,1:model_levels))
```

&

&

&

&



Finally, **treatment of SO₂ emissions** (if aerosol chemistry):

•Add (3-D volcanic + high-level anthropogenic) emissions to SO₂ tracer.

NOTE: The code removes direct sulphate fraction of emissions and converts from kg(S) to kg (SO₂)

•Add 3-D emissions from explosive volcanic eruptions (e.g. Pinatubo 1991 and 5 others):

CALL UKCA_VOLCANIC_SO2

Last note (only for ASAD framework). Some calls to output emission diagnostics:

CALL

asad_emissions_diagnostics

CALL

asad_3D_emissions_diagnostics



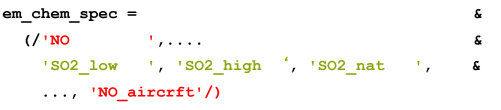
Advantages

Well tested for all UKCA configurations:

- Works with all chemistry schemes & with GLOMAP-mode
- UKCA_EMISSION_CTL accepts all online emissions available in UKCA

Limitations

Cannot easily treat emissions from independent source sectors separately.
 Example:



• Units in emission fields need to be as indicated in UKCA_EMISSION_CTL. However the UKCA code cannot check whether that is the case in your ancillary emission file!



3. NetCDF emission system



Available since UM vn8.6

Advantages

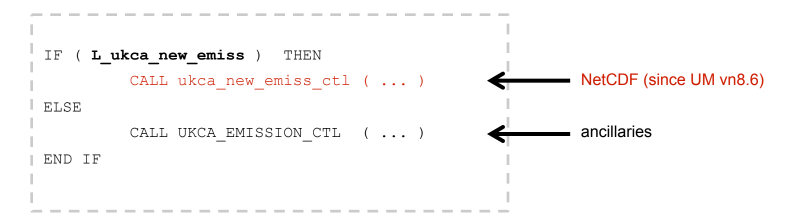
- Allow the use of different emission fields to account for independent source sectors for any given tracer
- Emissions can be injected at different altitudes and with different temporal variability
- Avoid inconsistencies in the units of the emission fields (to comply with CF conventions units should be "kg m⁻² s⁻¹")

Limitations

- Currently tested only for RAQ & Strat-Trop chemistry
- Needs to be extended for:
 - aerosol emissions (I_ukca_mode .OR. I_ukca_aerchem .OR. I_ukca_nr_aqchem)
 - iBVOC emissions (L_ukca_ibvoc .AND. L_bvoc_emis)



- As before "em_chem_spec" is allocated & filled within ukca_setd1defs.F90 Each element of the array could correspond to 1 or more emission fields
- From ukca_main1-ukca_main1.F90:



- ukca_new_emiss_ctl.F90 includes calls to many routines in order to:
 - Go through NetCDF emission files to automatically identify emission fields present in them
 - Initialise an "emissions" structure which will hold all emission fields
 - Update the "emissions" structure when needed
 - Output emission diagnostics

Met Office (Only available at UM vn8.6)

dlna.f: "cp ansyq (AQU 🚊 🗆 🗙

Include Y/N

Edit

UKCA

files

1es

Close

on UKCA Emiss. Job dlna.f.

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onoose the relevant section release	´ ● <1A> UK0	CA included.								
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 Stan 	dard Troposphe	eric(BE)								
Select Chemical Scheme RAQ	(BE)									
• Trop	ospheric+Isopre	ene								
 Stan 	dard Stratosphe	eric							ਜ UKCA NetC	DF Emission System : Jol
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Set Backward Euler Solver Sett	ings to non-defa	ault values?								
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🗆 Use a cyclic, monthly-va	rying 'backgrou	nd' aerosol field	d instead of time	eseries					FUSITORGA LO	go to the parent window
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NAMELIST input through Rose (I)

Met Office

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Section 33 - Free T Section 34 - UKCA:	🕼 l_ukca_intdd	✓ true	
Section 35 - Stocha	UKCA interactive dry deposition scheme	— — .	
Section 39 - Nudgi	🕼 l_ukca_new_emiss	□ false	
Short term logicals	Turn on the new emission system in UKCA (NetCDE)		
JULES Science Setting			
Data Assimilation	Specify which tracers have lateral boundary condition data in the LBC input file.		
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NAMELIST input through Rose (II)

Met Office

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Reconfiguration and A Number of wavelengths to be used								
IO System Settings O Added Input and Output Cut-off Pressure for tabulated photolysis	20.0							
r Model input and outp								
UM Science Settings General Physics Og Method above cut-off level	only lookup table							
Planet Constants	Fast-JX and lookup table							
Section 01 - 02 - B	O Fast_X only							
Section 03 - Bound Section 03 - Bound	◎ I_ukca_prescribech4							
Section 04 - Microp								
Section 05 - Conve UKCA 03 in radiation scheme	🗉 false							
Section 06 - Gravit	🗉 false							
Section 09 - Large VKCA_radcn4 VKCA CH4 in radiation scheme	Tabe							
 Sections 10 11 12 - Section 13 - Diffusi Lukca_intdd 	✓ true							
Section 14 - Energy								
Section 17 - Aerosc I_ukca_new_emiss	✓ true							
Section 21 - Thunde Turn on the new emission system in UKCA (N	etCDF)							
Section 26 - River f 👘 ukca_em_dir	'\$INPUT_DATA/AQUM'							
Section 33 - Free Tr Directory pathname for NetCDF emission file			_					
Section 34 - UKCA: 0 wkca_em_files	⁻ qrclim.ukca.surf.C2H4.nc	qrclim.ukca.surf.C2H6.nc						
Section 35 - Stocha Section 39 - Nudgir	qrclim.ukca.surf.C3H6.nc	qrclim.ukca.surf.C3H8.nc						
Short term logicals	qrclim.ukca.surf.C4H10.nc	qrclim.ukca.surf.C5H8.nc						
JULES Science Setting	qrclim.ukca.surf.CH3OH.nc	qrclim.ukca.surf.CH4.nc						
Data Assimilation	qrclim.ukca.surf.CO.nc	qrclim.ukca.surf.H2.nc						
	qrclim.ukca.surf.HCHO.nc	qrclim.ukca.surf.ME2CO.nc						
	qrclim.ukca.surf.MECHO.nc	qrclim.ukca.surf.NO.nc						
	qrclim.ukca.surf.OXYLENE.nc	qrclim.ukca.surf.TOLUENE.nc						
	qrclim.ukca.airc.nc	< >						



Maximum nr of emission files: 40

ukca_option_mod.F90:

INTEGER, PARAMETER :: nr_cdf_files = 40
CHARACTER (LEN=120) :: ukca_em_dir
CHARACTER(LEN=50) :: ukca_em_files(nr_cdf_files)

Each NetCDF file may include one or several emission fields

- Should comply with:
 - CF conventions (<u>http://cfconventions.org/</u>)
 - Met Office NetCDF conventions

However UM code could still work if files don't comply.

• Next slides: **Metadata attributes** needed in the files so that the UM code works!



Global attributes needed in each emission file:

 update_freq_in_hours: frequency in hours at which all emission fields present in that file should be read.

If you want 5 days (as done for ancillaries in many UMUI jobs):

update_freq_in_hours = 120

- emission type (follows same conventions as for ancillary files):
 - 0: Single time (not allowed yet)
 - 1: Time series
 - 2: Periodic time series



Met Office Some metadata attributes required for each emission field:

- **name:** Name of the emission field (80 characters, only for debugging)
- tracer_name: This has to be equal to one of the names in the list of emissions for the given chemical scheme, i.e. em_chem_spec
- **long_name:** Compulsory if there is no 'standard_name' available. Example: "tendency of atmosphere mass content of nitrogen monoxide due to emission"
- **units** = "kg m-2 s-1"

• hourly_scaling, daily_scaling & vertical_scaling: Characters read by the UM to apply corresponding numeric scaling factors



What UKCA code does with NetCDF attributes

Met Office

! Emission Data structure				
TYPE ukca_em_struct				
CHARACTER (LEN=256)	:: file_name			
CHARACTER (LEN=80)	:: var_name			
CHARACTER (LEN=10)	:: tracer_name			
CHARACTER (LEN=256)	:: std_name			
CHARACTER (LEN=256)	:: lng_name			
CHARACTER (LEN=30)	:: units			
INTEGER	:: update_freq			
INTEGER	:: update_type			
LOGICAL	:: l_update			
LOGICAL	:: three_dim			
REAL	:: base_fact			
REAL, POINTER	<pre>:: vert_scaling_3d (:,:,:)</pre>			
CHARACTER (LEN=20)	:: hourly_fact			
CHARACTER (LEN=20)	:: daily_fact			
CHARACTER (LEN=30)	:: vert_fact			
INTEGER	:: lowest_lev			
INTEGER	:: highest_lev			
REAL, POINTER	:: values (:,:,:)			
REAL, POINTER	:: diags (:,:,:)			

END TYPE ukca_em_struct

! Super array of emissions TYPE (ukca_em_struct), ALLOCATABLE :: emissions (:) 1. The UKCA code creates an "emissions" structure (derived type variable) which contains objects representing emissions. Each object can be distinguished by some components: emission name, values, units, vertical/temporal profiles, ...

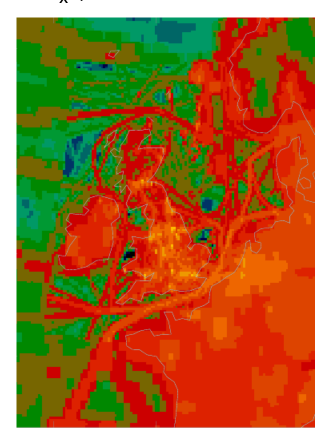
2. Then some of the NetCDF attributes are assigned to those components

Attribute in NetCDF files	Corresponding name in emissions structure	
	within the UKCA code	
name	emissions(:)%var_name	
standard_name	emissions(:)%std_name	
long_name	emissions(:)%lng_name	
tracer_name	emissions(:)%tracer_name	
units	emissions(:)%units	
hourly_scaling	emissions(:)%hourly_fact	
daily_scaling	emissions(:)%daily_fact	
vertical_scaling	emissions(:)%vert_fact	
lowest_level	emissions(:)%lowest_lev	
highest_level	emissions(:)%highest_lev	
update_freq_in_hours (global)	emissions(:)%update_freq	
emission_type (global)	emissions(:)%update_type	



Example of time / vertical profiles for RAQ (1)

Typical 12-monthly emission field for NO_x (all source sectors combined)

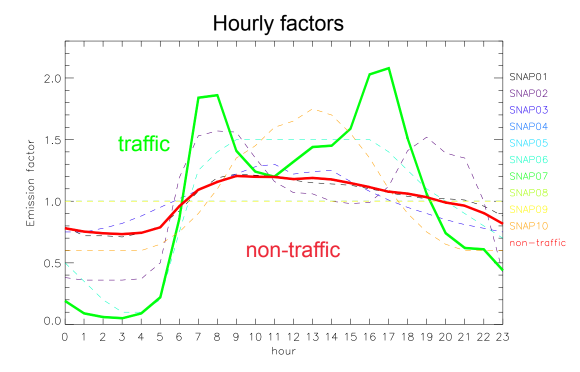


Better approach: Use separate SNAP source sectors

- 1 Combustion in energy and transformation industries
- 2 Non-industrial combustion plants
- 3 Combustion in manufacturing industry
- 4 Production processes
- 5 Extraction and distribution of fossil fuels
- 6 Solvents
- 7 Road transport ("traffic")
- 8 Other mobile sources (e.g. shipping)
- 9 Waste treatment and disposal
- 10 Agriculture
- 11 Natural and biogenic sources



- Hourly & daily factors to account for daily and weekly variability in emissions
- Based on data provided by TNO for the MACC project





Module UKCA_EMISS_FACTORS - Subroutine HOURLY_EMISS_FACTORS

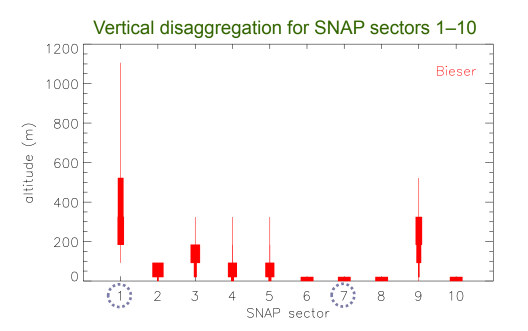
```
SELECT CASE (TRIM (hourly fact)) 

← NetCDF attribute hourly scaling
    ! No hourly factors applied unless specified
    CASE ('none', '')
      hourly scaling =
                                                                &
         (/1.00, 1.00, 1.00, 1.00, 1.00, 1.00, 1.00, 1.00,
                                                                &
           1.00, 1.00, 1.00, 1.00, 1.00, 1.00, 1.00, 1.00,
                                                                δ
           1.00, 1.00, 1.00, 1.00, 1.00, 1.00, 1.00, 1.00/)
    ! Hourly factors of emissions for Europe. Calculated by TNO
    ! for the MACC project.
    . . .
    CASE ('TNO MACC EU SNAP07')
      hourly scaling =
                                                                &
         (/0.19, 0.09, 0.06, 0.05, 0.09, 0.22, 0.86, 1.84,
                                                                &
           1.86, 1.41, 1.24, 1.20, 1.32, 1.44, 1.45, 1.59,
                                                                &
           2.03, 2.08, 1.51, 1.06, 0.74, 0.62, 0.61, 0.44/)
   . . .
END SELECT
```

0 1 2 3 4 5 6 7 8 9 10 11 12 13 14 15 16 17 18 19 20 21 22 23



Vertical profiles tested for anthropogenic emissions in RAQ chemistry scheme





Example of time / vertical profiles for RAQ (2b)

Module UKCA_EMISS_FACTORS - Subroutine VERTICAL_EMISS_FACTORS

```
! Altitude of the interfaces in metres
interf_ref = (/0.0, 20.0, 92.0, 184.0, 324.0, 522.0, 781.0, 1106.0/)
```

```
! Average vertical profiles for various SNAP source sectors.
CASE ('Bieser_modified_SNAP01')
vert scaling ref (:) = (/ 0.0, 0.0, 0.25, 51.0, 45.3, 3.25, 0.2 /)
```

```
CASE ('Bieser_modified_SNAP07')
vert_scaling_ref (:) = (/ 100.0, 0.0, 0.0, 0.0, 0.0, 0.0, 0.0 /)
```

•••

. . .

```
! Surface emiss in lowest model lev unless something different specified
CASE ('surface', '')
  vert scaling 3d (:, :, 1)
                                 = 1.0
                                                        1200
  vert scaling 3d (:, :, 2:) = 0.0
                                                                                 Rieser
                                                        1000
! 3D emissions over all model levels
                                                         800
                                                       altitude (m)
CASE ('all levels', '3D')
                                                         600
  vert scaling 3d (:,:,:) = 1.0
                                                         400
. . .
                                                         200
```

0

1 2 3 4

5 6 7 8 9 10

SNAP sector



• Attribute units in all NetCDF emission fields:

```
units = "kg m-2 s-1" or units = "kg/m2/s"
```

• If you want to express in kg(N), kg(C) or kg(S):

Check http://cfconventions.org/Data/cf-standard-names/27/build/cf-standard-name-table.html

- See code in 2 routines within the module **ukca_emiss_factors**:
 - Strings automatically detected by **base_emiss_factors**
 - Conversions done in get_base_scaling

You might need to adapt them



New emission diagnostics (1)

Emission field	Item number
(in em_chem_spec)	(in Sect. 50)
'NO '	156
'CH4 '	157
'CO '	158
'HCHO'	159
'C2H6 '	160
'C3H8 '	161
'Me2CO '	162
'MeCHO '	163
'C5H8 '	164
'C4H10 '	165
'C2H4 '	166
'C3H6 '	167
'TOLUENE '	168
'oXYLENE '	169
'CH3OH '	170
'H2 '	171
'NO_aircrft'	172

ukca_add_emiss_mod.F90

emissions(I)%diags (:, :,1) filled Column integrated & with time profiles applied

ukca_emdiags_struct_modF90: Declares a structure for emission diagnostics:

```
TYPE emdiags struct
   ! Flags indicating if diagnostics selected
  LOGICAL :: l em no
  LOGICAL :: 1 em ch4
   . . .
   ! Pointers to hold emission diagnostics
  REAL, POINTER :: em no
                            (:,:)
  REAL, POINTER :: em ch4 (:,:)
  REAL, POINTER :: em no air (:,:,:)
END TYPE emdiags struct
```

ukca_update_emdiagstruct_mod.F90

Update emission diagnostic values and flags in the "emdiags struct"

ukca_emiss_diags_mod.F90 Output emission diagnostics

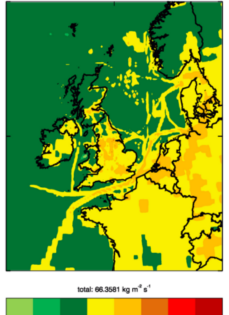
- Available for RAQ & other schemes
- New items can be added in S50



Emission diagnostics for NO:

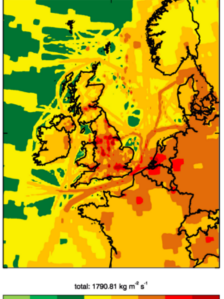
- 1. With temporal factors applied
- 2. Column integrated (always for 2-D emissions)
- 3. In kg (NO) m⁻² s⁻¹





1.0E-09 1.0E-06 1.0E-03 1.0E-02 1.0E-01 5.0E-011.0E+00

em_S50_NO (kg m⁻² s⁻¹) 10/06/2010 08Z



1.0E-09 1.0E-06 1.0E-03 1.0E-02 1.0E-01 5.0E-01 1.0E+00



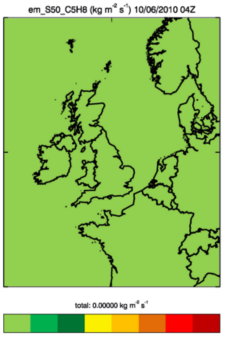
New emission diagnostics (3)

Met Office

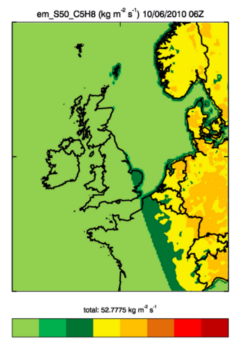
Emission diagnostics for (non-interactive) isoprene emissions, with netCDF attributes: tracer_name = 'C5H8'

hourly_scaling = 'diurnal_isopems'

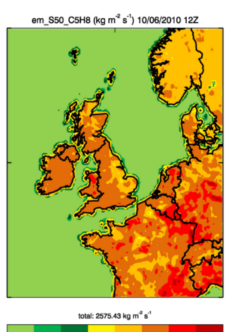
→ diurnal cycle applied via call to ukca_diurnal_isop_ems.F90



1.0E-09 1.0E-06 1.0E-03 1.0E-02 1.0E-01 5.0E-01 1.0E+00









NetCDF emission system will be extended for iBVOC emissions of C_5H_8 and other species



IF (I_first) THEN

! Read emission NetCDF files and look for the the emission fields ! in them to allocate all variables in the emissions structure. CALL ukca_emiss_init (...)

END IF

! Check if it is time to update the emiss fields that
 ! we read from NetCDF files (depending on time step
 ! and update frequency). If needed then update the files.

CALL ukca_emiss_update (...)

! **Deal with online emissions**. Always updated at each time step ! For the moment only NO_x from lightning and CH_4 from wetlands



! Update all fields in the emissions super array:

- !* Do conversions so that emissions are given as 'kg(tracer) m-2 s-1'
- !* Update isoprene emissions if they are diurnally varying

DO I = 1, num_em_flds

IF (emissions(I)%I_update) THEN

CALL base_emiss_factors

update "emissions(I)%values"

END IF

For **isoprene**: CALL ukca_diurnal_isop_ems to apply diurnal cycle

END DO

! Inject emissions and do tracer mixing CALL ukca_add_emiss (...)

! -----

! Call the emission diagnostics code if any of the diagnostics present ! in the routine GET_EMDIAG_STASH has been selected via stash.

CALL ukca_emiss_diags (...)



4. Final notes



- This talk is based on UM vn9.2. Some functionalities not available for older UM versions.
- I have not covered some topics in detail (aerosol emissions, BVOC emissions, ...).

Useful reading material

Online UKCA tutorial

http://www.ukca.ac.uk/wiki/index.php/UKCA_Chemistry_and_Aerosol_Tutorial_5

- UKCA UMDP 84
- Chapter 10: Emissions (ancillary system)
- Chapter 11: New NetCDF emission system

https://code.metoffice.gov.uk/doc/um/vn9.2/umdp.html https://code.metoffice.gov.uk/doc/um/vn10.0/umdp.html



Code in ukca_setd1defs.F90

Complicated block of code further down in this routine

J = n_use_tracers IF (n_chem_emissions+n_3d_emissions+n_mode_emissions > 0) THEN DO i=1, n chem emissions + n 3d emissions UkcaD1Codes(J+i)%section = 0 UkcaD1Codes(J+i)%item = n emiss first+i-1 UkcaD1Codes(J+i)%len dim1 = row length UkcaD1Codes(J+i)%len_dim2 = rows IF (.NOT. (L ukca new emiss)) THEN UkcaD1Codes(J+i)%required = .TRUE. END IF UkcaD1Codes(J+i)%prognostic = .TRUE. Special cases, emissions already available in UM IF (em_chem_spec(i)(1:7) == 'S02_low') THEN UkcaD1Codes(J+i)%item = 58 ELSEIF (em_chem_spec(i)(1:7) == 'S02_nat') THEN UkcaD1Codes(J+i)%item = 121 UkcaD1Codes(J+i)%len dim3 = tr levels ELSEIF (em_chem_spec(i)(1:8) == 'S02_high') THEN UkcaD1Codes(J+i)%item = 126 ELSEIF (em chem spec(i)(1:3) == 'NH3') THEN UkcaD1Codes(J+i)%item = 127 ELSEIF (em_chem_spec(i) == 'BC_fossil ') THEN UkcaD1Codes(J+i)%item = 310 ELSEIF (em_chem_spec(i) == 'BC_biofuel') THEN UkcaD1Codes(J+i)%item = 311 ELSEIF (em chem spec(i) == 'OC fossil ') THEN UkcaD1Codes(J+i)%item = 312 ELSEIF (em chem spec(i) == 'OC biofuel') THEN UkcaD1Codes(J+i)%item = 313 ELSEIF (em chem spec(i) == 'Monoterp ') THEN UkcaD1Codes(J+i)%item = 314 ELSEIF (em_chem_spec(i) == 'NVOC ') THEN UkcaD1Codes(J+i)%item = 315 ELSEIF (em chem spec(i) == 'BC biomass') THEN UkcaD1Codes(J+i)%item = 322 UkcaD1Codes(J+i)%len dim3 = tr levels ELSEIF (em chem spec(i) == 'OC biomass') THEN UkcaD1Codes(J+i)%item = 323 UkcaD1Codes(J+i)%len dim3 = tr levels ELSEIF (em_chem_spec(i) == 'SO2_biomas') THEN UkcaD1Codes(J+i)%item = 324 UkcaD1Codes(J+i)%len_dim3 = tr_levels ELSEIF (em_chem_spec(i)(1:3) == 'DMS') THEN UkcaD1Codes(J+i)%section = 17 UkcaD1Codes(J+i)%item = 205 UkcaD1Codes(J+i)%prognostic = .FALSE. IF (.NOT.L sulpc dms)UkcaD1Codes(J+i)%required=.FALSE. ! DMS internally calculated if CLASSIC is OFF ELSEIF (em chem_spec(i)(1:7) == 'NO_airc') THEN UkcaD1Codes(J+i)%item = 340 UkcaD1Codes(J+i)%len_dim3 = tr_levels ENDIF ENDDO ENDIF

Meaning

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- STASH 301–309 in same order as em_chem_spec
- STASH items defined for some emissions (e.g. items 322, 323, 324 reserved for 3D emissions)

Main thing to remember:

- single-level: items 301-320
- multi-level: items 321-340



5. Getting hold of some data



So where/how do we get information on emissions?

• By far the best source of information is GEIA:

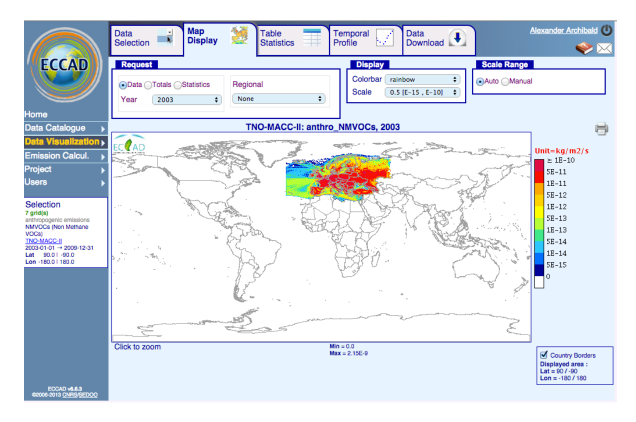
	ECCAD - THE GEIA DATABASE				
	LOGIN Enter Not yet registered?	😣 😒 😒			
ECCAD	Emissions of atmospheric Compounds & Compilation of Ancillary Data Data Catalogue Data Visualization Emission Calculation Emissions Inventories				
Home		Anthropogenic Biomass burning Natural			
Data Catalogue	GLOBAL INVENTORIES	REGIONAL INVENTORIES			
Data Visualization Emission Calcul.	MACCity ACCMIP RCPs EDGARv4.2 PEGASOS_PBL-v2 EDGARv3.2FT2000 RETRO	TNO-MACC-II (Europe) TNO-MACC (Europe)			
Project 🕨	ECLIPSE_GAINS_4a Junker-Liousse HYDE1.3 Andres_CO2_v2013 AMAP_Mercury	EMEP (Europe) Assamoi-Liousse (Africa)			
Users 🕨	GFASv1.0 GFED3 GFED2 GICC AMMABB	India_NOx (India) SAFAR-India (India) REAS (Asia)			
Newsletter #1	MEGAN-MACC MEGANv2 MEGANv2-CH3OH GEIAv1 POET				
	Developed for ongoing projects	Developed for ongoing projects			
Partners		ChArMEx (Mediterranean)			
	GUESS-ES GUESS-ES-Scenario				
	Ancillary Datasets				
	LAND COVER FIRES POPULATION	GEOGRAPHICAL INFORMATION			
	UMD CLM3 GLC2000 WFA GBA2000 Geoland2_BAv1_Africa GPW3_Population	GPW3 Region_IMAGE2.4 Pixel_Area			
ECCAD v6.6.3 ©2006-2013 <u>CNRS/SEDOO</u>					



So where/how do we get information on emissions?

• By far the best source of information is GEIA:

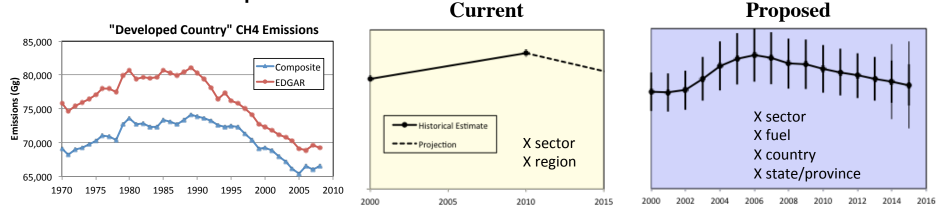
Easily select data for different compounds, from different datasets for different time periods and plot, download, visualize etc.





What about the future of emissions?

• Uncertainty quantification will be more important.



 Community Emissions Data System (CEDS) will be key for providing info for CMIP6 – you can get involved!