2024/03/29 13:28 1/11 PMIP4 and the CMIP6 DRS

PMIP4 and the CMIP6 DRS



All the *attributes* have to be defined properly when creating data for the CMIP6 database, but you will find below details about some attributes that are especially relevant for PMIP4

Some key concepts...

- attribute: a global attribute (e.g. in a NetCDF file) used to describe the data
- CV: sometimes the value of a given attribute has to be taken from a predefined set of values, known as a Controlled Vocabulary (CV)
- DRS = Data Reference Syntax: the DRS is used to identify experiments, simulations, ensembles of experiments, atomic datasets and is used, for example, in file names, directory structures, the further info url, and in facets of some search tools
- facet = a category or attribute you can put a search constraint on, when doing a faceted search



Example: the experiment_id attribute is used in the DRS, and its value has to be chosen from a CV ([piControl, past1000, lgm, ...]). You can put a search constraint on the Experiment facet by clicking on Experiment on the IPSL CMIP5 search node and then selecting lgm and clicking on </WRAP> ===== CMIP6 official specifications =====

The following CMIP6 document is still *in prep* (as of July 14th 2016)



This document specifies all the global attributes that are defined for CMIP6. It also indicates how a subset of those relate to the Data Reference Syntax (DRS) and are used in file names and directory structures. Controlled vocabularies are defined for some global attributes (e.g., source_type and grid_resolution).

* CMIP6 document: CMIP6 Global Attributes, DRS, Filenames, Directory Structure, and CV's (version 1.0) * Legacy CMIP5 documents: * CMIP5 Data Reference Syntax (DRS) and Controlled Vocabularies (Version 1.3.1 - 13 June 2012) * CMIP5 Model Output **Requirements: File Contents and Format, Data** Structure and Metadata (7 January 2010) ===== **Project identification attributes ===== *** activity id = activity labels * mip era = activity's associated CMIP cycle * Note: The project id used in CMIP5 is being replaced in CMIP6 with two global attributes: 1) an activity id, and 2) a mip era (a label indicating the cycle of CMIP that this activity falls under which will be set to "CMIP6" for the 6th CMIP cycle). In a few cases it may be appropriate to include multiple activities in the activity id (with multiple activities allowed, separated by single spaces). An example of this is "LUMIP AerChemMIP" for one of the land-use change experiments. ^ Project ^ activity id ^ mip era ^ Note ^ | CMIP6 | CMIP | CMIP6 | | | PMIP4-CMIP6 | CMIP

"CMIP PMIP"?? | CMIP6 | Should we use CMIP or "CMIP PMIP" for PMIP4 experiments that are part of CMIP6? This is confusing | PMIP4 | PMIP | PMIP4??



CMIP6??| Use this for non-CMIP6 experiments, or groups that are not part of CMIP6 Should we use PMIP4 because it is the 4th phase of PMIP, or CMIP6 because we will be using CMIP6 format specifications? | ==== Experiment names ==== You can find all the referenced experiment names on es-doc search site: select Project=CMIP6-DRAFT and Type=Experiment * experiment id = root experiment identifier * experiment = short expt. description * sub experiment id=none ⇒ needed for CMIP6 hindcast and forecast experiments to indicate "start year". For other experiments, this should be set to none * sub experiment=none ⇒ needed for CMIP6 hindcast and forecast experiments. For other experiments, this should be set to none ==== DECK and historical experiments ====



an historical experiment is the true continuation of a past1000 experiment?

We probably need to use parent experiment id=past100

2024/03/29 13:28 3/11 PMIP4 and the CMIP6 DRS



O in the files' metadata, as well as parent_activity_id, parent_mip_era=CMIP6, parent_source_id, branch_time_in_ parent and other related parent_* variables. We can probably also agree on a specific variant_label that will appear in the file names.

^ experiment id ^ experiment ^ | amip | Atmospheric Model Intercomparison Project | | piControl | Pre-Industrial Control | | abrupt-4xCO2 | abrupt quadrupling of CO2 | | 1pctCO2 | 1 percent per year increase in CO2 | | historical | all-forcing simulation of the recent past | ==== PMIP4-CMIP6 experiments ==== You can find specific details about the experiments by visiting the PMIP4 experimental design section, or by directly clicking on one of the experiments below ^ experiment id ^ experiment ^ | past1000 | past 1000 years | | mid-Holocene | mid-Holocene | | Igm | last glacial maximum | | lig127k | last interglacial | | midPlioceneeoi400 | mid-Pliocene | ==== PMIP4 only experiments ==== == Guidelines for creating new PMIP4 experiment id values === * Allowed characters: CMIP6 experiment id values are similar to CMIP5 (...so the permitted characters will be: a-z, A-Z, 0-9, and "-"), but a compound structure is allowed (segments separated by hyphens; e.g., "abrupt-4xCO2"); in a few cases multiple sub-experiments will be defined belonging to this root and constituting an ensemble. * Being wise when creating/proposing a new name * <code>Global attributes that label experiments are needed to construct file names and directories and can generally be used as search facets. Together, they should have the following characteristics: * Uniquely label each experiment within CMIP6 and distinguish experiments with specified conditions that differ in any way * **Easily be interpreted and remembered * Facilitate** representations of groups of experiments that are closely related (e.g., same forecast conditions but different start dates, or experiment with an "offline" model driven by output from various models)</code> * Planning for groups of related simulations * <code>Often several simulations will be performed that satisfy the conditions specified for each experiment. For example simulations of the historical period can branch from various points in a control run, and each of these will satisfy the conditions defining the experiment. Together such simulations constitute a "conforming ensemble" with member all

satisfying the same "root" experiment specifications. There are also occasional cases where the experiment designers (MIP leaders) define a family of related simulations and choose to label these with a common "root" experiment name. An example of this is the set of decadal prediction hindcasts that are all run similarly but started from different start dates (with each simulation identified by a different sub-experiment label). Such "defined ensembles" of experiments will be labeled with a "root" experiment name, and a "sub-experiment_id" will be used to distinguish among members in the ensemble.</code>
* Ensemble of simulations usually share a common experiment_id and have different ripf variant labels.
=== Proposed PMIP4 experiment id values ===



The following suggested PMIP4 experiment_id values should be considered as a *work in progress*, till they are validated!



You can find specific details about the experiments by visiting the PMIP4 experimental design section, or by directly clicking on one of the experiments below $^$ experiment_id $^$ experiment $^$ Status $^$ | LDv1-LGMspin | Last Glacial Maximum spinup | Work | | LDv1-transpin | Transient orbit and trace gases spinup (26-21 ka) | Work | LDv1 | Transient deglaciation (21-0 ka) | Work | ==== Handling groups of simulations ===== CMIP5 ensemble member ==== aka r<N>i<M>p<L> or rip

The definitions below have been superseded by CMIP6 specifications, but it is still useful to remember them. They have been copied from:



- a) CMIP5 Data Reference Syntax (DRS) and Controlled Vocabularies (Version 1.3.1 -13 June 2012)
- b) CMIP5 Model Output Requirements: File Contents and Format, Data Structure and Metadata (7 January 2010)

https://pmip4.lsce.ipsl.fr/ Printed on 2024/03/29 13:28

* b) ⇒ realization = an integer (≥1) distinguishing among members of an ensemble of simulations (e.g., 1, 2, 3, etc.). If only a single simulation was performed, then it is recommended that realization=1.

For fields appearing in table "fx" in the CMIP5 Requested Output, set realization=0 (violating the general rule that it should be a positive definite integer).

Note that if two different simulations were started from the same initial conditions, the same realization number should be used for both simulations. For example if a historical run with "natural forcing" only and another historical run that includes anthropogenic forcing were initiated from the same point in a control run, both should be assigned the same realization. Also, each so-called RCP (future scenario) simulation should normally be assigned the same realization integer as the historical run from which it was initiated. This will allow users to easily splice together the appropriate historical and future runs. A similar convention should be followed, when appropriate, with other simulations (e.g., the decadal simulations).

Note that the realization can be used in constructing the "ensemble member" called for by the DRS document; it is the value of N in r < N > i < M > p < L >.



[Note that for the "Transpose AMIP" project, the "realization" number is used to distinguish among the 16 members of each of 4 suites of runs (i.e., the 4 "seasons") generated from different observed conditions, spaced 30 hours apart. So, for example, the 16-member ensemble of runs initialized at 00Z on 15 Oct 2008, 06Z 16 Oct 2008, 12Z 17 Oct 2008, and so-on, would be assigned "r1", "r2", "r3", etc.] * b) ⇒ initialization_method = an integer (≥1) referring to the initialization method used or different observational datasets used to initialize. If only a single method and dataset was used to initialize the model, then this argument should

normally be given the value 1.

For fields appearing in table "fx" in the CMIP5

Requested Output, set initialization_method=0

(violating the general rule that it should be a positive definite integer).

See the DRS document for guidance on assigning initialization_method. Note that the initialization_method is used in constructing the "ensemble member" called for in the DRS document; it is the value of M in $r<N>i<M>p<L>.*b) \Rightarrow$ physics_version = an integer (≥ 1) referring to the physics version used by the model If there is only one physics version of the model, then this argument should be normally given the value 1. Note that model versions that are substantially

different should be given a different "model_id"; assigning a different "physics_version" should be reserved for closely-related model versions (e.g., as in a "perturbed physics" ensemble) or for the same model, but with different forcing or feedbacks active. In CMIP5, one would distinguish, for example, among runs forced by different combinations of "forcing" agents (as called for under the "historicalMisc" experiment - experiment 7.3) by assigning different values to physics version.

For fields appearing in table "fx" in the CMIP5 Requested Output, set physics version=0 (violating the general rule that it should be a positive definite integer). Note that the physics version is used in constructing the "ensemble member" called for by the DRS document; it is the value of L in r<N>i<M>p<L> * a) ⇒ Ensemble member (r<N>i<M>p<L>) = <code>This triad of integers (N, M,L), formatted as shown above (e.g., "r3i1p21") distinguishes among closely related simulations by a single model. All three are required even if only a single simulation is performed. The so-called "realization" number (a positive integer value of "N") is used to distinguish among members of an ensemble typically generated by initializing a set of runs with different, but equally realistic, initial conditions. CMIP5 historical runs initialized from different times of a control run, for example, would be identified by "r1", "r2", "r3", etc.). The data supplier must assign a realization number to each atomic dataset. It is generally recommended that the numbers be assigned sequentially starting with 1 (but other recommendations, specified below, may override this recommendation). In CMIP5, time-independent variables (i.e., those with frequency="fx") are not expected to differ across ensemble members, so for these N should be invariably assigned the value zero ("r0"). For TAMIP ("the Transpose AMIP activity), the "realization" number is used to distinguish among the 16 members of each of 4 ensembles (one for each of 4 "seasons") generated from different observed conditions, spaced 30 hours apart. So, for example, the 16-member ensemble of runs initialized at 00Z on 15 Oct 2008, 06Z 16 Oct 2008, 12Z 17 Oct 2008, and so-on, would be assigned "r1", "r2", "r3", etc. Models used for forecasts that depend on the initial conditions might be initialized from observations using different methods or different observational datasets. These should be distinguished by assigning different positive integer values of "M" in the "initialization method indicator" (i<M>). For CMIP5



this indicator might in some cases be needed to distinguish among runs in the decadal-prediction suite of experiments (1.1-1.6). The data supplier must assign an initialization method number to each atomic dataset. It is recommended that the numbers be assigned sequentially starting with 1. In CMIP5, timeindependent variables (i.e., 6 those with frequency="fx") are not expected to differ across ensemble members, so for these M should invariably be assigned the value zero ("i0"). A key (i.e., a table) should be made available that associates each value of M with a particular initialization method and/or observational dataset. If there are many closely related model versions, which, as a group, are generally referred to as a perturbed physics ensemble (e.g., QUMP or climateprediction.net ensembles), then these should be distinguishable by a "perturbed physics" number, p<L>, where the positive integer value of L is uniquely associated with a particular set of model parameters (e.g., r3i1p78 is a third realization of the seventy-eighth version of the perturbed physics model). If there are different "forcing" combinations prescribed in experiment 7.3 in CMIP5 (the "historicalMisc" runs), then each of these different runs are also assigned different values of L (in "p<L>"). Note that the data supplier must assign a physics version number to each atomic dataset. It is recommended that the numbers be assigned sequentially starting with 1. In CMIP5, timeindependent variables (i.e., those with frequency="fx") are not expected to differ across ensemble members, so for these L should always be assigned the value zero ("p0"). A key (i.e., a table) should be made available that associates each value of L with a particular set of model parameter values and/or, in the case of the "historicalMisc" experiment, a particular suite of "forcing" agents. Note that for a single model and experiment N, M, and L should be interpretable independently; for all members of the ensemble, the correspondence between the values of N, M, and L and the simulation characteristics they represent should be consistent. For example the two different ensemble members, r3i1p7 and r3i1p8, should both be initialized from exactly the same initial conditions using the same method (because the "r" and "i" values are identical) although the subsequent evolution of the simulations will presumably differ since they were produced by two different "perturbed physics" versions of the same model. Note that there may be cases where "gaps" could occur in the list of ensemble members. If, for example, two different



initialization procedures were used, but the second procedure was tested with only a subset of the initial condition cases of the first procedure (say, every other case). Then the list of ensemble members would look like: r1i1p1, r2i1p1 r3i1p1, r4i1p1, r5i1p1, r6i1p1, ..., r1i2p1, r3i2p1, r5i2p1, ... A recommendation for CMIP5 is that each so-called RCP (future scenario) simulation should when possible be assigned the same realization integer as the historical run from which it was initiated. This will allow users to easily splice together the appropriate historical and future runs. Thus, for example, suppose a 3member ensemble of historical runs of a model exists, and a single rcp45 simulation was produced, initialized from the third member of the historical ensemble. The recp45 simulation would be designated "r3" (rather than "r1"), even though it is the only existing ensemble member, in order to indicate that it was spawned from member 3 of the historical ensemble. A similar convention should be followed, when appropriate, with other simulations (e.g., the decadal simulations).</code> ==== CMIP6 variant label ==== aka r<k>i<l>p<m>f<n> or ripf * realization index = realization number (integer >0) * initialization index = index for variant of initialization method (integer >0) * physics index = index for model physics variant (integer >0) * forcing index = index for variant of forcing (integer >0) * Note: the information stored in the forcing attribute in CMIP5 may in CMIP6 appear in the variant info attribute * variant label = a label constructed from 4 indices stored as global attributes * <code>r<k>i<l>p<m>f<n> where k = realization index I = initialization index m = physics index n = forcing index</code> * variant info = brief description of what is unique about this ripf variant * Example: "forcing: black carbon aerosol only", "realization 1", "realization 1; initialized using anomaly approach (method 2)" ==== PMIP4 and variant label notes ==== Reminder: each option in r<k>i<l>p<m>f<n> has to be a strictly positive integer === realization index r<k> === The long PMIP4 simulations are going to require both a lot of processing power and a lot of storage. It is quite likely that there will be only one realization, for a given set of i<l>p<m>f<n> and that the variant label will always start with r1 === forcing index f<n> === **Depending on available resources, the PMIP4 groups** may choose to perform several simulations for the same experiment, using different combinations of **forcings.** The forcings used will have to be carefully



2024/03/29 13:28 9/11 PMIP4 and the CMIP6 DRS

described in the documentation (and in the metadata inside each NetCDF file) and be encoded in the integer value of the forcing index. There are several ways to proceed. The easiest way is to let each group choose its own way of numbering the forcings combinations (and document it!), but all groups should try to use a common scheme for and associate the same combination of forcings with the same integer == **Sequential numbering scheme == The contact people** for each experiment determine which forcings combinations are most likely to be used and associate them with a predefined number. If necessary, a group can later ask for a new forcing combination to be registered ^ Forcings ^ fforcing index ^ | Recommended default, or most likely combination, or mandatory simulation | f1 | | forcing1='on', forcing2='off', etc | f2 | | Some other combination | fN | == Hierarchical numbering scheme == The following scheme will create bigger integers, but the values will be more meaningful If there are 10 or less options for each type of forcing, we can assign a power of 10 to each type, multiply it with the forcing option and add everything Tentative example for the Igm experiment: ^ Power ^ Forcing ^ Options ^ | 2 | Ice sheet | 1=*ICE*-6*G*-*C* 2=GLAC-1D | | 1 | Aerosols | 1=Hopcroft et al 2=Albani et al | 0 | Vegetation | 1=interactive vegetation 2=interactive carbon cycle 3=prescribed | Example: GLAC-1D + Hopcroft et al + interactive vegetation = $2 * 100 + 1 * 10 + 1 \Rightarrow$ f211 ==== variant label constraints for PMIP4 experiments ==== === historical === historical simulations that are the continuation of a past1000 simulation should use 1000 for the initialization method ⇒ i1000 === past1000 === **Prix Me!** === mid-Holocene === **Prix Me!** === lgm === **Prix Me!** === lig127k === **Prix Me!** === === **Prix Me!** === **LD**-transpin === **Prix Me!** === LD === Fix Me! ==== PMIP4-CMIP6 directory structure and file names ==== The DRS defines (among other things) how the different attributes will be combined to generate unambiguous directories and file names, in the ESGF distributed database <code>Directory structure = <mip era>/ <activity id>/ <institution id>/ <source id>/ <experiment id>/ <member id>/ ←= variant label <table_id>/ <variable_id>/ <grid_label>/ <version>/ file name = <variable id> <experiment id

>_<source_id>_<member_id>_<grid_label>[_<time_ra
nge>].nc </code> For PMIP4, we have
sub_experiment_id == none (because we don't use
forecast and hindcast), and therefore member_id ==
variant_label ^ Used in
dir? ^ Used in
file? ^ Attribute
name ^ Value for PMIP4-CMIP6 ^ | Y | N | mip_era |
CMIP6
PMIP4 ? | | Y | N | activity_id | CMIP
PMIP

Note: "CMIP PMIP" becomes CMIP (If multiple activities are listed in the global attribute, the first one is used in the directory structure) | | Y | N | institution_id | institution label (IPSL, ...) | | Y | N | version | vYYYYMMDD (e.g., v20160218), indicating a representative date for the version

Note: the *version* is not stored in the NetCDF files and not** used in the file names, because it is only specified when *publishing* (eg storing the data in ESGF) the NetCDF files |



Υ	Y	source_id	source label (e.g. the model name/version using only authorized characters)
Υ	Υ	experiment_id	See the Experiment names section
Y	Υ	member_id	PMIP4 does not use sub_experiment_id, so the value of member_id is equal to the variant_label: r <k>i<l>p<m>f<n> (see the CMIP6 variant label section)</n></m></l></k>
Υ	Υ	table_id	CMOR table label (Amon,)
Υ	Υ	variable_id	variable identifier (tas, pr,)
Υ	Υ	grid_label	gn: output is reported on the native grid gr: output is regridded by the modeling group to a "primary grid" of its choosing gr1, gr2,: output is regridded on another grid than the primary grid (that was already different from the native grid)
N	Υ	time_range	the last segment of the file name indicates the time-range spanned by the data in the file, and is omitted when inappropriate. The format for this segment is the same as in CMIP5

Examples:

• Directory =

https://pmip4.lsce.ipsl.fr/

2024/03/29 13:28 11/11 PMIP4 and the CMIP6 DRS

CMIP6/CMIP/NCAR/CCSM2-1/1pctC02/rli1p1f1/
Amon/tas/gn/v20150320/

File =

tas_Amon_CCSM2-1_1pctC02_r1i1p1f1_gn_2020 01-202912.nc



 Directory = CMIP6/DCPP/NCAR/CCSM2-1/dcppA-hindcast/s1960r1i2p1f1/Amon/tas/gr/v20150320/ File = tas_Amon_CCSM2-1_hindcast_s1960r1i2p1f1 gn 198001-198412.nc

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